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**The enigmatic morula: mechanisms of development, cell fate determination, self-correction and implications for ART**

Running title: The morula: implications for development and ART

Giovanni Coticchio<sup>1,3</sup>, Cristina Lagalla<sup>1</sup>, Roger Sturmey<sup>2</sup>, Francesca Pennetta<sup>1</sup>, and Andrea Borini<sup>1</sup>

<sup>1</sup> 9.Baby, Family and Fertility Center, Via Dante 15, 40125 Bologna, Italy

<sup>2</sup> Centre for Atherothrombosis and Metabolism, Hull York Medical School, University of Hull, Hull, United Kingdom

<sup>3</sup> Corresponding author. Email: [giovanni.coticchio@9puntobaby.it](mailto:giovanni.coticchio@9puntobaby.it)

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40 **Background:** Assisted reproduction technology offers the opportunity to observe the very  
41 early stages of human development. However, due to practical constraints, for decades  
42 morphological examination of embryo development has been undertaken at a few isolated  
43 time points at the stages of fertilisation (day 1), cleavage (day 2-3) and blastocyst (day 5-6).  
44 Rather surprisingly, the morula stage (day 3-4) has been so far neglected, despite its  
45 involvement in crucial cellular processes and developmental decisions.

46  
47 **Objective and rationale:** The objective of this review is to collate novel and unsuspected  
48 insights into developmental processes occurring during formation of the morula, highlighting  
49 the key importance of this stage for a better understanding of preimplantation development  
50 and an improvement of ART.

51  
52 **Search method:** PubMed was used to search the MEDLINE database for peer-reviewed  
53 English-language original articles and reviews concerning the morula stage in mammals.  
54 Searches were performed by adopting 'embryo', 'morula', 'compaction', 'cell fate' and  
55 'IVF/assisted reproduction' as main terms, in association with other keywords expressing  
56 concepts relevant to the subject (e.g. cell polarity). The most relevant publications, i.e. those  
57 concerning major phenomena occurring during formation of the morula in established  
58 experimental models and the human species, were assessed and discussed critically.

59  
60 **Outcomes:** Novel live cell imaging technologies and cell biology studies have extended our  
61 understanding of morula formation as a key stage for the development of the blastocyst and  
62 determination of the inner cell mass (ICM) and the trophectoderm (TE). Cellular processes,  
63 such as dynamic formation of filopodia and cytoskeleton-mediated zippering cell-to-cell  
64 interactions, intervene to allow cell compaction (a geometrical requisite essential for  
65 development) and formation of the blastocoel, respectively. At the same time, differential  
66 orientation of cleavage planes, cell polarity and cortical tensile forces interact and cooperate  
67 to position blastomeres either internally or externally, thereby influencing their cellular fate.  
68 Recent time lapse microscopy (TLM) observations also suggest that in the human the  
69 process of compaction may represent an important checkpoint for embryo viability, through  
70 which chromosomally abnormal blastomeres are sensed and eliminated by the embryo.

71  
72 **Wider implications:** In clinical embryology, the morula stage has been always perceived  
73 as a 'black box' in the continuum of preimplantation development. This has dictated its virtual

74 exclusion from mainstream ART procedures. Recent findings described in this review  
75 indicate that the morula, and the associated process of compaction, as a crucial stage not  
76 only for the formation of the blastocyst, but also for the health of the conceptus. This  
77 understanding may open new avenues for innovative approaches to embryo manipulation,  
78 assessment and treatment.

79

80

81 **Keywords:** morula; compaction; embryo; inner cell mass; trophectoderm.

82 **Introduction**

83

84 In assisted reproduction technology (ART), embryos are routinely assessed and selected  
85 for transfer at isolated time points on day 2, 3 and, in case of culture to the blastocyst stage,  
86 day 5/6. Embryos may also be observed, but are rarely selected, on day 4, since embryo  
87 transfer (ET) at this stage does not imply the practical advantages of transfer on days 2-3  
88 (reduced costs and management) or day 5/6 (higher self-selection and increased cycle  
89 efficiency). Day 4 ET is also hindered by difficulties in assessing accurately the morphology  
90 of embryos at this stage of development.

91

92 In normal undisturbed development, day 4 embryos have already started, and have often  
93 completed, the transition from the cleavage stages, in which blastomeres appear as well-  
94 distinct approximately spherical units, to the compacted morula stage, characterised instead  
95 by tightly interconnected cells with ill-defined margins, localised either internally or externally  
96 in the embryo (ESHRE Atlas of Human Embryology). At the sub-cellular level, this phase of  
97 development is also when the blastomeres undergo cell polarisation. Cells localised  
98 internally are characterised by a nucleus positioned approximately centrally and organelles  
99 and other specialised structures distributed in the cell environment without an apparent  
100 pattern. Cells distributed externally respond instead to a more defined pattern: the nucleus  
101 is positioned basally, cell adhesion structures and stable acetylated microtubules are  
102 localised baso-laterally, while more dynamic microtubules, actin filaments and microvilli are  
103 found in the apical region. The regulation of the process of polarisation remains unclear,  
104 with significant efforts on unpicking the origin of cell polarity in mammalian embryos.  
105 Furthermore, the chromosomal status of the day 4 embryo has received little attention,  
106 despite the hypothesis that during development from the cleavage to the blastocyst stage,  
107 mosaic embryos can implement mechanisms of self-correction to reduce the aneuploidy  
108 load.

109

110 Being a developmental phase less amenable to observation and morphological  
111 classification, knowledge of the human embryo on day 4 is limited compared to other stages  
112 of development, especially concerning the relationship between morphology and  
113 developmental ability. Consequently, this review will highlight the relevance of the morula  
114 relative to developmental processes that occur during preimplantation development and  
115 discuss downstream implications of such processes for practices used in human ART.

116

**117 Methods**

118

119 PubMed was systematically searched for peer-reviewed original articles and reviews  
120 identified by relevant keywords, such as ‘embryo’, ‘blastomere’, ‘morula’, ‘compaction’,  
121 ‘blastocyst’, ‘blastocoel’, ‘cell fate’, ‘cell lineage’, ‘cell determination’, ‘cell polarity’, ‘cell  
122 junctions’, ‘IVF’, ‘assisted reproduction’, ‘inner cell mass’ and ‘trophectoderm’. Keywords  
123 were used in multiple and overlapping combinations in order to identify those publications  
124 strictly relevant to the morula. Additional studies were identified by thorough analysis of  
125 reference lists from relevant publications. The ‘English language’ limit was applied. The most  
126 relevant publications, i.e. those concerning major phenomena occurring during the  
127 developmental morula stage were assessed and discussed critically to offer a  
128 comprehensive description of the process bridging early segmentation and the blastocyst.  
129 Concerning animal studies, priority was given to publications relevant to species more  
130 consistently used as experimental models for the human.

131

**132 The morula stage in the continuum of preimplantation development**

133

134 The crucial phases of preimplantation development have been described in meticulous  
135 detail; days 1 (fertilisation and first cleavage), days 2-3 (second and third rounds of  
136 cleavage) and days 5-7 (blastocyst stage) of human development have been assessed  
137 morphologically in thousands of studies. In contrast, the morula stage has received  
138 astonishingly little attention, as illustrated by an authoritative source describing the morula  
139 simply as “an indistinguishable mass of cells on day 4 of development” (ESHRE Atlas of  
140 Human Embryology). In the human, compaction typically occurs usually between the 8- and  
141 10-cell stage, although sporadically can be observed as early as the 4-cell stage (Iwata *et*  
142 *al.*, 2014). Qualitative assessment of the morula also appears rather vague; a “good quality  
143 morula” is defined as “composed of 16-32 blastomeres and all the blastomeres should be  
144 included in the compaction process” (ESHRE Atlas of Human Embryology). Exclusion of  
145 one or more cells from the compacted morula is thought to be associated with reduced  
146 developmental competence (Ebner *et al.*, 2009). However, the morula remains poorly  
147 described. This is due in part to the lack of well-defined morphological markers that can be  
148 considered typical of this stage. In addition, compaction and formation of the morula are  
149 dynamic processes, difficult to assess in IVF routine practice if observation is carried out at

150 isolated time points. Nevertheless, overlooking this "Cinderella" stage of development is  
151 naïve, since recent data indicate that the morula is pivotal for blastocyst formation,  
152 establishment of the first cell lineages and, therefore, the entire developmental process. This  
153 role is underpinned by molecular, metabolic and cellular changes, which are temporally and  
154 spatially regulated, and a transition from a group of individual cells to a co-ordinated,  
155 responsive structure (Brison *et al.*, 2014). Stage-specific modulations of gene expression  
156 and metabolism are indicative of such changes.

157

158 The early stages of embryo development are largely under the molecular control of  
159 maternally-inherited mRNAs and regulatory proteins stockpiled throughout oogenesis. The  
160 function of genes, referred to as "maternal effect genes", encoding for such regulators was  
161 initially ascertained in invertebrates and lower vertebrates, but knowledge of their action in  
162 mammals, including the human, is emerging (Condic, 2016). This initial maternal control of  
163 development is gradually superseded as the embryo expresses its own genes. The first  
164 studies in the human reported that embryonic genome activation (EGA) began at 4 to 8-cell  
165 stage (Braude *et al.*, 1988), but details of the overall dynamics of gene expression during  
166 preimplantation development remained unclear for several years. More recently,  
167 sophisticated DNA technologies have revealed a more detailed and complex picture of EGA,  
168 which has relevance for the morula stage. Briefly, we now know that the first signs of  
169 transcription in the human embryonic genome are detectable at the 2-cell stage, much  
170 earlier than previously thought. Notably, analysis of the whole preimplantation period  
171 showed that EGA occurs in a multiphasic fashion at designated stages: a pattern seen  
172 mirrored in a range of mammalian embryos (Svoboda *et al.*, 2015). A major wave of  
173 transcription activation occurs at the 4-6 cell stage, involving mainly genes with a role in the  
174 translation machinery. This sets the stage for the two following and more extensive  
175 transcriptional bursts, which take place towards the end of day 3 of development (from the  
176 8-10 cell stage interval), i.e. shortly before compaction of the morula (Vassena *et al.*, 2011).  
177 Many genes transcribed in these phases encode for proteins related to the metabolism of  
178 lipids, proteins, amino acids and carbohydrates. Embryo compaction is therefore preceded  
179 by changes in gene expression that are suggestive of major shifts in embryo metabolism.

180

181 Indeed, it is well documented both in model species and the human that during  
182 preimplantation development, metabolism undergoes important phase-dependent  
183 modifications in response to specific requirements of the embryo (Leese, 2012). During the

184 early cleavage stages, the embryo is comparatively quiescent (Smith and Sturmey, 2013).  
185 Cell divisions occur at a moderate pace, net mass does not increase and most of the energy  
186 requirements are focused on general homeostatic functions, with limited need for  
187 biosynthetic processes. Consequently, metabolic needs, in terms of ATP during the  
188 cleavage stages, are satisfied largely from low levels of oxidation of pyruvate and lactate,  
189 which are abundantly present in the surrounding environment (Smith and Sturmey, 2013).  
190 In addition, a high ATP:ADP ratio negatively regulates the flux of glucose through the  
191 glycolytic pathway prior to compaction (Rozell *et al.*, 1992). This picture changes in the  
192 second half of preimplantation development, corresponding to a period when gene  
193 transcription is more sustained, the rate of cell division increases and embryo mass  
194 increases. Importantly, considerable energy is invested in Na<sup>+</sup>/K<sup>+</sup> ATPase, an enzyme  
195 needed for an active transport of those ions from the surrounding environment to intercellular  
196 spaces (Martin and Leese, 1999; Houghton *et al.*, 2003). A high ion concentration in the  
197 intercellular space facilitates movement of water down a concentration gradient from the  
198 extraembryonic environment, leading ultimately to the formation of the blastocoel (Watson  
199 and Barcroft, 2001). The energy demands for such activities are satisfied by a marked  
200 increase in metabolic function. The rate of oxygen consumption rises significantly, as a  
201 range of metabolic fuels including pyruvate, glutamine and fatty acids are oxidised  
202 (Houghton *et al.*, 1996)(Thompson *et al.*, 1996)(Houghton *et al.*, 2003)(Sturmey and Leese,  
203 2003)(Lopes *et al.*, 2005). In parallel, there is a marked, significant rise in glucose  
204 consumption via aerobic glycolysis (Gardner *et al.*, 2001)(Krisher and Prather, 2012).  
205 Importantly, aerobic utilisation of glucose can support multiple biosynthetic pathways, as  
206 well as providing ATP (Krisher and Prather, 2012). In addition to aerobic glycolysis, the  
207 increased uptake of glucose is important for sustaining the Pentose Phosphate Pathway,  
208 essential for the provision of ribose sugars necessary for nucleotide synthesis (Smith and  
209 Sturmey, 2013). Crucially, genes whose products, such as the pyruvate dehydrogenase  
210 complex, functionally linking glycolysis to the activity of the oxidative TCA cycle, may be  
211 regulated at the transcriptional level and expressed in a timely fashion. Therefore, as the  
212 embryo evolves from a low-energy, low activity, almost automatic system to one that is  
213 highly energetic, highly proliferative and developmentally active, the morula stage  
214 represents a key regulatory landmark during which a pyruvate-based metabolism  
215 characterised by low oxidative levels gives way to a more active aerobic utilisation of  
216 glucose, combined with other metabolic fuels, in response to modified developmental needs.  
217 Unfortunately, detailed knowledge of metabolism of the morula per se is sparse, compared



218 to the wealth of data on cleavage and blastocyst stage embryos, mostly due to the  
219 challenges described above.

220

### 221 **Inner/outer cells and morula geometry**

222

223 During preimplantation development, the cells of the embryo make their first developmental  
224 decision, i.e. whether to contribute to the formation of the inner cell mass (ICM) or the  
225 trophoctoderm (TE) of the blastocyst. After implantation, the former will develop mainly into  
226 the soma of the future organism, while the latter will give rise to extra-embryonic tissues  
227 supporting the embryo proper. Classical studies carried out in the mouse have suggested  
228 the concept (referred to as inside-outside theory) that, as cleavage progresses, blastomeres  
229 that become totally internalised within the structure of the embryo will exclusively form the  
230 ICM, while blastomeres that remain external and exposed to the surrounding environment  
231 will contribute mainly to the TE. These conclusions were drawn by simple, but ingenious,  
232 experiments in which outer cells of 16-cell mouse embryos were labelled with fluorescent  
233 microparticles and labelling patterns of ICM and TE were observed following development  
234 of morulae into blastocysts (Fleming, 1987). However, the precise timing and mechanism of  
235 the fate decision remains unclear and answers to the questions of how blastomeres acquire  
236 an inner or outer position, how inner and outer cells become different and how positional  
237 cues are translated into developmental decisions still elude us. In a fascinating recent study,  
238 Biase *et al.* (2018) used an RNASeq approach, combined with cellular barcoding, to indicate  
239 that lineage differences were apparent as early as the two-cell stage in mouse embryos,  
240 indicating that the fate of the morula may be determined after the first cleavage. The morula  
241 and the process of compaction are particularly relevant to such questions, because they  
242 represent the functional manifestation of repositioning the inner and outer cells (as  
243 discussed below).

244

245 The creation of the morula presents a geometrical challenge, in which cells are required to  
246 distinctly occupy internal or external positions. Up until the third round of divisions, all cells  
247 (up to eight) are approximately spherical and are all exposed to the surrounding  
248 environment, in addition to being in contact with each other. During the fourth round of  
249 cleavage, cell number increases from 8 to 16, and, on average, 5 blastomeres become  
250 positioned in the interior of the mouse embryo (Morris *et al.*, 2010). If blastomeres were to  
251 retain a spherical shape and remain in mutual contact, for mere geometrical rules (the

252 Newton's kissing number), only a single cell would find allocation internally in the embryo at  
253 the 13-cell stage (White *et al.*, 2017). Beyond such a stage, the addition of further cells in  
254 the interior of the embryo would be impossible while maintaining blastomere and embryonic  
255 sphericity and cell-to-cell contact. In the mouse, compaction occurs normally between the  
256 8- and 16-cell stage, concomitantly with the formation of inner and outer cells. Cell  
257 compaction is therefore the geometrical, functionally-essential, solution that permits the  
258 development of populations of inner and outer cells in a definite proportion and to preserve  
259 its overall shape of a homogeneous sphere within the constraints of the zona pellucida. In  
260 the majority of cases in the human, compaction also occurs during the same developmental  
261 interval (Iwata *et al.*, 2014), although the overall process of commitment of inner and outer  
262 cells in the formation of ICM and TE, respectively, is less certain (see below).

263

### 264 **The forces that shape the morula in the process of compaction**

265

266 Geometr is however only one of the several obstacles that the developing embryo has to  
267 overcome to ensure the generation of two cell lines. One of the most topical questions that  
268 concern the morula stage and indeed preimplantation development revolves around the  
269 forces that make compaction possible. Until the 8-cell stage, cleavage unfolds apparently  
270 uneventfully. Compaction marks a dramatic change in cell behaviour and shape, so  
271 profound that it can be observed by a simple transmitted light microscope. At compaction,  
272 blastomeres lose their sphericity to acquire a rather flattened epithelial-like shape in which  
273 cell contour is difficult to discriminate. This has prompted the hypothesis that cellular  
274 specialisations that ensure cell-to-cell contact and adhesion play an essential role in  
275 compaction. Early investigations on the Ca<sup>2+</sup>-dependent cell adhesion molecule E-cadherin  
276 were consistent with this hypothesis. In the mouse, E-cadherin is uniformly distributed in the  
277 cell membrane during the first cleavages but from the 8-cell stage, its localisation becomes  
278 restricted to intercellular adherens junctions, also referred to as zonulae adherens  
279 (Vestweber *et al.*, 1987), which ensure mutual lateral adhesion between epithelial cells and  
280 contribute to the maintenance of epithelial cell polarity (Table 1). In the human, E-cadherin  
281 follows a similar pattern of redistribution, with a preferential localisation in adherens junctions  
282 from day 3.5-4.0 of development (Campbell *et al.*, 1995). The initial finding that mouse  
283 compacted morulae exposed to functionally interfering anti-E-cadherin antibodies undergo  
284 decompaction (Vestweber and Kemler, 1985) inspired subsequent gene targeting studies.  
285 Indeed, mouse morulae homozygous for a null mutation of the E-cadherin-encoding gene

286 were unable to persist in a compacted state, progressively losing cohesiveness and  
287 reverting to a decompacted morphology, with cells still able to cleave but not mutually  
288 adhesive and unable to organise themselves into blastocyst (Riethmacher *et al.*, 1995). One  
289 explanation for this is that the early embryo is endowed with E-cadherin molecules of  
290 maternal origin that can initiate compaction, but new, zygotic-derived E-cadherin is  
291 necessary to complete the process. Interestingly, loss of E-cadherin function and  
292 compaction is accompanied by intracellular alterations, particularly the loss of cell polarity  
293 (Riethmacher *et al.*, 1995). However, compaction and cell polarity are not strictly mutually  
294 dependent because under specific experimental conditions, one process can occur in the  
295 absence of the other (Stephenson *et al.*, 2010).

296

297 In the mouse, other molecules interacting with E-cadherin in the formation of adherens  
298 junctions have been implicated in the process of compaction, further supporting a role for  
299 cell adhesion in this developmental phase. The intracellular domain of E-cadherin is  
300 connected with  $\beta$ - or  $\gamma$ -catenin. This complex is in contact with  $\alpha$ -catenin, through which it is  
301 anchored to the actin cytoskeleton (Perez-Moreno *et al.*, 2003). In mice, experimental  
302 ablation of  $\alpha$ -catenin produces a phenotype similar to that of the previously described E-  
303 cadherin null mutants, with embryos able to initiate but not complete compaction (Torres *et*  
304 *al.*, 1997). By contrast, in  $\beta$ -catenin knockout embryos, compaction occurs unperturbed, with  
305 developmental anomalies emerging only at gastrulation, perhaps as an effect of redundancy  
306 due to the presence of other catenins or protracted action of maternal molecules (Haegel *et*  
307 *al.*, 1995). Ephitin, a transmembrane serine protease, is another example of a cell adhesion  
308 molecule that appears to be involved in compaction. It co-localises with E-cadherin at areas  
309 of the cell membrane of contact between blastomeres of compacted mouse 8-cell embryos.  
310 When its expression is ablated in RNAi experiments, cell adhesion is lost and embryo death  
311 occurs shortly afterwards (Khang *et al.*, 2005). Importantly, and unlike the mouse, the culture  
312 conditions may also affect the stability and function of junctional complexes occurring  
313 between blastomeres in human embryos, with implications for the process of compaction  
314 and ultimately embryo viability in vitro (Eckert *et al.*, 2007).

315

316 Recent advances in in-vivo labelling and time lapse imaging techniques have extended  
317 our understanding of the mechanisms of compaction and the role of E-cadherin and  
318 associated molecules. It is now possible to view E-cadherin distribution in live cells, by  
319 microinjecting RNA molecules encoding E-cadherin with a GFP fluorescent tag. This

320 approach has revealed a complex and regulated cell-to-cell interaction occurring at the time  
321 of compaction (Fierro-González *et al.*, 2013). Before the 8-cell stage, E-cadherin is found  
322 throughout the cell membrane, although particularly abundant at adherens junctions. During  
323 the 8- to 16-cell stage interval, concomitant with compaction, E-cadherin-rich filopodia  
324 become visible. Originating from an area situated between the adherens junctions and the  
325 cell apical domain, these filopodia have a length of 10-12  $\mu\text{m}$  and extend over the apical  
326 membrane of adjacent cells. These blastomere protrusions are positive for F-actin and  
327 Myo10, but do not appear to contain  $\alpha$ -tubulin, consistent with typical filopodia structure  
328 (Table 1). Detailed 4D analysis during the 8- to 16-cell interval unveiled an astonishing  
329 regulation of filopodia (Fierro-González *et al.*, 2013). They are detected only in 55-60% of  
330 all blastomeres. Each blastomere projects 5-6 filopodia over the apical membrane of 2-3  
331 adjacent cells; however, adjacent blastomeres never project filopodia reciprocally. Filopodia  
332 are dynamic, appearing to extend and retract in coordination with cell division. Remarkably,  
333 they are retracted before a cell division is initiated, while cells receiving filopodia do not  
334 divide as long as filopodia extend over their membranes. In addition, approximately two  
335 thirds of blastomeres with filopodia undergo symmetric division and remain to the outer of  
336 the embryo organization, while the remaining one third divide asymmetrically giving rise to  
337 one inner and one outer cell. Notably, such a 7:3 ratio of symmetric/asymmetric divisions is  
338 believed to occur during compaction (Morris *et al.*, 2010). The observation that, before  
339 division, retraction of filopodia is followed by a change in cell shape from elongated to  
340 rounded, has inspired further experiments confirming a causal role of filopodia in the process  
341 of compaction. When filopodia were ablated by laser micromanipulation, the juxtaposed  
342 membranes of two adjacent cells retract immediately, indicating that filopodia impose  
343 mechanical forces between cells that can influence membrane tension and therefore cell  
344 shape. The same effect was not observed when adherens junctions were ablated (Fierro-  
345 González *et al.*, 2013).

346

347 Molecular manipulations have confirmed a role for filopodia in determining cell elongation  
348 behaviours required by compaction. In effect, mouse blastomeres microinjected with E-  
349 cadherin short interfering RNA (siRNA) form few filopodia, remain rounded and fail to  
350 integrate in a compacted structure formed by non-injected cells. Knock-down of  $\alpha$ -catenin  
351 and  $\beta$ -catenin, which co-localise with E-cadherin in filopodia, also cause a drastic reduction  
352 in filopodia number and inability of the knocked-down blastomeres to elongate and compact  
353 (Fierro-González *et al.*, 2013). Together, these findings confirm an instrumental role for E-

354 cadherin and associated proteins in embryo compaction, while indicating filopodia, but not  
355 adherens junctions, as the prominent cell specialisation required for a change in cell shape  
356 from rounded to elongated.

357

358 Collectively, this evidence indicates a critical role for cell adhesion in the process of  
359 compaction. However, cell adhesion appears to have other significant implications for the  
360 function of the morula and initiation of blastocyst formation. In particular, by preventing the  
361 free diffusion of membrane proteins, adherens junctions maintain the integrity of the baso-  
362 lateral and apical membrane domains. Such a difference represents a prominent aspect of  
363 cell polarity. Differentiations in the composition and function of cell membrane domains are  
364 central for the mechanism of accumulation of intercellular fluid which leads to formation of  
365 the blastocoel. As a further consequence of the localisation and function of adherens  
366 junctions, the Na<sup>+</sup>/K<sup>+</sup> ATPase system is localised only at the baso-lateral membrane domain  
367 of outer cells of mouse embryos (Watson and Kidder, 1988). By hydrolysis of one  
368 molecule of ATP, the enzyme transports three Na<sup>+</sup> ions out of the cell, in exchange for two  
369 K<sup>+</sup> ions. This allows the establishment of an ion concentration gradient between the  
370 intercellular spaces of the morula, which become enriched in Na<sup>+</sup> ions, and the surrounding  
371 environment. As a result, and assisted by increased and coordinated expression of  
372 aquaporins (Barcroft *et al.*, 2003), water flows unidirectionally from the exterior through the  
373 epithelial-like outer cells, accumulating extracellularly and thereby forming the blastocoel  
374 (Watson and Barcroft, 2001). Concomitantly at this stage, E-cadherin-dependent formation  
375 of tight junctions (zonulae occludens) creates a belt-like impermeable structure located at  
376 the upper-lateral sides of adjacent outer cells (Table 1). This specialisation of cell-to-cell  
377 contact contributes to the retention of water in the intercellular spaces of the embryo, as  
378 shown in the mouse, the human and other species (Eckert and Fleming, 2008). Live cell  
379 imaging used to study the development of mouse embryos has revealed that the creation of  
380 a barrier that seals the interior of the mouse embryo from the outside is assisted by an actin-  
381 zipper mechanism. Before formation of the blastocoel, actin-rich rings form at the apical  
382 side of outer cells. These rings do not have contractile ability but instead expand in size and  
383 diameter until they reach cell-to-cell areas of contact. At this level, they interact with and  
384 stabilise cell junctions by recruiting the relevant components. This is followed by focused  
385 association with myosin II (Table 1). In this fashion, the newly formed actomyosin ring  
386 localised at the margins of adjacent cells produces the tension forces required for the

387 zippering mechanism that acts in coordination with cell junctions to seal the intercellular  
388 contact between outer cells (Zenker *et al.*, 2018).

389

### 390 **Cellular arrangements in the nascent morula**

391

392 If indeed compaction, encompassing cell elongation and flattening, represents cellular  
393 "trickery" to accommodate inner and outer cells in the structure of the morula beyond the 8-  
394 cell stage, it poses the question of how these two cell populations are generated in the first  
395 place. Collective evidence suggests that modalities by which inner and outer cell are  
396 generated are multiple and probably not mutually exclusive. One such modality, the first to  
397 be identified in studies on mouse embryo development, is based on two major elements, i.e.  
398 cell polarity and symmetry of cell division (Mihajlović and Bruce, 2017). As mentioned above,  
399 between the 8-cell and the 16-cell stage, blastomeres acquire a well-defined polarised  
400 organisation, with nuclear (Reeve and Kelly, 1983), cytoskeletal (Johnson and Maro, 1984;  
401 Houlston and Maro, 1989) and cell membrane components (Fleming and Pickering, 1985;  
402 Korotkevich *et al.*, 2017) distributed differently in the baso-lateral (internal) and apical  
403 (external) domains. Blastomere polarisation was also observed in human embryos from the  
404 8-cell stage (Nikas *et al.*, 1996). Different orientations of cleavage planes will therefore  
405 determine alternative destinies of daughter cells (Johnson and Ziomek, 1981a). Planes  
406 oriented parallel to the cell baso-apical axis will produce two symmetric daughter cells, both  
407 remaining in an external position and inheriting approximately the same polarised  
408 organisation of their progenitor cell. On the contrary, planes oriented orthogonally to the cell  
409 baso-apical axis will generate two asymmetric cells, one external cell inheriting the  
410 organisation and molecules of the apical domain and one internal cell characterised by baso-  
411 lateral attributes of the mother cell. These two types of cell division are also referred to as  
412 conservative and differentiative, respectively, depending on whether they preserve and  
413 reproduce a pre-existing cellular status or generate diversity in the position (external or  
414 internal) and inheritance of cellular organisation (apical or baso-lateral) between daughter  
415 cells (Sutherland *et al.*, 1990). Recent studies (Korotkevich *et al.*, 2017) indicate that  
416 orientation of the cleavage plane at the 8-cell stage is not a random event, but rather seems  
417 regulated by innate cell contact-independent factors. For example, in isolated blastomeres  
418 of mouse embryos which already show signs of cell polarisation, the mitotic spindle aligns  
419 preferentially to the basal-apical axis producing asymmetric cells in more than 80% of cases  
420 (Korotkevich *et al.*, 2017). In addition, experimental disruption of the apical domain in

421 isolated blastomere of 8-cell embryos is associated with a random spindle (and cleavage  
422 plan) orientation, suggesting that the cortical domain can regulate spindle positioning and  
423 ultimately symmetric or asymmetric division. This hypothesis is in line with micro-  
424 transplantation experiments in which the cortical domain of polarised cells integrated in non-  
425 polarised cells was able to induce asymmetric cell division (Korotkevich *et al.*, 2017). The  
426 polarised cortical domain seems therefore not only necessary, but also sufficient to control  
427 the orientation of the cleavage plan and the occurrence of asymmetric division.  
428 Consequently, the geometry of cell division at the 8- 16-cell stage introduces a new  
429 structural dimension (inner-outer) in the multicellular organisation of the mouse embryo and  
430 creates diversity among blastomeres (Johnson, 2009). Notably, asymmetric division  
431 unequally redistributes cell fate determining factors (localised especially in the apical  
432 domain) in inner and outer cells, making them not only positionally, but also functionally,  
433 different (see below). Importantly, the frequency of symmetric and asymmetric divisions has  
434 implications for the relative proportion of inner and outer cells and, as a consequence,  
435 abundance of ICM and TE cells in the ensuing blastocyst (Bischoff *et al.*, 2008). Studies on  
436 isolated blastomeres of 8-cell mouse embryos suggest that the probability by which  
437 symmetric and asymmetric divisions can occur may be an intrinsic cell characteristic at this  
438 stage (Johnson and Ziomek, 1981b). In particular, symmetric divisions, which preserve  
439 polarity in both daughter cells, are more likely to occur in blastomeres with larger apical  
440 domains (Pickering *et al.*, 1988) and higher levels of expression of apical-specific  
441 determinants.

442

443 More recent studies on cell allocation in the mouse embryo have shown that the geometry  
444 of cell division (symmetric/asymmetric) is not the exclusive, and perhaps not even the most  
445 important, modality by which the two populations of inner and outer cells are formed. Such  
446 studies have been made possible by an advanced and highly sophisticated live imaging  
447 technique. This approach involves embryos that are microinjected with RNA encoding for a  
448 fluorescent protein (mCherry) targeting the cell membrane. The emitted fluorescence can  
449 be then detected by two-photon confocal microscopy and used to track cell positioning in  
450 4D at a high resolution, following analysis by a methodology referred to as computational  
451 membrane segmentation (Fierro-González *et al.*, 2013). Mouse embryos at the 8 to 16-cell  
452 stage studied with this approach showed unexpected properties (Samarage *et al.*, 2015). In  
453 the first place, it was observed that more than 80% of inner cells derive from symmetric  
454 divisions and that 60% of embryos produce inner cells without undergoing asymmetric

455 divisions. Therefore, asymmetric division, although important in the process of cell allocation  
456 as demonstrated by early studies, is not a frequent or "sine qua non" condition to determine  
457 the position of the first inner cells to be formed. Rather, computational membrane  
458 segmentation has revealed a different morphogenetic mechanism causing cell  
459 internalisation. Starting from the 12-cell stage, some cells undergo a well-defined change in  
460 shape that involves constriction of the apical portion and expansion of the baso-lateral  
461 domain, with a consequent gradual repositioning towards the geometrical centre of the  
462 embryo. Once positioned centrally, after the 16-cell stage these cells divide to increase their  
463 number, although more inner cells can be generated through the classical mechanism of  
464 asymmetric division (Samarage *et al.*, 2015).

465

466 Further observations clarified how apical constriction can occur in cells destined to  
467 internalisation (Samarage *et al.*, 2015). In principle, cell constriction can result from different  
468 morphogenetic forces acting between adjacent cells, such as adhesion, involving E-  
469 cadherin, and cortical tension, derived by contractility of actomyosin structures. In 8- to 16-  
470 cell mouse embryos, E-cadherin is localised preferentially in the baso-lateral region of all  
471 cells, while no differences in its distribution are seen in the apical domain of constricting and  
472 non-constricting cells (Fierro-González *et al.*, 2013). Modulation in cell-to-cell adhesion is  
473 therefore unlikely to generate apical constriction. On the contrary, myosin II, but not actin,  
474 distribution can differ between adjacent cells (Figure 1). In particular, Myosin II is more  
475 abundant at the borders between constricting and non-constricting cells, tending to  
476 accumulate as apical constriction progresses (Samarage *et al.*, 2015). Micromanipulation  
477 experiments involving laser ablation of areas of the actomyosin organisation situated at the  
478 margins between constricting and non-constricting cells have produced data informing on  
479 directionality and magnitude of tensile interactions. Such information is consistent with a  
480 biomechanical model, according to which forces acting at the border of constricting cells  
481 promote their internalisation, while forces surrounding the same cells act against apical  
482 constriction and cell internalisation (Table 1). Myosin II is essential in this mechanism. This  
483 is shown by the evidence that, when Myosin II is experimentally downregulated in some, but  
484 not all, cells of an 8- to 16 cell embryo, control cells with normal levels of Myosin II fail to  
485 constrict and undergo internalisation if they are delimited by three or more knockdown cells.  
486 Therefore, tensile forces acting in non-constricting cells are also important for internalisation  
487 (Samarage *et al.*, 2015).

488



489 The overall choreography by which cell shape changes to allow the accommodation of an  
490 adequate number and positioning of inner and outer cells in the growing mouse morula is  
491 therefore complex. Adherens junctions and filopodia drive modification of the cell shape from  
492 approximately spherical, until the 8-cell stage, to elongated and flattened at later stages to  
493 overcome the geometrical limitations imposed by the Newton's kissing number. At the  
494 beginning of compaction, opposite tensile forces produced by actomyosin accumulation at  
495 the cell borders induce some cells to undergo apical constriction coupled to basal  
496 broadening. As a result of these changes, such cells reposition internally. At later stages of  
497 compaction, inner and outer cells are also generated by differential orientation of the  
498 cleavage planes by which outer cells divide, with planes oriented parallel or orthogonal to  
499 the cell basal-apical axis giving rise to only outer or both inner and outer cell, respectively.  
500 Finally, actomyosin rings localised at the areas of contact between outer cells generate  
501 tensile forces required to seal the barrier formed from adherens junctions, making it  
502 impermeable. This ultimately allows accumulation and retention of fluid in the blastocoel.

503

#### 504 **Roles of cell positioning and polarity in cell fate determination**

505

506 Experimental removal and spatial rearrangements of blastomeres have shown that early  
507 mouse embryos are highly plastic. When one blastomere is removed from a 2-cell embryo,  
508 the remaining half embryo can develop to term without significant consequences. Also,  
509 blastomeres of 4- to 8-cell embryos, still morphologically and positionally identical, can be  
510 disassembled and reassembled to form chimaeric associations able to support normal pre-  
511 and postimplantation development (Morris *et al.*, 2012). During the 8- to 16-cell interval,  
512 inner and outer cells are formed, developing preferentially into the ICM and TE  
513 compartments of the blastocyst, respectively. This could suggest a change in developmental  
514 rules by which cell potency is progressively restricted. While in general this concept is  
515 correct (Suwińska *et al.*, 2008), the reality is more complex. Elegant cell disassociation-  
516 reassociation experiments have revealed that all blastomeres of mouse 16-cell embryos  
517 maintain the ability to contribute to the formation of both the ICM and the TE (Ziomek and  
518 Johnson, 1982; Suwińska *et al.*, 2008). Blastomeres of 32-cell embryos appear to have lost  
519 this plasticity, developing only into ICM or TE depending on whether they have an inner or  
520 outer origin, respectively (Suwińska *et al.*, 2008). Therefore, under undisturbed conditions  
521 during the 8- 16-cell transition, blastomeres appear to have made 'decisions' about position  
522 and ultimate fate, but these seem not to be irreversible commitments to a specific lineage at

523 this stage. What make blastomeres different during the 8- 16-cell interval is their inner/outer  
524 position and apolar/polar symmetry. Historically, both these conditions have been proposed  
525 to cause cell determination at this stage (reviewed in Mihajlović and Bruce, 2017). According  
526 to one model, by definition, inner cells are totally engaged in cell-to-cell contacts throughout  
527 their surface, while outer cells have also cell-to-cell contact-free surfaces exposed to the  
528 surrounding environment. These two statuses are believed to coincide with different micro-  
529 environmental cues that determine the ICM and TE cell fate, respectively. In the polarity  
530 model, at the 8-cell stage blastomeres become polarised acquiring a basolateral domain  
531 oriented towards the interior and an apical domain facing the exterior. Depending on whether  
532 cleavage occurs parallel or orthogonal to the baso-apical axis, daughter cells lose or  
533 preserve their polarity, respectively, and inherit differently distributed regulatory molecules  
534 that commit them into the ICM or TE pathway. The two models are not mutually exclusive,  
535 though, because predominantly inner cells are apolar and outer cells are polar. These  
536 concepts were already described in this review, but notably several more recent studies  
537 point toward an interplay between polarity, position and a third major regulator of cell fate,  
538 the Hippo signaling pathway. Hippo is a tumour-suppressor pathway described initially in  
539 *Drosophila* and conserved across species (Mihajlović and Bruce, 2017)(reviewed in Harvey  
540 *et al.*, 2013). Its activation depends on several stimuli among which is, importantly in the  
541 case of embryos, cell contact. When Hippo is active, transcriptional co-activator proteins  
542 Yap (Yap1 and Wwtr1) are phosphorylated by Lats protein kinase. In its phosphorylated  
543 form, Yap is retained in the cytoplasmic compartment and degraded. By contrast, inactivity  
544 of Hippo allows nuclear localisation of unphosphorylated Yap. In the nucleus, Yap can bind  
545 several transcription factors, among which are TEA domain transcription factors (TEAD).  
546 Then, the binary complex TEAD-Yap targets several effector genes (Ota and Sasaki, 2008;  
547 Zhao *et al.*, 2008) (Table 1). Clues that the TEAD and Hippo systems are involved in  
548 embryonic cell fate determination derives from experiments showing that in mutated TEAD4  
549 embryos, caudal-type homeoboxprotein-2 (Cdx2), which commit cells into the TE fate, is  
550 largely downregulated and all blastomeres acquire ICM characteristics (Yagi *et al.*, 2007;  
551 Nishioka *et al.*, 2008) (Table 1). However, because TEAD4 is ubiquitously expressed in the  
552 embryo, in normal development a modality must exist that prevents its action in blastomeres  
553 that develop into ICM. The key regulatory factor of this network is indeed Hippo (Figure 2)  
554 (Nishioka *et al.*, 2009). Its signaling pathway is activated only in the inner cells, where Yap  
555 becomes phosphorylated, and so remains in the cytoplasm meaning that TEAD4 cannot  
556 trigger the TE pathway. At the same time, in the outer cells, unphosphorylated Yap can be

557 transported into the nucleus and activate TEAD4 (Hirate *et al.*, 2012). Finally, TEAD4 directly  
558 promotes the transcription of TE-determining genes, such as the previously mentioned  
559 Cdx2, and GATA-binding protein 3 (Gata3) (Nishioka *et al.*, 2009; Ralston *et al.*, 2010)  
560 (Table 1).

561

562 At this stage, cell polarity and position assume importance. In outer cells, polarity is imposed  
563 by mutual regulatory influences by which the Par3-Par6-aPKC (Par-aPCK) system promotes  
564 the organisation of the apical domain (Table 1), while Par1 induces the configuration of the  
565 baso-lateral domain (Suzuki and Ohno, 2006). Par-aPCK appears to have the ability to  
566 inhibit the Hippo pathway, which acts against the specification of TE characteristics. This is  
567 indicated by the finding that at the 32-cell stage in Par-aPCK-mutated embryos showing loss  
568 of polarity in outer cells, Hippo is activated and Yap remains localised in the cytoplasm in all  
569 blastomeres, not only in inner cells (Hirate *et al.*, 2013). On another hand, Hippo signaling  
570 requires strong cell-cell adhesion, which is particularly enhanced in inner cells; this is  
571 confirmed by the fact that Hippo is not activated in dissociated blastomeres from polarity-  
572 disrupted embryos irrespective of the original inner or outer cell position (Hirate *et al.*, 2013).

573

574 In the human, studies on the role of Yap in cell determination are lacking. However, cell  
575 programming experiments showed that Yap is activated when embryonic fibroblasts are  
576 induced to become pluripotent stem cells, suggesting that the hippo regulatory pathway is  
577 involved in the expression of pluripotency (Lian *et al.*, 2010).

578

579 Some details on how, at the molecular level, polarity and cell adhesion interact to regulate  
580 Hippo activation are known (Figure 2). Hippo activation depends on angiominin (Amot)  
581 proteins (Table 1). These regulatory proteins are also crucial for cell fate determination  
582 because their loss is sufficient to commit blastomeres into the TE fate irrespective of their  
583 position or polarity (Hirate *et al.*, 2013; Leung and Zernicka-Goetz, 2013). Amot has a  
584 different intracellular distribution in inner and outer cells. In the former it is found associated  
585 with the adherens junctions throughout the membrane, while in the latter it is selectively  
586 localised at the apical domain (Hirate *et al.*, 2013; Leung and Zernicka-Goetz, 2013). Amot  
587 also has binding activities which are differentially regulated in inner and outer cells. At the  
588 adherens junctions of inner cell, one region of the protein binds the cytoplasmic segment of  
589 E-cadherin and the junction-associated protein Nf2. Another subdomain of Amot has F-actin  
590 binding activity. In the same inner cells when the Amot-E-cadherin-Nf2 complex is formed,

591 phosphorylation of this site by Lats reduces the affinity for F-actin, promotes interaction with  
592 Lats itself and ultimately stimulates the ability of Lats to activate Hippo (Hirate *et al.*, 2013).  
593 In outer cells, polarisation generated by Par-aPKC sequesters Amot in the apical domain  
594 bound to F-actin and prevents the interaction of the same protein with E-cadherin at the  
595 level of adherens junctions. In this fashion, Amot is not phosphorylated by Lats and Hippo  
596 cannot be activated (Hirate *et al.*, 2013). These conditions are permissive for a nuclear  
597 localisation of Yap and expression of TE-determining genes. Taken together, regulation of  
598 the Hippo signaling exemplifies how cell-cell adhesion, whose intensity depends on cell  
599 position and cell polarity, can influence cell fate. This scenario, however, does not rule out  
600 that other factors can affect the Hippo-Yap pathway and ultimately cell fate. For example, in  
601 compacted mouse morulae, when biomechanical forces of outer blastomeres are  
602 experimentally weakened by knocking down the maternal myosin gene (*Myh9*), loss of cell  
603 contractility alone can lead to increased Yap phosphorylation and cytoplasmic localization.  
604 As a consequence of reduced intranuclear YAP presence, although such blastomeres  
605 remain external, they fail to express the TEAD-Yap-controlled genes that are responsible  
606 for the acquisition of TE characteristics (Maître *et al.*, 2016). These experiments  
607 demonstrate an extreme integration of mechanical, positional and molecular cues in a single  
608 regulative network of cell fate determination.

609

## 610 **Molecular control of cell fate determination**

611

612 As described, Hippo signaling is a major regulatory mechanism which ultimately prevents  
613 expression of transcription factors that elicit the TE phenotype. However, several other  
614 questions concern the pathways that promote the specification of the TE and ICM lineages  
615 and establish mutual regulatory loops. Notably, decisions on cell fate are not necessarily  
616 made at the same time by the cells of the same compartment, making the understanding of  
617 regulatory networks particularly arduous. In addition, the large majority of data have been  
618 generated in the mouse model (see below), while information concerning the human  
619 embryos remains scant.

620

### 621 TE specification

622

623 As discussed above, *Cdx2* is a crucial component for specification of TE characteristics. In  
624 the mouse 8-cell stage embryo, this regulator is expressed only in some blastomeres but as

625 the morula forms, it is found in a higher proportion of cells, to become expressed in outer  
 626 cells of the morula and finally in elements of the TE (Jedrusik *et al.*, 2008; Ralston and  
 627 Rossant, 2008). *Cdx2* <sup>-/-</sup> is embryonic lethal, due to the inability to form a TE. Embryos  
 628 begin to cavitate but the blastocoel do not form properly because outer cells do not acquire  
 629 their typical epithelial phenotype and fail to form a selective barrier which is instrumental for  
 630 inward transport of water and expansion of the embryo (Strumpf *et al.*, 2005). In the absence  
 631 of *Cdx2* expression, a lack of epithelial characteristics is associated with specific molecular  
 632 features, such as repression of *Eomesodermin* (*Eomes*) and *Hand1*, while ICM-determining  
 633 genes are expressed ectopically (Strumpf *et al.*, 2005). *Eomes* is a T-box-specific factor  
 634 (Ciruna and Rossant, 1999). In null mutants, its absence coincides with the inability of  
 635 embryos to develop a functional TE. This occurs in association with normal levels of *Cdx2*  
 636 while *Hand1* and *P11*, typical of epithelial differentiation, are undetectable (Strumpf *et al.*,  
 637 2005). Combined, these data suggest that the relative position of *Eomes* in the pathway that  
 638 specifies the TE fate is downstream of *Cdx2* and upstream of more terminal markers of TE  
 639 differentiation. Another key regulator of TE formation is the zinc-finger transcription factor  
 640 *Gata3*. Loss of function of this factor, through RNAi action, leads to a phenotype very similar  
 641 to *Cdx2* null mutants, in which blastocyst formation is impeded and development arrests at  
 642 the blastocyst stage (Ralston *et al.*, 2010). While there is consensus on the opinion that  
 643 *Gata3* is not governed by the Hippo pathway, the question of the hierarchical position with  
 644 respect to *Cdx2* is more controversial. However, a comparable temporal pattern of  
 645 expression between *Gata3* and *Cdx2* suggest that the two regulators act in parallel (Ralston  
 646 and Rossant, 2008), with no upstream influence of one over the other. Indeed, *Gata3* is co-  
 647 expressed with *Cdx2* already at the 8-cell stage and is found in TE, but not ICM cells, at the  
 648 blastocyst stage (Ralston *et al.*, 2010).

649

#### 650 ICM specification

651

652 As the inner cells commit to their lineage, they must not only not express TE characteristics  
 653 as an effect of hippo activation, but also, they must initiate the ICM pathway. In such cells,  
 654 the master regulatory factor is the octamer-binding transcription factor 3/4 (*Oct4*, also known  
 655 as *Pou5f1*) (Table 1). *Oct4* positively regulates gene transcription by recognising the  
 656 sequence ATGCAAAT situated in the promoters and enhancer regions of several genes  
 657 (Schöler *et al.*, 1990). While its expression is observed throughout preimplantation  
 658 development, as the embryo develops its presence becomes restricted to the ICM. *Oct4*

659 appears downregulated in embryonic stem cells following differentiation induced by  
660 leukaemia inhibitory factor (LIF) (Palmieri *et al.*, 1994), while its absence in Oct4 *-/-* mutant  
661 embryos does not prevent decidualisation but causes peri-implantation death (Nichols *et al.*,  
662 1998). This factor is therefore considered instrumental for pluripotency. Other factors  
663 contribute to pluripotency of the ICM and are differentially regulated in a cell- and stage-  
664 specific fashion. For example, in the mouse, Nanog (Chambers *et al.*, 2003) is expressed in  
665 the inner cells of the morula and becomes restricted to the epiblast of the ICM in the  
666 blastocyst, from which the embryo proper derives (Chambers *et al.*, 2003). Indeed,  
667 reprogramming experiments of somatic cells into induced pluripotent stem cells illustrate the  
668 critical importance of both Oct4 and Nanog in maintaining pluripotency (Takahashi and  
669 Yamanaka, 2006; Okita *et al.*, 2007).

670

671 The binding domain-containing family of transcription factors, Sox2, is a member of the High  
672 Mobility Group (HMG) protein family that is associated with maintenance of pluripotency in  
673 the embryo (Table 1). It is detectable throughout mouse preimplantation development, from  
674 the zygote until the morula stage, when it disappears in the cells that will form the TE (Avilion  
675 *et al.*, 2003). The observations that Sox2 *-/-* embryos have an unchanged pattern of protein  
676 expression and that Sox2 mRNA starts to be produced from the morula stage indicate that  
677 this factor is accumulated in the oocyte during oogenesis (Avilion *et al.*, 2003). Sox2 appears  
678 crucial for ICM and in particular the epiblast at the time of implantation (Avilion *et al.*, 2003).  
679 Furthermore, an interaction between Sox2 and Oct4 appears essential for the expression  
680 of downstream pluripotency genes, such as Fgf4 (Ambrosetti *et al.*, 1997), Utf1 (Okuda *et al.*  
681 *et al.*, 1998) and Nanog (Rodda *et al.*, 2005) (Table 1). Again, in the human, the timing of  
682 expression may differ; indeed, nuclear localization of Sox2 was observed only from the  
683 compacted morula stage, i.e. much later than in the mouse embryo (Cauffman *et al.*, 2009).

684

### 685 Mutual negative regulation

686

687 Several lines of evidence converge towards a model involving mutual interaction at the  
688 intracellular level between the ICM and TE regulatory systems. Indeed, Cdx2 knockout  
689 mouse embryos are unable to promote positive regulation of TE specific genes such as  
690 Eomes and Hand1 in outer cells, but instead express Oct4 and Nanog, which are typically  
691 associated with inner cells. At the molecular level, this cross-regulation is explained by the  
692 fact that at the blastocyst stage Cdx2 and SWI/SNF chromatin remodelling factor co-operate

693 to bind the promoter region of Oct4 in cells that will develop into TE, thus inhibiting the action  
694 of this pluripotency factor. This replicates a scheme observed in other types of epithelial  
695 cells (Wang *et al.*, 2010).

696

697 There are also hints of an inverse, mutual regulation between the Cdx2 and Oct4 pathways.  
698 Indeed, in embryonic stem cells, Oct4 was found to interact with Cdk1. The two proteins  
699 form a complex that is required to repress the transcription of Cdx2 (Li *et al.*, 2012).  
700 Therefore, in the mouse, not only do the Cdx2 and Oct4 regulatory pathways specify the TE  
701 and ICM cell fates, respectively, but also are involved in an elegant interplay of mutual  
702 regulation in which one represses the other in its own embryonic compartment. Consistent  
703 with the mouse embryo, in the human at the 8-cell stage, Oct4 and Nanog are initially  
704 expressed in all blastomeres; afterwards their localization remains limited to the ICM  
705 (Kimber *et al.*, 2008; Niakan *et al.*, 2012). By contrast, human embryos show a different  
706 temporal pattern of Cdx2 expression. In fact, this regulator is detectable only after formation  
707 of the blastocoel and is initially co-expressed with Oct4 (Niakan *et al.*, 2012). It seems,  
708 therefore, that specification of the two cell lines occurs significantly later than in the mouse  
709 and does not necessarily require the Oct4/Cdx2 antagonism. This opens the possibility that  
710 the fate of human blastomeres can remain plastic until relatively advanced stages of  
711 development. Indeed, human outer cells from compacted morulae or early blastocysts can  
712 contribute to formation of the ICM if experimentally repositioned inside the embryo (De  
713 Paepe *et al.*, 2013) . Furthermore, after isolation and reaggregation, human outer cells are  
714 able to reconstitute an embryo able to cavitate and form an ICM (De Paepe *et al.*, 2013).  
715 Therefore, while mouse and human development share some common features of cell fate  
716 determination, further studies are warranted to ascertain possible interspecies differences.

717

## 718 **The morula stage in clinical embryology**

719

### 720 Embryo selection and transfer at the morula stage

721

722 Clinical data on the reproductive performance of day 4 embryos are scarce. In clinical IVF,  
723 embryos are usually transferred in the uterus at early cleavage (days 2/3) or blastocyst (days  
724 5/6) stages. Much less commonly, they are replaced on day 4 when they are expected to  
725 have reached the morula stage, owing in part to the challenges described above.  
726 Furthermore, where such embryos are used, they will often be transferred together with day

727 3 or day 5 embryos, making the association between embryonic stage and ability to implant  
728 and develop to term unachievable. However, there is isolated evidence that embryo transfer  
729 (ET) can be successfully implemented on day 4. In the late nineties, preliminary reports  
730 indicated that morula stage embryos could be used for ET following blastomere biopsy on  
731 day 3 in preimplantation genetic testing (PGT) cycles (Grifo *et al.*, 1998; Gianaroli *et al.*,  
732 1999). In one of the few specific studies carried out on the morula stage, Tao *et al.* (2002)  
733 compared retrospectively the relative clinical performance of day 3 and day 4 embryos. In  
734 all cases, day 4 ET were associated with higher implantation and pregnancy rates,  
735 suggesting that culture to day 4 could offer better chances for embryo selection compared  
736 to day 3. A prospective randomised study published several years later and including 350  
737 couples (Pantos *et al.*, 2008) showed that day 4 ET was associated with implantation and  
738 clinical pregnancy rates that were comparable to day 3 ET (22.0 vs. 21.0 and 49.7% vs.  
739 45.3%, respectively). A further retrospective analysis (Skorupski *et al.*, 2007) claimed that  
740 the transfer of day 4 embryos was compatible with high rates of implantation (45% to 34%)  
741 and live birth (55% to 33%) across a large spectrum of female age ( $\leq 34$  to 40 years).  
742 However, the absence of a control group and scoring criteria, as well as the fact that some  
743 of the embryos had undergone assisted hatching, means that these data need careful  
744 interpretation. The clinical outcome of day 4 and day 5 ET was the object of another  
745 retrospective study (Feil *et al.*, 2008). This analysis adopted a single embryo transfer (SET)  
746 approach and reported comparable overall ongoing pregnancy rates between day 4 and day  
747 5 transfer (38.7% and 32.1%, respectively). In addition, sub-analysis of the day 4 group  
748 revealed that morulae entirely or partially compacted, and otherwise normal, implanted with  
749 comparable rates (40.0% and 37.1%, respectively); however, in cases where partial  
750 compaction was associated with large vacuoles and extensive fragmentation, the  
751 implantation rate dropped dramatically. Thus, day 4 embryos at the morula stage can be  
752 discriminated morphologically and developmentally to some degree and, in the best-case  
753 scenario, appear to have implantation rates comparable to that achieved by day 5 transfers.  
754 Such conclusions are supported by a further retrospective study by Kang *et al.*, (2012). For  
755 all major clinical outcomes, i.e. pregnancy rate (51.5% vs. 51.8%), implantation rate (52.3%  
756 vs. 52.5%) and live birth rate (39.2% vs. 44.7%), results were comparable between morula  
757 and day 5 ET, respectively, although the miscarriage rate tended to be higher in the day 4  
758 group. Together, these data suggest that ET of morulae on day 4 can be contemplated as  
759 an option in case of intense laboratory workload and necessity to redistribute ETs more  
760 uniformly over consecutive days.



761

762 A recent report specifically focused on the predictive value of morphology of day 4 embryos  
763 (Fabozzi *et al.*, 2015). Embryos were considered at the morula stage if showing at least 14  
764 cells and scored according to four morphological classes (A-D, best-worst), depending on a  
765 combination of the degrees of compaction and integrity. Out of 393 embryos, the proportions  
766 of grade A-D morulae were 47.84%, 26.72%, 20.36% and 5.09%, respectively. The degree  
767 of compaction and integrity was positively associated with the blastocyst formation rate  
768 which was 87.2%, 63.8%, 41.3%, and 15.0% for the A-D classes, respectively. Importantly,  
769 this classification system was found to be a better predictor of blastocyst formation and  
770 quality compared with a conventional grading system performed on day 3. Overall, these  
771 data confirm previous reports suggesting that the scoring systems based on degree of  
772 compaction and integrity of day 4 embryos have the ability to predict the rate and quality of  
773 blastocyst formation (Ebner *et al.*, 2009; Ivec *et al.*, 2011).

774

#### 775 The use of morula stage embryos for preimplantation genetic testing

776

777 Over the course of the last decade, PGT has undergone major changes. Progress in embryo  
778 culture and cryopreservation have shifted the preference for the time of embryo biopsy from  
779 day 3 (6-8 cell stage) to day 5 or 6 (blastocyst stage). This means that more cells are  
780 available for analysis (3-10 instead of 1-2) and, in general, embryos appear more resilient  
781 to the biopsy procedure. However, Zakharova *et al.* (2014) reported on embryo biopsies  
782 performed on 709 day 4 morulae from 215 PGT cycles. To achieve loosening of intercellular  
783 contacts, compacted morulae were exposed to a Ca<sup>2+</sup>-free medium. Three to seven  
784 blastomeres were biopsied from each embryo and analysed by fluorescent in-situ  
785 hybridization (FISH). After return to standard medium containing Ca<sup>2+</sup>, more than 90% of  
786 biopsied embryos reached the blastocyst stage by day 6. In an impressive range of  
787 endpoints, including postnatal follow up, no differences were apparent between the PGT  
788 and non-PGT groups. Therefore, the authors concluded that embryo biopsy is technically  
789 feasible and safe to be carried out at the compacted morula stage. Although encouraging,  
790 caution must be exercised, since the relatively small numbers of embryos from a single-  
791 centre study are unlikely to offer solid conclusions on the safety of the day 4 approach.  
792 Concerns derive not only from the possible impact of removal of 3-7 cells from an embryo  
793 usually formed from 12-32 blastomeres, but also from potential undefined physiological  
794 effects arising from the disruption, as we have illustrated above, of highly complex and

795 important morphogenetic events occurring at the morula stage. Thus, exposure to Ca<sup>2+</sup>-free  
796 medium, functional ablation of intercellular contacts and consequent reversion of cell shape  
797 from elongated to spherical, although transient, could potentially perturb embryo physiology  
798 with long-term developmental consequences. Nonetheless, embryo biopsy at the morula  
799 stage remains an interesting option, implicating advantages such as i) the recovery of a  
800 higher number of cells compared to day 3 biopsy, ii) the possibility to have one full day for  
801 chromosome analysis and therefore perform fresh ET of unaffected embryos on day 5, and  
802 iii) the recovery of intact biopsied cells, whose integrity is amenable to FISH, in cases where  
803 this analytical method is preferable.

804

#### 805 Cryopreservation of human morulae

806

807 Cryopreservation at the morula stage has been reported in numerous studies conducted in  
808 animal models, including mouse, rat, cow, pig and goat. Therefore, it appears to be  
809 technically possible. However, evidence from animal studies cannot be directly applied to  
810 human IVF. Membrane permeability to the various cryoprotectants, on which successful  
811 cryopreservation depends, occurs by simple diffusion or via specific channels (aquaporins)  
812 according to kinetics that vary from stage to stage often in a species-specific fashion. Early  
813 experiments suggested that human blastocysts vitrified using fine plastic capillaries as  
814 storage devices gave acceptable rates of survival (Cremades, 2004). Recent progress in  
815 vitrification has allowed improvement in survival rates of both intact and biopsied morulae  
816 (92.0% and 87.5%, respectively) (Zhang *et al.*, 2009). Isolated cases of healthy live births  
817 (one twin and one single) were also reported after the transfer of compacted morulae  
818 cryopreserved by slow freezing (Tao *et al.*, 2001). In a larger case series, 54 day 4 morulae  
819 were warmed after vitrification; 38 embryos survived to 24 hours after thawing, while 30  
820 developed to the blastocyst stage. In 18 ETs, implantation and birth rates for ET were 20.0%  
821 (6/30) and 27.8% (5/18), respectively (Vanderzwalmen *et al.*, 2002). Therefore, while  
822 cryopreservation at the morula stage can be applied, the limited available data suggest a  
823 reduced efficacy compared with cryopreservation at the cleavage or blastocyst stage. Again  
824 however, caution should be exercised as this is an underinvestigated area of treatment.

825

#### 826 Emerging data from TLM

827

#### 828 *Timing of the morula stage*

829

830 For decades, the morula has been described as characteristic of day 4, but little data was  
831 available on the specific timing of cleavage events beyond the 8-cell stage or the dynamics  
832 of the process of compaction (ESHRE Atlas of Human Embryology). Isolated reports  
833 indicated that starting compaction 'early' was associated with a higher implantation potential  
834 (Skiadas *et al.*, 2006), although the precise timing of initiation of this process was not  
835 described.

836

837 The introduction of TLM has provided new opportunities to observe the morphokinetics of  
838 the morula stage. In one of the first TLM studies describing the morula stage, Campbell *et al.*  
839 *et al.*, (2013) reported the median time of start of compaction in euploid embryos was 79.7  
840 hours post ICSI, a value similar to that of embryos affected by a single aneuploidy (80.7  
841 hours), but significantly different to that of embryos carrying multiple aneuploidies (85.1  
842 hours). Median times of full morula formation in the three study groups were not statistically  
843 different (83.5, 87.9 and 88.2 hours, respectively). Another study investigated the formation  
844 of the fully compacted morula in embryos of PGT cases, reporting mean times of 94.4 and  
845 95.3 hours in euploid and aneuploid embryos, respectively (Minasi *et al.*, 2016). Times of  
846 full compaction appears different in the two studies, although it should be noted that they  
847 were reported using different parameters (median and mean, respectively). In a study  
848 specifically focussed on stage of compaction, Iwata *et al.* (2014) found that compaction can  
849 occur at any stage between the 4- and the 16-cell stage, although the majority of embryos  
850 initiate compaction at the 8-cell stage or later. In such embryos, almost half go on to form  
851 good quality blastocysts, while embryos starting compaction before the 8-cell stage form  
852 good blastocysts in less than 20% of cases. Interestingly, early compacting embryos appear  
853 to be characterised by cell cycle anomalies, like multinucleation caused by cytokinesis  
854 failure (Iwata *et al.*, 2014).

855

856 Technically, the determination of the time of full compaction is challenging because of its  
857 dynamic nature, occurring over several hours. Therefore, significant intra- and inter-operator  
858 variability cannot be excluded. Nonetheless, it is realistic that time differences between  
859 studies, at the morula and all developmental stages, can reflect intrinsic differences between  
860 cohorts of embryos and/or extrinsic influences, such as culture conditions. Indeed, while  
861 mean times of formation of fully compacted morulae are very similar in males and females  
862 embryos (91.2 and 91.9 hours, respectively) (Bronet *et al.*, 2015), embryos from

863 overweight/obese women have an overall faster developmental kinetics. This is also  
864 reflected in times of morula formation 17 hours shorter than women of normal weight (Leary  
865 *et al.*, 2015). Likewise, in another study, it was shown that embryos undergoing repeated  
866 contractions and expansions during blastocyst formation, a phenomenon indicated as a  
867 negative marker of implantation potential, achieve full compaction approximately 5 hours  
868 earlier (80.9 vs. 86.3 hours, respectively) compared with those that do not exhibit cyclical  
869 contractions/expansions (Marcos *et al.*, 2015). Laboratory manipulations can also impact  
870 the morula stage. Indeed, an analysis carried out in embryos produced in PGT cycles  
871 described that day 3 embryos of at least 8 cells and subject to removal of 1-2 blastomeres  
872 progressed to the morula stage with a 5-hour delay and implanted at a lower rate compared  
873 with embryos of a control group (Bar-El *et al.*, 2016). These findings are consistent with a  
874 previous study reported by Kirkegaard *et al.* (2011). Notwithstanding such limitations, the  
875 time of compaction alone appears to be associated with embryo implantation. This notion  
876 derives from a recent study showing that in a selected population of embryos full compaction  
877 in accomplished 14 hours earlier in those that implant (86.4 vs. 100.3 hours, respectively)  
878 (Motato *et al.*, 2016).

879

#### 880 *Morphokinetics and chromosomal status of the morula*

881 Data from a recent investigation (Lagalla *et al.*, 2017) suggest that the morula stage could  
882 be an important checkpoint for embryo quality during preimplantation development and that  
883 the process of compaction could be involved in mechanisms of self-correction. The study  
884 confirmed that diverse cleavage anomalies (e.g. direct cleavage of one cell into three, rapid  
885 cleavage, reverse cleavage) were associated with a 50% reduction in the chances of an  
886 embryo developing into a good quality blastocyst. Interestingly, the study also showed that  
887 blastocysts derived from such abnormally cleaving embryos exhibited a higher rate of  
888 euploidy compared with blastocysts developed from embryos without cleavage anomalies  
889 (75.0% vs, 49.2% respectively). Even more interestingly, it was found that all euploid  
890 blastocysts derived from abnormally cleaving embryos were characterised by the  
891 phenomenon of partial compaction. Partial compaction was observed also in embryos with  
892 no apparent cleavage anomalies, but at a much lower rate. Taken together, these findings  
893 suggest that, on one hand, blastomeres that have become aneuploid as a consequence of  
894 cleavage anomalies (e.g. direct cleavage) may in some cases prevent development to  
895 blastocyst. On the other hand, it might be possible that such aneuploid blastomeres are  
896 excluded at compaction, with a rescue of the original condition of euploidy of the embryo as

897 a whole. The latter hypothesis appears to be confirmed by comparative chromosome  
898 analysis of cells excluded from compaction. In the majority of cases, in the same embryo  
899 the chromosome constitution of these cells was more affected by segregation errors  
900 compared with cells of the trophectoderm, or they could not be analysed due to extensive  
901 DNA degradation, a hallmark of apoptosis often triggered in response to complex  
902 aneuploidy. Therefore, it is plausible to propose that during compaction, aneuploid cells are  
903 excluded from the embryos allowing a restoration of euploidy. This is in line with the well-  
904 established notion that mosaicism is more frequent in earlier stages of development than at  
905 the blastocyst stage. If true, this suggests a startling critical role for the morula in terms of  
906 embryo rescue and elimination of mosaicism, as well as the more widely-described functions  
907 relating to morphogenetic events and determination of cell fate. The notion of the existence  
908 of mechanisms of self-correction during preimplantation development is also emerging from  
909 other observations. For example, a study on embryos displaying direct unequal cleavage  
910 (i.e. division of one cell into three or more daughter cells) indicates that extrusion from the  
911 compacted morula of blastomeres affected by such a cleavage anomaly is strongly  
912 associated with the formation of good quality blastocysts (Zhan *et al.*, 2016). Such  
913 information might be of crucial relevance in the application of PGT at the morula stage.

914

915 *A plea for more research on the morula stage and compaction process*

916

917 Recent investigations confirm the necessity of intensifying research efforts on the morula  
918 stage. Data from Mayer et al (2018) described the phenomenon of late onset of cell  
919 vacuolisation at the morula stage. In particular, they reported that the implantation ability of  
920 good quality blastocysts is significantly reduced in case of formation of vacuoles at the  
921 morula stage. The morula stage also appears relevant to the management of slow-growing  
922 embryos. In fact, Tannus et al. (2018) observed that embryos reaching the morula stage on  
923 early day 5 and achieving the blastocyst stage on day 6 can implant with higher frequency  
924 if cryopreserved and transferred in a frozen embryo replacement, instead of being used  
925 fresh, confirming the importance of endometrial receptivity and synchronisation. Even more  
926 intriguing information seems to derive from detailed observation of the mechanism of  
927 compaction and elimination of presumably abnormal cell from the mass that then organises  
928 itself into blastocyst. In a preliminary report, Lagalla and colleagues (2018) proposed the  
929 existence of two distinct mechanisms of cell elimination resulting in partial compaction: a)  
930 exclusion of cells from the outset, before the beginning of compaction, or b) extrusion of

931 cells following their initial involvement in the compaction process. Interestingly, in partially  
932 compacted embryos, exclusion was found to occur more frequently in embryos of younger  
933 women, while the opposite was observed in case of extrusion. This finding, together with  
934 the recently reported differences in chromosome constitution between the embryo and cells  
935 eliminated during compaction (Lagalla et al., 2017), is suggestive of the morula stage as an  
936 important checkpoint stage during formation of a viable blastocyst. However, research in  
937 clinical embryology alone is unlikely to have the capacity to reveal the intimate secrets of  
938 the morula stage. Strategic collaboration with other disciplines and approaches will be  
939 crucial to ensure progress in this field. Being a dynamic process and extremely difficult to  
940 observe even with the aid of TLM, the morula stage could be amenable to investigation by  
941 exploiting machine learning approaches, which has already found application to predict  
942 embryo implantation ability at the blastocyst stage (Zaninovic et al., 2018). Further progress  
943 could derive from the analysis of extruded/excluded cells to reveal their cellular and  
944 molecular constitution and gain more in-depth insights on possible mechanisms of self-  
945 correction. Live cell imaging, especially if developed in a truly non-invasive fashion, could  
946 also give a crucial impulse to research on the morula stage. Three-dimensional modelling  
947 approaches have already begun to reveal how cell internalisation, and therefore formation  
948 of the two main cell lineages, occurs only when differences in cell surface contractility  
949 exceed a predictable threshold (Maître et al., 2016). Molecular, cellular and metabolic  
950 analysis of spent media exposed to embryos of different developmental stages could shed  
951 new light on the morula and better explain its unique characteristics.

952

## 953 **Conclusions**

954

955 The morula stage and the associated process of compaction have received relatively little  
956 attention in ART due to their highly dynamic nature combined with the ambiguous criteria of  
957 morphological classification. Morula stage embryos therefore remain scarcely integrated in  
958 the mainstream ART procedures, such as embryo quality assessment, ET, cryopreservation  
959 and embryo biopsy. This is in striking conflict with the importance that the construction of  
960 the morula has for preimplantation development and indeed the entire developmental  
961 process. For example, during the temporal span of the morula stage, newly expressed  
962 genes of the embryonic genome mediate the transition from a low-activity, low-energy  
963 metabolic state characterising the early cleavage stages to a more active glucose-based  
964 aerobic capacity required for the formation and expansion of the blastocyst (Leese, 2012).

965 From a developmental standpoint, the morula stage is when the first crucial cell-fate decision  
966 is taken, whereby blastomeres commit themselves to contribute to either the ICM or TE  
967 compartment (Mihajlović and Bruce, 2017). As early as the 1980s, positional information  
968 (internal or external localisation) was recognised as a determining factor for the  
969 establishment of an ICM or TE cell fate (Fleming, 1987). Early studies also suggested that,  
970 starting from the 8-10 cell stage, differential intracellular distribution of cell determinants  
971 cooperates with the geometry of cleavage planes to generate functional diversity between  
972 internal and external cells (Johnson and Ziomek, 1981a; 1981b). These notions remain valid  
973 today. However, recent studies, aided by extraordinary progress in live cell imaging  
974 techniques, have expanded our understanding of the biology of the morula (Table 1). We  
975 now know that compaction is assisted not only by junctional structures, but also by fine and  
976 dynamic regulation of specialised cellular projections, filopodia (Fierro-González *et al.*,  
977 2013). We have also learnt that generation of inner and outer cells occurs not only as a  
978 consequence of alternative orientation of cleavage planes, but also as an effect of  
979 differences in cortical tension between adjacent outer blastomeres. In addition, new  
980 information has revealed the crucial role of an actin-zippering mechanism in ensuring  
981 intercellular sealing and establishing a permeability barrier required by the process of  
982 blastocoel expansion. New light has also been shed on the mutually regulated pathways of  
983 gene expression that commit cells into distinct developmental destinies (Jedrusik, 2015).  
984 Although obtained in the mouse model, such a wealth of information has reignited interest  
985 in the specific merit of the morula stage for clinical embryology. Information on possible  
986 influences of in-vitro culture on developmental decisions taken at the morula stage are  
987 lacking in human ART. Indeed, recent morphokinetic data suggest that compaction could  
988 represent an opportunity to sense chromosomal anomalies in individual blastomeres and  
989 implement self-correction mechanisms aimed at excluding abnormal cells from the  
990 developing embryo (Lagalla *et al.*, 2017). Overall, such a knowledge casts new light on this  
991 crucial developmental stage. It also indicates new avenues for future research in  
992 preimplantation development in the context of clinical embryology.

993

**994 Authors' roles**

995 Giovanni Coticchio: MS design, literature search, MS writing critical discussion and editing

996 Cristina Lagalla: MS design; literature search; MS writing, critical discussion and editing

997 Roger Sturmey: MS writing

998 Francesca Pennetta: MS design, critical reading and editing; Figures design and  
999 production

1000 Andrea Borini: MS design and critical discussion

1001

1002

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1007 None declared

1008

1009

1010



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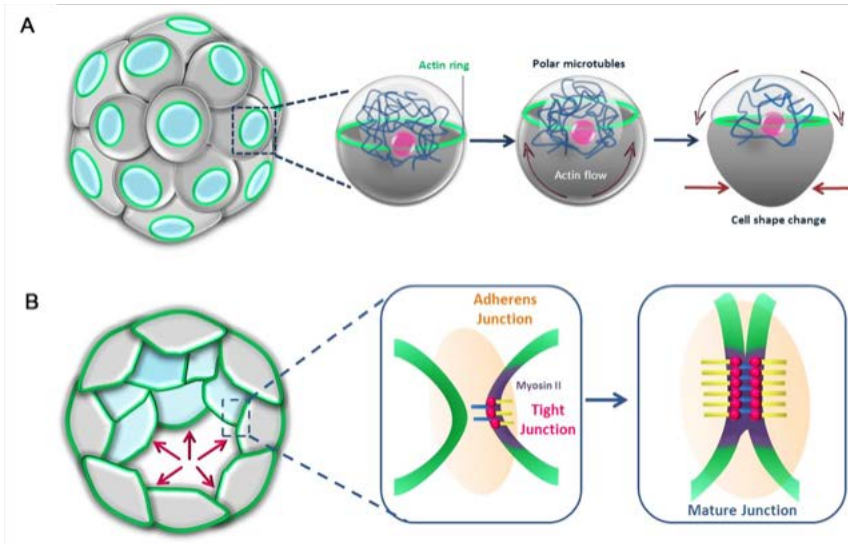
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1343 **Figures and table legends**

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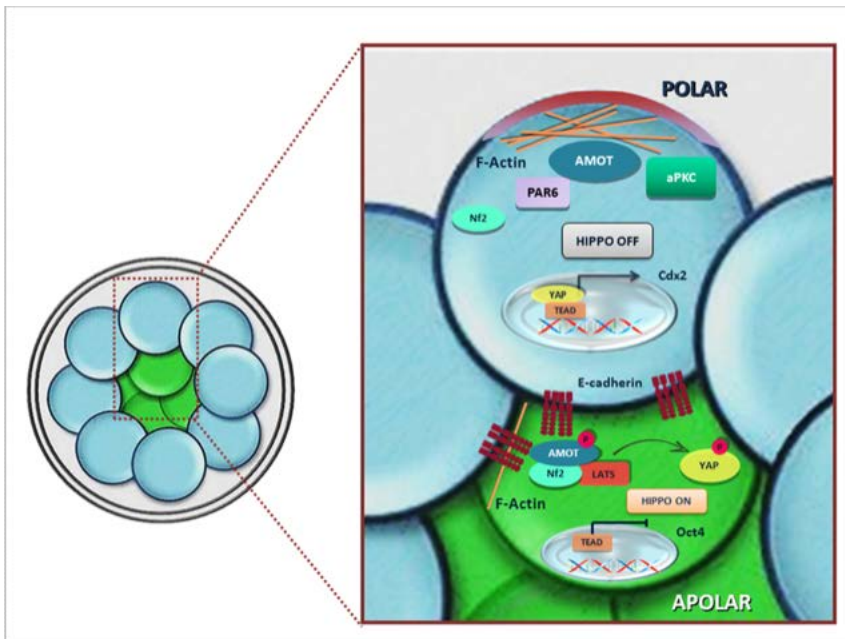
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**Figure 1. Forces that shape blastomeres and secure sealing of intercellular spaces at the morula stage.** A) Flows of cortical cytoplasm and a network of polar microtubules cooperate to generate a ring of actin in the apical domain of outer cells. This ring expands reaching the cell boundaries, recruits myosin II, becomes contractile and allows cell shape remodelling. B) At cell boundaries, expanded actin rings cooperate with tight and adherens junctions to achieve intercellular zippering and sealing to finally assist accumulation of intercellular fluid and formation of the blastocoel.



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1360 **Figure 2. Example of integration of positional, polarity and molecular cues to achieve**

1361 **differential gene expression and determination of alternative cell fates.** In inner cells

1362 (in green), Amot is associated with the adherens junctions throughout the cell membrane

1363 and has reduced activity for F-actin. This condition promotes interaction with and

1364 phosphorylation by Lats and ultimately activation of the Hippo regulatory pathway, which

1365 prevents intranuclear localisation of Yap and expression of TE determining genes. In outer

1366 cells (in blue), polarisation generated by Par-aPKC sequesters Amot in the apical domain

1367 bound to F-actin and prevents the interaction of the same protein with E-cadherin at the

1368 level of adherens junctions. In this fashion, Amot is not phosphorylated by Lats and Hippo

1369 cannot be activated (Hirate et al., 2013).

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Protein	Developmental role	References
E-cadherin Catenins	Contribute to the formation of adherens junctions (zonulae adherens), which ensure mutual lateral adhesion between epithelial cells, thus contributing to the maintenance of epithelial cell polarity	Vestweber et al., 1987; Haegel et al., 1995; Riethmacher et al., 1995; Campbell et al., 1997; Torres et al., 1997; Perez-Moreno et al., 2003; Kang et al., 2005.
E-cadherin F-actin Myo10	Contribute to the organization and function of filopodia, which extend between adjacent blastomeres and determine cell flattening during compaction	Fierro-González et al., 2013.
E-cadherin	Intervenes in the formation of tight junctions (zonulae occludens), which create belt-like impermeable structures located at the upper-lateral sides of adjacent outer cells	Eckert and Fleming, 2008.
Actin Myosin II	Co-operate to lead to the formation of an actomyosin ring localized at the margins of adjacent cells. This ring produces the tension forces required for the zippering mechanism that acts in coordination with cell junctions to seal the intercellular contact between outer cells	Zenker et al., 2018.
Myosin II	Generates tensile forces at cell borders and constriction of the apical domain, by which blastomeres are internalized	Samarage et al, 2015.
Hippo Yap Lats proteins	Co-operate in the formation of a regulatory mechanism involving signaling of Hippo for the phosphorylation and consequent cytoplasmic localization of Yap. Absence of nuclear localization of Yap prevents the activation of the transcription regulator TEAD. Inactivation of Hippo leads to localization of dephosphorylated Yap in the nucleus and activation of TEAD	Ota and Sakaki, 2008; Zhao et al., 2018; reviewed in Harvey et al., 2013; Nishioka et al., 2009; Hirate et al., 2012.
TEAD	Upon activation by unphosphorylated Yap, TEAD initiates the transcription of several effector genes, among which those that determine the trophectoderm cell fate (e.g. Cdx2 and Gata3)	Yagi et al. 2007; Nishioka et al., 2008; Nishioka et al., 2009; Ralston et al., 2010.
Par3-Par6-aPKC (Par-aPCK) Par1	Interact mutually to establish apical (Par-aPCK) and baso-lateral domains in outer cells	Suzuki and Ohno, 2006.

Par3-Par6-aPKC (Par-aPCK)	Inhibit the Hippo pathway, which acts against the specification of trophectoderm characteristics	Hirate et al., 2013.
Angiomotin (Amot) proteins Lats	Cooperate to activate Hippo	Hirate et al., 2013; Leung and Zernicka-Goetz;
Cdx2 Gata3	Once expressed as an effect of positive regulation by TEAD, induce the expression of trophectoderm characteristics.	Jedrusik 2008; Ralston and Rossant, 2008; Ralston et al, 2010
Oct 4 Sox2	Act as apical regulators for the maintenance of cell pluripotency	Chambers et al. 2003; Takahashi and Yamanata, 2006; Okita et al., 2007
Fgf4 Nanog Utf1	Are regulated by Oct4 and Sox2, contributing to the preservation of pluripotency	Ambrosetti et al., 1997; Okuda et al., 1998; Cauffman et al., 2009.

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**Table 1**

1387 **Table 1. Schematic description of function of proteins in key developmental**  
1388 **processes that characterise the morula stage.** These include compaction, intercellular  
1389 sealing to allow blastocoel formation, positioning of inner and outer cells, establishment of  
1390 cell polarity and cell fate determination.

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