# Kent Academic Repository Full text document (pdf)

### **Citation for published version**

Victor, Andrea R. and Griffin, Darren K. and Brake, Alan J and Tyndall, Jack C and Murphy, Alex E and Lepkowsky, Laura T and Lal, Archana and Zouves, Christo G and Barnes, Frank L and McCoy, Rajiv C and Viotti, Manuel (2019) Assessment of aneuploidy concordance between clinical trophectoderm biopsy and blastocyst. Human Reproduction, 34 (1). pp. 181-192. ISSN

## DOI

https://doi.org/10.1093/humrep/dey327

### Link to record in KAR

https://kar.kent.ac.uk/72159/

### **Document Version**

Author's Accepted Manuscript

#### Copyright & reuse

Content in the Kent Academic Repository is made available for research purposes. Unless otherwise stated all content is protected by copyright and in the absence of an open licence (eg Creative Commons), permissions for further reuse of content should be sought from the publisher, author or other copyright holder.

#### Versions of research

The version in the Kent Academic Repository may differ from the final published version. Users are advised to check http://kar.kent.ac.uk for the status of the paper. Users should always cite the published version of record.

#### Enquiries

For any further enquiries regarding the licence status of this document, please contact: **researchsupport@kent.ac.uk** 

If you believe this document infringes copyright then please contact the KAR admin team with the take-down information provided at http://kar.kent.ac.uk/contact.html





1	Assessment of aneuploidy concordance between clinical trophectoderm
2	biopsy and blastocyst
3	
4	
5	Running Title:
6	Aneuploidy concordance within blastocysts
7	
8	Authors:
9	Andrea R. Victor <sup>1,2</sup> , Darren K. Griffin <sup>2</sup> , Alan J. Brake <sup>1</sup> , Jack C. Tyndall <sup>1</sup> , Alex E.
10	Murphy <sup>1</sup> , Laura T. Lepkowsky <sup>1</sup> , Archana Lal <sup>1</sup> , Christo G. Zouves <sup>1,3</sup> , Frank L.
11	Barnes <sup>1,3</sup> , Rajiv C. McCoy <sup>4</sup> and Manuel Viotti <sup>1,3,*</sup>
12	
13	<sup>1</sup> Zouves Fertility Center, Foster City, California, USA;
14	<sup>2</sup> School of Biosciences, University of Kent, Canterbury, United Kingdom;
15	<sup>3</sup> Zouves Foundation for Reproductive Medicine, Foster City, California, USA.
16	<sup>4</sup> Department of Ecology and Evolutionary Biology, Princeton University,
17	Princeton, New Jersey, USA.
18	
19	*Corresponding author information: Manuel Viotti
20	manuel@zouvesfoundation.org
21	
22	
23	

24	
25	
26	ABSTRACT
27	
28	Study question:
29	Is a clinical trophectoderm (TE) biopsy a suitable predictor of chromosomal
30	aneuploidy in blastocysts?
31	
32	Summary answer:
33	In the analyzed group of blastocysts, a clinical TE biopsy was an excellent
34	representative of blastocyst karyotype in cases of whole chromosome
35	aneuploidy, but in cases of segmental (sub-chromosomal) aneuploidy, a TE
36	biopsy was a poor representative of blastocyst karyotype.
37	
38	What is known already:
39	Due to the phenomenon of chromosomal mosaicism, concern has been
40	expressed about the possibility of discarding blastocysts classified as aneuploid
41	by pre-implantation genetic testing for aneuploidy (PGT-A) that in fact contain a
42	euploid Inner Cell Mass (ICM). Previously published studies investigating
43	karyotype concordance between TE and ICM have examined small sample sizes
44	and/or have utilized chromosomal analysis technologies superseded by Next
45	Generation Sequencing (NGS). It is also known that blastocysts classified as
46	mosaic by PGT-A can result in healthy births. TE re-biopsy of embryos classified

47	as aneuploid can	potentially uncover	new instances	of mosaicism, b	out the
----	------------------	---------------------	---------------	-----------------	---------

- 48 frequency of such blastocysts is currently unknown.
- 49

#### 50 Study design, size, duration:

- 51 45 patients donated 100 blastocysts classified as uniform aneuploids (non-
- 52 mosaic) using PGT-A by NGS (n=93 whole chromosome aneuploids, n=7
- 53 segmental aneuploids). In addition to the original clinical TE biopsy used for
- 54 PGT-A, each blastocyst was subjected to an ICM biopsy as well as a second TE
- 55 biopsy. All biopsies were processed for chromosomal analysis by NGS, and
- 56 karyotypes compared to the original TE biopsy.
- 57

#### 58 **Participants/materials, setting, methods:**

- 59 Single IVF center with in-house PGT-A program and associated research
- 60 laboratory.
- 61

#### 62 Main results and the role of chance:

- 63 When one or more whole chromosomes were aneuploid in the clinical TE biopsy,
- 64 the corresponding ICM was aneuploid in 90 out of 93 blastocysts (96.8%). When
- 65 the clinical TE biopsy only contained segmental (sub-chromosomal)
- aneuploidies, the ICM was aneuploid in 4 out of 7 cases (57.1%). Blastocysts
- 67 showing an uploidy concordance between clinical TE biopsy and ICM were also
- 68 aneuploid in a second TE biopsy in 86 out of 88 cases (97.7%). In blastocysts

69	displaying clinical TE-ICM discordance, a second TE biopsy was aneuploid in
70	only 2 out of 6 cases (33.3%).

- 71
- 72

#### 73 Limitations, reasons for caution:

All embryos in this study had an initial classification of 'aneuploid' and not

<sup>75</sup> 'euploid' or 'mosaic'. Therefore, the findings of this study refer specifically to a TE

76 biopsy predicting aneuploidy in the remaining blastocyst, and cannot be

extrapolated to deduce the ability of a TE biopsy to predict euploidy in the

78 blastocyst. No conclusions should be drawn from this study about a mosaic TE

biopsy's ability to predict the karyotype of the corresponding blastocyst. Caution

should be exercised in generalizing the findings of the sample group of this study

to the general IVF blastocyst population. The segmental aneuploidy group only

82 contains 7 samples.

83

#### 84 Wider implications of the findings:

The high rate of intra-blastocyst concordance observed in this study concerning whole chromosome aneuploidy contributes experimental evidence to the validation of PGT-A at the blastocyst stage. Concomitantly, the results suggest potential clinical value in reassessing blastocysts deemed aneuploid by TE rebiopsy in select cases, particularly in instances of segmental aneuploidies. This

90 could impact infertility treatment for patients who only have blastocysts classified

91 as aneuploid by PGT-A available.

q	2
~	_

#### 93 Study funding/competing interest(s):

- 94 This study was supported by the Zouves Foundation for Reproductive Medicine
- 95 and Zouves Fertility Center. The authors have no competing interest to disclose.

96

- 97 Trial registration number:
- 98 Not applicable
- 99
- 100 Keywords: Aneuploidy, Concordance, Blastocyst, PGT-A, Mosaic

### 102 INTRODUCTION

104	A number of clinical trials have reported improved IVF outcomes following the
105	vetting of embryos for chromosomal abnormalities (Forman et al., 2013; Rubio et
106	al., 2017; Scott et al., 2013; Yang et al., 2012), and yet the IVF community is still
107	debating the appropriate use of preimplantation genetic testing for aneuploidy
108	(PGT-A, previously called PGS) (Practice Committees of the American Society
109	for Reproductive et al., 2018; Sermon et al., 2016). Skeptics of the technology
110	condemn the assumption that a 5-10 cell biopsy is representative of the
111	remaining embryo (Esfandiari et al., 2016; Gleicher and Orvieto, 2017). Indeed,
112	the phenomenon of mosaicism, the condition of containing two or more cell lines
113	with distinct chromosomal content (Taylor et al., 2014), provides a biological
114	rationale for that concern. A karyotypic categorization of the trophectoderm (TE),
115	the precursor to the placenta, might therefore not always be predictive of the
116	inner cell mass (ICM), which gives rise to the fetus.
117	One of the potential consequences of misclassification of embryos during
118	PGT-A is the deselection of viable embryos when a blastocyst is deemed
119	aneuploid by TE biopsy but in fact contains a euploid ICM (Esfandiari et al.,
120	2016; Schoolcraft et al., 2017; Vera-Rodriguez and Rubio, 2017). Some patients
121	are only capable of producing embryos classified as aneuploid by PGT-A even
122	after repeated IVF cycles, particularly with advanced age (Franasiak et al., 2014).
123	Such cases invariably lead to the abandonment of infertility treatment.

124	Previous studies investigating rates of TE-ICM chromosomal concordance
125	(expertly reviewed by Capalbo and Rienzi), while extremely valuable, have relied
126	on limited sample sizes or methodologies that have recently been superseded by
127	higher resolution genetic testing platforms (Capalbo and Rienzi, 2017). Next
128	Generation Sequencing (NGS), has been heralded as a PGT-A technique with
129	superior sensitivity for chromosomal mosaicism compared to aCGH, qPCR or
130	SNP array (Fragouli et al., 2017; Harton et al., 2017; Maxwell et al., 2016; Munne
131	et al., 2017; Munne and Wells, 2017) and has also been reported as highly
132	effective in detecting segmental (i.e. sub-chromosomal) losses and gains with
133	higher precision than previous methods (Lai et al., 2017).
134	The purpose of this study was specifically to test the hypothesis that a
135	blastocyst embryo classified as aneuploid by NGS-based PGT-A correctly
136	predicts the ploidy of the ICM in the majority of cases. Furthermore, by analyzing
137	a second TE biopsy we determined the frequency of blastocysts originally
138	classified as aneuploid that could be redefined as mosaic by re-biopsy.
139	

#### 141 MATERIALS AND METHODS

142

#### 143 Embryos and clinical PGT-A analysis by NGS

144

145 Blastocysts derived from patients seeking infertility treatment were 146 generated by in vitro fertilization and embryo culture as previously described 147 (Victor et al., 2017), and were evaluated using the Gardner system (Gardner and 148 Schoolcraft, 1999). As part of the embryo selection process, a clinical 5-10 cell 149 TE biopsy was collected and blastocysts were vitrified. The clinical TE biopsies 150 were subjected to whole genome amplification (WGA) with SurePlex reagents 151 (Illumina) followed by NGS-based PGT-A using Illumina's VeriSeq kit (Illumina) 152 on a MiSeq system (Illumina) according to the manufacturer's protocol and 153 described in detail elsewhere (Vera-Rodriguez et al., 2016). For guality control, 154 only samples satisfying the following cutoffs were used: number of Reads 155 Passing Filter: >0.25M; Average Q-Score: >30; Alignment Score: >30; DLR 156 (derivative log ratio): <0.4. Karyotype profiles were evaluated independently by 157 three analysts and consensus determined. Copy number variation (CNV) for 158 each chromosome was scored in Bluefuse Multi Analysis Software (Illumina) 159 according to guidelines defined by the Preimplantation Genetic Diagnosis 160 International Society (PGDIS), accessible at 161 'http://www.pgdis.org/docs/newsletter 071816.html': Profiles with copy number 162 scale values <1.2 and >2.8 were recorded as aneuploid, those with values

163	between 1.8 and 2.2 were recorded as euploid, and all others were recorded as
164	mosaic. These guidelines reflect the detection range of mosaicism by NGS PGT-
165	A, validated in various cell- and DNA-mixing experiments (Fragouli et al., 2017;
166	Maxwell et al., 2016; Munne et al., 2017; Munne and Wells, 2017). The resolution
167	of VeriSeq NGS is validated to detect segmental (sub-chromosomal)
168	aneuploidies of 20Mb or larger by the manufacturer, although detection of
169	regions down to 1.81Mb have been reported using this platform (Zheng et al.,
170	2015). In our center, we consider 'aneuploidy' to encompass both whole and
171	segmental chromosome abnormalities.
172	Supernumerary blastocysts classified as 'aneuploid' (no mosaics) by PGT-
173	A were donated to science by signed informed consent by 45 patients (average
174	age of $36.5 \pm 5.7$ ) and de-identified. This study was approved by the institutional
175	review board of the Zouves Foundation for Reproductive Medicine (OHRP
176	IRB00011505).
177	
178	
179	ICM and Second TE Biopsy Collection and Analysis
180	
181	ICM biopsies were isolated from vitrified-warmed blastocysts as outlined in the
182	legend for Fig. 1A, basing the technique on a protocol described previously
183	(Taylor et al., 2016) but omitting the exposure of samples to Ca <sup>2+</sup> /Mg <sup>2+</sup> -free
184	medium. Immediately following ICM biopsy an additional TE biopsy was
185	collected. All biopsies were washed three times to clear any loose cells or cellular

186	debris, and subsequently stored at -80°C until further processing. Biopsies were
187	subjected to NGS-based PGT-A (as detailed above), and results evaluated
188	independently by three analysts blinded to the analysis profile of the original,
189	clinically reported TE biopsy.
190	For transparency, all karyotype profiles of every biopsy analyzed in this
191	study are shown in the main or supplemental figures of the manuscript.
192	
193	
194	Immunofluorescence
195	
196	Whole blastocysts or biopsies were immersed in fixation buffer containing 4%
197	paraformaldehyde (EMS #15710) and 10% fetal bovine serum (FBS) (Seradigm
198	1500-050) in phosphate buffered saline (PBS) (Corning MT21040CM) for 10
199	minutes (min) at room temperature (rt), followed by three 1 min washes at rt in
200	stain buffer, composed of 0.1% Triton X-100 (TX-100) (Sigma X100-100ML) and
201	10% FBS in PBS. Samples were then immersed in permeabilization buffer (0.5%
202	TX-100, 10% FBS in PBS) for 30 min at rt, followed by three washes in stain
203	buffer. Samples were then exposed to stain buffer containing both primary
204	antibodies each in 1:200 concentrations over night at 4°C rocking on a nutator.
205	Primary antibodies were mouse anti-human GATA3 (Thermo Fisher MA1-028)
206	and rabbit anti-human OCT4A (Cell Signaling #2890). The next day, after three
207	washes in stain buffer, samples were immersed in stain buffer containing both
208	secondary antibodies each in 1:500 concentrations for 2-3 hours at rt. Secondary

209	antibodies were goat anti-mouse IgG AlexaFluor488 (Thermo Fisher A11029)
210	and goat anti-rabbit IgG AlexaFluor647 (Thermo Fisher A21245). After three
211	washes in stain buffer, samples were exposed to nuclear stain (Hoechst 33342,
212	Thermo Fisher H3570) diluted at 1:1000 in stain buffer for 30 min at rt, followed
213	by three more washes in stain buffer. Samples were placed in glass bottom
214	dishes (MatTek P35G-1.5-20-C) in small drops of stain buffer overlaid with
215	mineral oil (Sigma M5904), and imaged with a LSM 780 Confocal microscope
216	(Zeiss).
217	
218	
219	Analysis of tissue relatedness
220	In cases of clinical TE-ICM karyotype discordance we confirmed tissue
221	relatedness by a DNA fingerprinting method that utilizes SNP analysis and
222	linkage disequilibrium known as 'Tilde' (Vohr et al., 2015). A full explanation of
223	the adaptation of this method to PGT-A samples with low coverage NGS is
224	detailed in Supplemental Data 1.
225	
226	Statistical analysis of correlation between morphology and karyotype
227	discordance
228	Analysis and graph preparation were performed in Prism 6 (GraphPad).
229	Differences between groups were assessed by Chi-square test for trend with
230	95% confidence levels. Significance was defined when $P < 0.05$ .
231	

#### 233 RESULTS

234

#### 235 Isolation of ICM and Second TE Biopsies

236 We adopted a modified ICM-biopsy procedure previously outlined (Taylor 237 et al., 2016), which permitted us to collect an ICM biopsy and subsequently a 238 second TE biopsy in blastocysts (Fig. 1A and Video 1). Immunofluorescence was 239 used to confirm accurate isolation of intended cells. In whole blastocysts the 240 pluripotency factor OCT4 was present at high levels in the ICM and low levels in 241 the TE, while GATA3 was exclusively expressed in cells of the TE as previously 242 shown (Deglincerti et al., 2016). Analysis of matched TE-ICM biopsies from 12 243 blastocysts indicated that both biopsy types exclusively contained cells of their 244 intended lineage and were devoid of contamination from the other cell type (Fig. 245 1B and Supplemental Fig. 1). Nuclear counterstain by Hoechst did not reveal any 246 cells with fragmenting or apoptotic nuclear material, suggesting that the biopsy 247 technique did not disrupt individual cells (Fig. 1B and Fig. S1). On average, TE 248 biopsies comprised 7.6 cells (+/- 1.3 SD) while ICM biopsies comprised 7.3 cells 249 (+/- 2.0 SD).

250

#### 251 Clinical TE-ICM Biopsy Concordant Blastocysts

252 Of the 100 blastocysts originally classified as aneuploid by clinical 253 (original) TE biopsy, 93 had ICMs that were also classified as aneuploid, which 254 we denote as aneuploid-aneuploid concordant (Fig. 2 and Table I). Importantly,

255	when only considering blastocysts with whole chromosomal aneuploidies (single
256	or multiple) in their clinical TE biopsies, aneuploidy in the ICM was present in 90
257	out of 93 cases (96.8%). On the other hand, when considering blastocysts with
258	only segmental (sub-chromosomal) aneuploidies in their clinical TE biopsies,
259	aneuploidy in the ICM was present in only 3 out of 7 cases (42.9%).
260	In aneuploid-aneuploid concordant blastocysts, analysis of second TE
261	biopsies showed aneuploidy in 86 out of 88 cases, equaling 97.7% (Table I). The
262	remaining two samples showed a mosaic pattern in their respective second TE
263	biopsies. In five samples a second TE biopsy could not be retrieved.
264	The 93 clinical TE-ICM aneuploid-aneuploid concordant blastocysts could
265	be further subdivided in two groups. 79 were blastocysts that had perfectly
266	matching karyotypes in the clinical TE and ICM biopsies (i.e., all the same
267	chromosomes possessed the same aneuploidies in both tissues), which we
268	denoted as aneuploid-aneuploid perfect concordant (Fig. 2, Table I, and for the
269	karyotypic profiles see Supplemental Fig. 2). Such instances are likely
270	consequences of meiotic errors, as the identical aneuploidy is present in both TE
271	and ICM tissues (see Supplemental Data 2 for more detailed interpretations).
272	The remaining 14 out of 93 blastocysts had dissimilar aneuploidies in the
273	clinical TE and ICM biopsies, which we denoted as aneuploid-aneuploid
274	imperfect concordant (Fig. 2, Table I, and for the karyotypic profiles see
275	Supplemental Fig. 2). Interestingly, most such blastocysts (10 out of 14) showed
276	the same aneuploid chromosome(s) in the ICM biopsy as the clinical TE biopsy

277	(presumed	consequence	of meiotic error)	, but contained	additional mosaic
-----	-----------	-------------	-------------------	-----------------	-------------------

events in the ICM (resulting from mitotic error), often segmental in nature.

279 See Supplemental Data 2 for interpretations of chromosomal error

etiologies in samples of the aneuploid-aneuploid *imperfect* concordant group.

281

#### 282 Clinical TE-ICM Biopsy Discordant Blastocysts

Of the 100 blastocysts tested, we observed two cases in which the clinical TE biopsy was uniformly aneuploid but the ICM was mosaic (Fig. 2, Table I, and for the karyotypic profiles see Fig. 3) Supplemental Data 2 contains more detailed interpretations of their karyotypes.

287 Five out of 100 blastocysts had euploid ICMs while their clinical TE 288 biopsies contained aneuploidies (Fig. 2, Table I, and for the karyotypic profiles 289 see Fig. 3). Blastocyst #96 was the only case in which the clinical TE biopsy had 290 a whole chromosomal aneuploidy (gain of chromosome 12, note that the 291 karyotype profile enters into the 2.8-3.0 copy number region) but displayed 292 euploidy in the ICM as well as in the second TE biopsy. 293 The remaining four samples (blastocysts #97-#100) contained segmental 294 aneuploidies in their original TE biopsies, but euploid ICM biopsies. For 295 blastocyst #97, the clinical and second TE biopsies contained the same 296 segmental aneuploidy, thereby suggesting euploidy confined to the ICM. This 297 would be consistent with a mitotic event happening before or at the time of 298 lineage segregation but in the progenitor cell of a large part of the TE. 299 Blastocysts #98 and #99 displayed mosaicism in their respective second TE

300	biopsies, revealing the occurrence of mitotic errors in the TE lineage. For one
301	blastocyst (#100), the second TE biopsy did not yield results due to a failed WGA
302	reaction. (Global WGA failure rate for this study is 1 out of 221, or 0.4%). In total,
303	of the clinical TE-ICM discordant blastocysts (aneuploid-euploid or aneuploid-
304	mosaic) yielding information in the second TE biopsy, only 2 out of 6 $(33.3\%)$
305	were uniformly aneuploid.
306	In cases of clinical TE-ICM biopsy discordance, there existed the
307	possibility of sample contamination or mislabeling. Notably, in the 100 embryos
308	tested, the sex chromosomes (XX or XY) were always concordant between
309	biopsies taken within the same blastocyst. Further, for each of the seven
310	blastocysts that produced discordant results, we performed DNA fingerprinting to
311	confirm that the clinical TE and ICM biopsies were derived from the same
312	respective embryos (Fig. 4 and Supplemental Fig. 3).
313	Finally, we determined whether poor blastocyst morphology impacted
314	karyotype discordance. The analysis indicated that neither blastocyst stage nor
315	ICM/TE grade affected the likelihood of intra-blastocyst karyotype inconsistencies
316	(Supplemental Fig. 4).
317	

318

#### 320 DISCUSSION

321

322 Some parties have argued that PGT-A should not be performed under any 323 circumstance and one of the criticisms of the technology questions whether a 324 clinical TE biopsy is a valid genetic representative of the embryo (Esfandiari et 325 al., 2016; Mastenbroek and Repping, 2014; Sermon et al., 2016). A study basing 326 its rationale on mathematical modeling has claimed that a typical TE cell biopsy 327 cannot determine embryo ploidy accurately enough for clinical use (Gleicher et 328 al., 2017). One of the ensuing concerns that has been expressed is the 329 possibility of erroneously discarding viable embryos (Practice Committees of the 330 American Society for Reproductive et al., 2018). Here however, we provide 331 experimental evidence using NGS that a TE biopsy classified as aneuploid is 332 commonly predictive of an uploidy in the ICM. In our experience, a whole 333 chromosome aneuploidy in a clinical TE biopsy is predictive of aneuploidy in the 334 ICM in 96.8% of cases (sample size n=93), although for a segmental aneuploidy 335 this decreases significantly to 42.9% (n=7). 336 A blastocyst with an aneuploid TE and ICM due to meiotic error is in 337 principle exceptionally unlikely to result in healthy pregnancy (Adashi and 338 McCoy, 2017). Although various corrective mechanisms for aneuploidies in 339 human embryos have been proposed (differential proliferation/depletion, 340 preferential lineage allocation, self-correction) (Capalbo and Rienzi, 2017; 341 McCoy, 2017) and have also been conceptually demonstrated in mouse embryos

342 (Bolton et al., 2016) and human embryonic stem cells (hESC) (Munne et al.,

343 2005), most models describe the out-competition of an euploid cells by euploid
344 cells in the mosaic setting, not the conversion of an entirely an euploid embryo to
345 an entirely euploid one.

346 The observation that segmentals had a drastically different rate of clinical 347 TE-ICM discordance compared to whole chromosome aneuploids highlights the 348 difference in mechanistic origins of these two types of aneuploidies. Whole 349 chromosome aneuploidies can arise during meiosis or mitosis by different 350 mechanisms that include non-disjunction, anaphase lag, and endoreplication 351 (Taylor et al., 2014), but the majority are believed to be derived from meiotic 352 errors in the oocyte (Nagaoka et al., 2012). The majority of segmental 353 aneuploidies on the other hand are mitotic in origin and are thought to arise 354 during the first few cell divisions after fertilization (Babariya et al., 2017). Cell 355 cycle control is thought to be more lax during the first days of embryogenesis due 356 to rapid mitoses primarily controlled by maternal RNA and proteins, leading to an 357 increased incidence of double strand breaks which upon faulty correction 358 mechanisms result in segmental duplications or deletions when left unresolved 359 by a strained cell cycle machinery (Babariya et al., 2017). Consequently, 360 segmental aneuploidies will often be represented in mosaic configurations at a 361 whole blastocyst level, likely translating in the high TE-ICM discordance rate 362 observed for the segmental aneuploidy group in this study. 363 Out of 93 blastocysts with whole chromosome aneuploidies (single or 364 multiple) in a clinical TE biopsy, three embryos had a discordant ICM: two

365 contained mosaic ICM biopsies, and one had a euploid ICM. Consequently, the

366 karyotype of these three blastocysts should be re-classified from aneuploid to 367 mosaic, since on a whole embryo level they contained aneuploid and euploid 368 cells. This re-categorization would have changed the status of the blastocysts 369 from 'not recommended for transfer' to 'possible transfer if no euploid embryos 370 available'. Mosaic embryos have recently been considered for transfer in several 371 clinics, producing healthy pregnancies albeit with considerably lower implantation 372 rates than blastocysts classified as euploid (Fragouli et al., 2017; Lledo et al., 373 2017; Munne et al., 2017; Spinella et al., 2018). 374 From a clinical standpoint, our findings may support re-biopsy of 375 blastocysts in patients that have only produced embryos classified as aneuploid 376 (particularly segmentals) by initial TE biopsy after repeated IVF cycles, an

377 occurrence that happens with relative frequency especially with advanced

378 maternal age (Franasiak et al., 2014). It could also affect those patients that have

379 unsuccessfully transferred their embryos classified as 'euploid' and 'mosaic', and

only have 'aneuploid' samples remaining. In our study, all blastocysts had an
initial, clinical TE biopsy that was uniformly aneuploid. When a second TE biopsy

382 was either mosaic or euploid, such a blastocyst had a 66% chance to contain an

383 ICM that was either mosaic or euploid as well. On the other hand, in cases where

the second TE biopsy was aneuploid, the ICM was mosaic in only 1.1% of cases,

385 and there were no euploid ICM instances. Therefore, our results suggest that TE

386 re-biopsy can reveal whether the ICM is mosaic or euploid, helping to identify

387 new blastocysts for possible clinical use that were originally not recommended

388 for transfer due to aneuploidy in the clinical TE biopsy. Importantly, while the act

of re-biopsy might negatively affect blastocysts, re-biopsied blastocysts can lead
to healthy pregnancies albeit with lower efficiency than single-biopsied
blastocysts (Bradley et al., 2017). Nevertheless, more research is necessary to
determine the short and long term effects of TE re-biopsy, and recommendation
of routine re-biopsy of blastocysts classified as aneuploid is undoubtedly
premature.

The confirmed existence of clinical TE-ICM discordant embryos could also help explain the rare accounts of healthy pregnancies resulting from transfer of embryos classified as aneuploid by PGT-A (Gleicher et al., 2016), although it must be pointed out that to our knowledge, there exist no reports of such events when using blastocyst stage NGS-based PGT-A.

400 It is important to note that our determined rates of clinical TE-ICM 401 concordance apply specifically to blastocysts classified as 'uniform aneuploid' (no 402 mosaics) by PGT-A. Having observed an overall 7% clinical TE-ICM discordance 403 rate in our samples, we cannot assume the inverse: that 7% of blastocysts 404 classified as euploid contain an aneuploid ICM. A further intriguing and clinically 405 important question is what a clinical TE biopsy showing mosaicism says about 406 the ICM. Unfortunately, our study cannot shed light on that point. 407 A further limitation of this study was that not all cells were analyzed for 408 intra-blastocyst karyotypic concordance. The ICM biopsies isolated (averaging

409 7.3 cells) collected the bulk of ICM cells but invariably left residual ICM cells

410 behind. On average, we collected 15 TE cells from the two combined TE biopsies

411 of each blastocyst, hence a substantial portion of the TE was left unanalyzed.

412 From the technical standpoint we were unable to isolate more cells from a

413 specific tissue without contamination from the other lineage. As a result,

414 instances of karyotype discordance could remain concealed.

415 While highly controversial, the concept of transferring embryos testing 416 aneuploid by PGT-A is a real subject of discussion in both scientific (Gleicher et 417 al., 2016) and mainstream media (Hall, 2017). The upheaval created by these 418 viewpoints has partly been bolstered by the yet unspecified capability of a single 419 clinical TE biopsy to reflect the state of the ICM and remaining TE. With regard to 420 this guestion our findings contribute experimental validation on the practice of 421 PGT-A at the blastocyst stage, considering the high intra-blastocyst aneuploidy 422 concordance rates, especially in the case of whole chromosome losses or gains. 423 If indeed the group of blastocysts analyzed in this study is representative of the 424 general body of IVF blastocysts, it would mean that when selecting an embryo 425 classified as 'aneuploid' by PGT-A for uterine transfer, it almost always contains 426 aneuploidy in the entire blastocyst. Unless robust self-correction mechanisms do 427 in fact exist, said embryo would invariable lead to failed implantation, 428 miscarriage, or chromosomally abnormal babies. Segmental aneuploidies on the 429 other hand are rarely concordant; if our observations are confirmed in a larger 430 sample group they should be regarded as their own distinct class when 431 prioritizing or de-selecting embryos for transfer in the clinic.

432

433

#### 435 Authors' Roles

- 436 A.R.V., D.K.G., A.L., C.G.Z., F.L.B., R.C.M., and M.V. designed the experiments.
- 437 A.R.V, A.J.B., J.C.T. A.E.M., L.T.L. R.C.M., and M.V. performed the experiments.
- 438 A.R.V., D.K.G., F.L.B., and M.V. wrote the manuscript.
- 439 A.J.B., J.C.T. A.E.M., L.T.L., A.L., C.G.Z., and R.C.M edited the manuscript.

440

#### 441 Acknowledgements

- 442 We would like to thank the entire ZFC staff for supporting this study in many
- 443 ways, including sample storage, reagent preparation, and discussions.

444

- 445 Funding
- 446 This study was supported by the Zouves Foundation for Reproductive Medicine
- 447 and Zouves Fertility Center.

- 449 **Conflict of Interest**
- 450 The authors have no conflict of interest to declare.

451		
452	FIGURE LEGE	NDS
453		
454	Figure 1	
455	Validatio	on of biopsy and PGT-A methods used in the study
456		
457	(A)	Stills from video depicting isolation of ICM biopsy in blastocysts.
458		The blastocyst is immobilized with a holding pipet touching the
459		polar TE (adjacent to the ICM), and laser pulses are
460		administered through the zona and mural TE opposite the ICM
461		creating an opening. A biopsy pipette is introduced and guided
462		to the ICM, which is suctioned out through the opening. Once a
463		portion of ICM cells are extracted past the zona, they are
464		exposed to laser pulses aimed at cell-cell junctions to isolate a
465		5-10 cell biopsy.
466		
467	(B)	Nuclear counterstain (Hoechst) and immunofluorescent stains
468		for the TE marker GATA3 and ICM marker OCT4 in a whole
469		human blastocyst and isolated TE and ICM biopsies. See
470		additional samples in Figure S1. Scale bars = $25\mu$ m.
471		
472		
473		

474	Figure 2.
475	Summary of paired clinical TE-ICM comparison results
476	
477	Dot plot displays results for all blastocysts, regardless of aneuploidy type.
478	Pie charts depict data stratified by nature of aneuploidy detected in the
479	original TE biopsy for each blastocyst.
480	
481	
482	Figure 3.
483	NGS-based PGT-A karyotype profiles for biopsies in blastocysts with
484	discordant clinical TE-ICM patterns
485	
486	See Table I for the interpretation of each profile.
487	
488	
489	Figure 4.
490	Log-likelihood ratios of relatedness between tissues in blastocysts
491	with clinical TE-ICM discordance
492	
493	In green, controls comparing biopsies from embryos derived from patients
494	expected to be unrelated, showing negative values. In red, comparisons
495	between biopsies from blastocysts derived from the same patient (full-
496	sibs) showing positive values. In purple, comparisons between clinical TE

497	and ICM biopsies for each blastocyst classified as discordant in the study,
498	showing highly positive log-likelihood ratios of relatedness.
499	
500	Supplemental Figure 1.
501	Validation of contamination-free ICM and TE biopsy technique
502	
503	Nuclear counterstain (Hoechst) and immunofluorescent stains for the TE
504	marker GATA3 and ICM marker OCT4 in matched isolated TE and ICM
505	biopsies. Numbers in nuclear stain panels indicate total number of cells for
506	each biopsy. Numbers in TE stain panels indicate incidence of cells
507	positive for GATA3. Numbers in ICM stain panels indicate incidence of
508	cells with high nuclear signal for OCT4. Scale bars = $25\mu$ m.
509 510	
511	Supplemental Figure 2.
512	Karyotype profiles analyzed in this study (continued from Fig. 3)
513	
514	Use Table I as a reference for resulting analysis.
515	
516	
517	Supplemental Figure 3.
518	Genotypes of discordant blastocysts visualized in reference ancestry
519	space
520	

Page 29 of 87

521	Supplemental Figure 4
522	Analysis of correlation between morphology and intra-blastocyst
523	karyotype discordance
524	Blastocysts were evaluated using the Gardner system, that assigns a
525	number score for blastocyst expansion stage (1-6 from least to most
526	progressed), and letter scores for ICM and TE grades (C-A from worst to
527	best quality) (Gardner and Schoolcraft, 1999). Stage of blastocyst was
528	analyzed when there was any intra-blastocyst karyotype inconsistency (A),
529	when an inconsistency was specific to the ICM (B), and when an
530	inconsistency was specific to the TE (C). Grade of the ICM was analyzed
531	when there was any karyotype inconsistency between clinical TE biopsy
532	and the ICM (D). Grade of the TE was analyzed when there was any
533	karyotype inconsistency between clinical TE and second TE biopsy (E).
534	Note that in graphs A, C, and D, sample numbers do not add up to 100
535	because in six cases the second TE biopsy was not processed (in five
536	cases the biopsy could not be collected, and in one case there was a
537	failed WGA reaction.
538	

539	
540	TABLE LEGENDS
541	
542	Table I. List of blastocysts, clinical TE, ICM, and second TE biopsies analyzed in
543	this study.
544	
545	
546	Supplemental Table I. Blastocysts and assigned 1000 Genomes super-
547	populations.
548	
549	Supplemental Data 1. Adaptation of 'Tilde' to PGT-A samples with low coverage
550	NGS.
551	
552	Supplemental Data 2. Additional interpretations of chromosomal error etiologies

553 in blastocysts.

#### 554 **BIBLIOGRAPHY**

555

- 557 Adashi, E.Y., and McCoy, R.C. (2017). Technology versus biology: the limits of
- 558 pre-implantation genetic screening: Better methods to detect the origin of
- aneuploidy in pre-implantation embryos could improve the success rate of
- 560 artificial reproduction. EMBO Rep *18*, 670-672.
- 561 Babariya, D., Fragouli, E., Alfarawati, S., Spath, K., and Wells, D. (2017). The
- 562 incidence and origin of segmental aneuploidy in human oocytes and
- 563 preimplantation embryos. Hum Reprod 32, 2549-2560.
- Bolton, H., Graham, S.J., Van der Aa, N., Kumar, P., Theunis, K., Fernandez
- 565 Gallardo, E., Voet, T., and Zernicka-Goetz, M. (2016). Mouse model of
- 566 chromosome mosaicism reveals lineage-specific depletion of aneuploid cells and
- 567 normal developmental potential. Nat Commun 7, 11165.
- 568 Bradley, C.K., Livingstone, M., Traversa, M.V., and McArthur, S.J. (2017). Impact
- 569 of multiple blastocyst biopsy and vitrification-warming procedures on pregnancy
- 570 outcomes. Fertil Steril 108, 999-1006.
- 571 Capalbo, A., and Rienzi, L. (2017). Mosaicism between trophectoderm and inner 572 cell mass. Fertil Steril *107*, 1098-1106.
- 573 Deglincerti, A., Croft, G.F., Pietila, L.N., Zernicka-Goetz, M., Siggia, E.D., and
- 574 Brivanlou, A.H. (2016). Self-organization of the in vitro attached human embryo.
- 575 Nature *5*33, 251-254.
- 576 Esfandiari, N., Bunnell, M.E., and Casper, R.F. (2016). Human embryo
- 577 mosaicism: did we drop the ball on chromosomal testing? J Assist Reprod Genet
- 578 33, 1439-1444.

579 Fo	orman, E.J.,	Hong, K.H.	, Ferry, K.M.	, Tao, X.,	Taylor,	D., Levy, B.	, Treff, N.R.,
--------	--------------	------------	---------------	------------	---------	--------------	----------------

and Scott, R.T., Jr. (2013). In vitro fertilization with single euploid blastocyst

transfer: a randomized controlled trial. Fertil Steril *100*, 100-107 e101.

- 582 Fragouli, E., Alfarawati, S., Spath, K., Babariya, D., Tarozzi, N., Borini, A., and
- 583 Wells, D. (2017). Analysis of implantation and ongoing pregnancy rates following
- the transfer of mosaic diploid-aneuploid blastocysts. Hum Genet *136*, 805-819.
- 585 Franasiak, J.M., Forman, E.J., Hong, K.H., Werner, M.D., Upham, K.M., Treff,

586 N.R., and Scott, R.T., Jr. (2014). The nature of aneuploidy with increasing age of

587 the female partner: a review of 15,169 consecutive trophectoderm biopsies

- evaluated with comprehensive chromosomal screening. Fertil Steril *101*, 656-663e651.
- 590 Gardner, D.K., and Schoolcraft, W.B. (1999). In vitro culture of human

591 blastocysts. . In Towards reproductive certainty: fertility and genetics beyond, J.

R., and M. D., eds. (Camforth, UK: Parthenon Publishing), pp. 378-388.

593 Gleicher, N., Metzger, J., Croft, G., Kushnir, V.A., Albertini, D.F., and Barad, D.H.

594 (2017). A single trophectoderm biopsy at blastocyst stage is mathematically

595 unable to determine embryo ploidy accurately enough for clinical use. Reprod596 Biol Endocrinol *15*, 33.

597 Gleicher, N., and Orvieto, R. (2017). Is the hypothesis of preimplantation genetic
598 screening (PGS) still supportable? A review. J Ovarian Res *10*, 21.

- 599 Gleicher, N., Vidali, A., Braverman, J., Kushnir, V.A., Barad, D.H., Hudson, C.,
- 600 Wu, Y.G., Wang, Q., Zhang, L., Albertini, D.F., et al. (2016). Accuracy of
- 601 preimplantation genetic screening (PGS) is compromised by degree of
- 602 mosaicism of human embryos. Reprod Biol Endocrinol *14*, 54.
- Hall, S.S. (2017). A new last chance. In New York Magazine.

604	Harton, G.L., Cinnioglu, C., and Fiorentino, F. (2017). Current experience
605	concerning mosaic embryos diagnosed during preimplantation genetic screening.
606	Fertil Steril 107, 1113-1119.

Lai, H.H., Chuang, T.H., Wong, L.K., Lee, M.J., Hsieh, C.L., Wang, H.L., and
Chen, S.U. (2017). Identification of mosaic and segmental aneuploidies by next-

609 generation sequencing in preimplantation genetic screening can improve clinical 610 outcomes compared to array-comparative genomic hybridization. Mol Cytogenet

- 611 *10*, 14.
- 612 Lledo, B., Morales, R., Ortiz, J.A., Blanca, H., Ten, J., Llacer, J., and Bernabeu,
- R. (2017). Implantation potential of mosaic embryos. Syst Biol Reprod Med 63,206-208.
- 615 Mastenbroek, S., and Repping, S. (2014). Preimplantation genetic screening:
- back to the future. Hum Reprod 29, 1846-1850.
- 617 Maxwell, S.M., Colls, P., Hodes-Wertz, B., McCulloh, D.H., McCaffrey, C., Wells,

D., Munne, S., and Grifo, J.A. (2016). Why do euploid embryos miscarry? A

619 case-control study comparing the rate of aneuploidy within presumed euploid

620 embryos that resulted in miscarriage or live birth using next-generation

- 621 sequencing. Fertil Steril *106*, 1414-1419 e1415.
- 622 McCoy, R.C. (2017). Mosaicism in Preimplantation Human Embryos: When
- 623 Chromosomal Abnormalities Are the Norm. Trends Genet *33*, 448-463.
- Munne, S., Blazek, J., Large, M., Martinez-Ortiz, P.A., Nisson, H., Liu, E.,
- Tarozzi, N., Borini, A., Becker, A., Zhang, J., *et al.* (2017). Detailed investigation
- 626 into the cytogenetic constitution and pregnancy outcome of replacing mosaic
- 627 blastocysts detected with the use of high-resolution next-generation sequencing.
- 628 Fertil Steril *108*, 62-71 e68.

- Munne, S., Velilla, E., Colls, P., Garcia Bermudez, M., Vemuri, M.C., Steuerwald,
- N., Garrisi, J., and Cohen, J. (2005). Self-correction of chromosomally abnormal
- embryos in culture and implications for stem cell production. Fertil Steril *84*,1328-1334.

Munne, S., and Wells, D. (2017). Detection of mosaicism at blastocyst stage with
the use of high-resolution next-generation sequencing. Fertil Steril *107*, 10851091.

- 636 Nagaoka, S.I., Hassold, T.J., and Hunt, P.A. (2012). Human aneuploidy:
- 637 mechanisms and new insights into an age-old problem. Nat Rev Genet 13, 493-
- 638 504.

639 Practice Committees of the American Society for Reproductive, M., the Society

640 for Assisted Reproductive Technology. Electronic address, A.a.o., Practice

641 Committees of the American Society for Reproductive, M., and the Society for

642 Assisted Reproductive, T. (2018). The use of preimplantation genetic testing for

643 aneuploidy (PGT-A): a committee opinion. Fertil Steril *109*, 429-436.

Rubio, C., Bellver, J., Rodrigo, L., Castillon, G., Guillen, A., Vidal, C., Giles, J.,
Ferrando, M., Cabanillas, S., Remohi, J., *et al.* (2017). In vitro fertilization with
preimplantation genetic diagnosis for aneuploidies in advanced maternal age: a
randomized, controlled study. Fertil Steril *107*, 1122-1129.

Schoolcraft, W., Meseguer, M., and Global Fertility Alliance. Electronic address,
a.t.i.c. (2017). Paving the way for a gold standard of care for infertility treatment:
improving outcomes through standardization of laboratory procedures. Reprod
Biomed Online *35*, 391-399.

- 652 Scott, R.T., Jr., Upham, K.M., Forman, E.J., Hong, K.H., Scott, K.L., Taylor, D.,
- Tao, X., and Treff, N.R. (2013). Blastocyst biopsy with comprehensive
- 654 chromosome screening and fresh embryo transfer significantly increases in vitro

- 655 fertilization implantation and delivery rates: a randomized controlled trial. Fertil656 Steril *100*, 697-703.
- 657 Sermon, K., Capalbo, A., Cohen, J., Coonen, E., De Rycke, M., De Vos, A.,
- 658 Delhanty, J., Fiorentino, F., Gleicher, N., Griesinger, G., et al. (2016). The why,
- the how and the when of PGS 2.0: current practices and expert opinions of
- 660 fertility specialists, molecular biologists, and embryologists. Mol Hum Reprod 22,
- 661 845-857.
- 662 Spinella, F., Fiorentino, F., Biricik, A., Bono, S., Ruberti, A., Cotroneo, E., Baldi,
- 663 M., Cursio, E., Minasi, M.G., and Greco, E. (2018). Extent of chromosomal
- 664 mosaicism influences the clinical outcome of in vitro fertilization treatments. Fertil
- 665 Steril *109*, 77-83.
- 666 Taylor, T.H., Gitlin, S.A., Patrick, J.L., Crain, J.L., Wilson, J.M., and Griffin, D.K.
- 667 (2014). The origin, mechanisms, incidence and clinical consequences of
- 668 chromosomal mosaicism in humans. Hum Reprod Update 20, 571-581.
- 669 Taylor, T.H., Griffin, D.K., Katz, S.L., Crain, J.L., Johnson, L., and Gitlin, S.
- 670 (2016). Technique to 'Map' Chromosomal Mosaicism at the Blastocyst Stage.
- 671 Cytogenet Genome Res *149*, 262-266.
- 672 Vera-Rodriguez, M., Michel, C.E., Mercader, A., Bladon, A.J., Rodrigo, L.,
- 673 Kokocinski, F., Mateu, E., Al-Asmar, N., Blesa, D., Simon, C., et al. (2016).
- 674 Distribution patterns of segmental aneuploidies in human blastocysts identified
- by next-generation sequencing. Fertil Steril *105*, 1047-1055 e1042.
- 676 Vera-Rodriguez, M., and Rubio, C. (2017). Assessing the true incidence of
- 677 mosaicism in preimplantation embryos. Fertil Steril *107*, 1107-1112.
- 678 Victor, A.R., Brake, A.J., Tyndall, J.C., Griffin, D.K., Zouves, C.G., Barnes, F.L.,
- and Viotti, M. (2017). Accurate quantitation of mitochondrial DNA reveals uniform

- 680 levels in human blastocysts irrespective of ploidy, age, or implantation potential.
- 681 Fertil Steril *107*, 34-42 e33.
- Vohr, S.H., Buen Abad Najar, C.F., Shapiro, B., and Green, R.E. (2015). A
- 683 method for positive forensic identification of samples from extremely low-
- 684 coverage sequence data. BMC Genomics *16*, 1034.
- Yang, Z., Liu, J., Collins, G.S., Salem, S.A., Liu, X., Lyle, S.S., Peck, A.C., Sills,
- 686 E.S., and Salem, R.D. (2012). Selection of single blastocysts for fresh transfer
- via standard morphology assessment alone and with array CGH for good
- 688 prognosis IVF patients: results from a randomized pilot study. Mol Cytogenet 5,689 24.
- Sheng, H., Jin, H., Liu, L., Liu, J., and Wang, W.H. (2015). Application of next-
- 691 generation sequencing for 24-chromosome aneuploidy screening of human
- 692 preimplantation embryos. Mol Cytogenet *8*, 38.
- 693

#### Clinical TE-ICM Aneuploid-Aneuploid Perfect Concordant

TABLE I

#### Clinical TE-ICM Aneuploid-Aneuploid Imperfect Concordant

Blastocyst Study id# Gardner Grade	Clinical TE Biopsy	ICM Biopsy	Second TE Biopsy
1 4BB	45,XY,-4	45.XY4	45,XY,-4
2 4BB	45,XX,-7	45,XX,-7	45,XX,-7
3 5BB	45,XX,-8	45,XX,-8	46,XX,-8*,+15*,+20*
4 3AB	45,XY,-8	45,XY,-8	45,XY,-8
5 5BC	45,XX,-10	45,XX,-10	45,XX,-10
6 4BB	45,XY,-11	45,XY,-11	45,XY,-11
7 5BB	45,XX,-13	45,XX,-13	45,XX,-13
0.580	40,A1,-13	45,X1,-15	1//d 45 VV 14
10 4PR	45 XY -14	45 XY -14	45 XY -14
11 2BB	45 XX -15	45 XX -15	45 XX -15
12 4BC	45 XX -15	45 XX -15	n/a
13 3BB	45,XX,-16	45,XX,-16	45,XX,-16
14 4BC	45,XX,-16	45,XX,-16	45,XX,-16
15 4BB	45,XX,-16	45,XX,-16	45,XX,-16
16 4BB	45,XY,-16	45,XY,-16	45,XY,-16
17 5CC	45,XY,-16	45,XY,-16	45,XY,-16
18 5CC	45,XX,-17	45,XX,-17	45,XX,-17
19 5AB	45,XY,-18	45,XY,-18	45,XY,-18,del(10)(q11.21q26.3)*
20 4BB	45,XX,-21	45,XX,-21	45,XX,-21
21 5BB	45,XX,-21	45,XX,-21	45,XX,-21
22 4CB	45,XY,-21	45,XY,-21	45,XY,-21
23 5BB	45,XX,-22	45,XX,-22	45,XX,-22
24 488	45,88,-22	45,88,-22	45,8,8,-22
20 0BB	40, A1, -22	45,X1,-22	45,X1,-22
20 400	45 XY -22	45 XV -22	45 XV -22
27 444	45,X1,-22	45,X1,-22	46 XX dup(X)(p22 33p21 1) dup(X)(p22 3p25)*
20 400	40,A	47 YY ±1	40,XX,dup(X)(p22.33p21.1),dup(X)(d22.3d23)
30 4BC	47 XY +4	47 XY +4	47,XX +4
31 4AB	47 XX +13	47 XX +13	47 XX +13
32 3BB	47 XX +13	47 XX +13	47 XX +13
33 5AA	47.XY.+15	47.XY.+15	47.XY.+15
34 4CB	47,XX,+16	47,XX,+16	47,XX,+16
35 3AB	47,XX,+16	47,XX,+16	47,XX,+16
36 5BB	47,XX,+16	47,XX,+16	n/a
37 5BB	47,XY,+16	47,XY,+16	47,XY,+16
38 4BB	47,XY,+16	47,XY,+16	47,XY,+16
39 4BC	47,XY,+16	47,XY,+16	47,XY,+16
40 5BB	47,XY,+17	47,XY,+17	47,XY,-10*,+17
41 4BB	47,XY,+18	47,XY,+18	47,XY,+18
42 4CB	47,XY,+18	47,XY,+18	47,XY,+18
43 5BB	47,XY,+19	47,XY,+19	47,XY,+19
44 5AB	47,XX,+20	47,XX,+20	47,XX,+20
45 480	47,XX,+21	47,XX,+21	47,XX,+21
40 SUB	46,XX,+21(X2)	48,XX,+21(X2)	48,XX,+21(X2)
47 400	47,XX,T22	47, XX, T22	47, XX, T22
40 548	47, XX ±22	47,XX,122	47, XX, 122
50 4BB	47 XY +22	47 XY +22	47,XX,+22
51 500	47 XY +22	47 XY ±22	47 XY +22
52 4BC	47.XY.+22	47.XY.+22	47.XY.+22
53 3CC	47,XY,+22	47,XY,+22	47,XY,+22,del(1)(q25.2q44)
54 3BB	44,XY,-10,-22	44,XY,-10,-22	44,XY,-10,-22
55 4BB	46,XX,-12,+16	46,XX,-12,+16	46,XX,-12,+16
56 4CC	44,XX,-14,-19	44,XX,-14,-19	44,XX,-14,-19
57 4BB	44,XY,-18,-21	44,XY,-18,-21	44,XY,-18,-21
58 4BC	46,XY,-19,+22	46,XY,-19,+22	n/a
59 5BB	46,XY,-21,+22	46,XY,-21,+22	46,XY,-21,+22
60 5CB	48,XY,+2,+3	48,XY,+2,+3	48,XY,+2,+3
61 4BB	40,X1,+3,-22	40,XY,+3,-22	n/a
62 4AA	48,XX,+5,+9	48,XX,+5,+9	48,XX,+5,+9
03 4AU	40,XX,+0,-10	40,XX,+0,-15	40,XX,+0,-15
65 3PP	40,X1,10,110	40,X1,10,110	40,X1,10,110,00(2)(021.1037.3)
66 3PP	48,XX, 13,110 48,XX +9 +22	48 XX +9 +22	48 XX +9 +22
67 4BB	48.XX.+10.+18	48.XX.+10.+18	48.XX.+10.+18
68 5AB	48.XY.+16.+21	48.XY.+16.+21	48.XY.+16.+21
69 5BB	46,XX,+18,-22	46,XX,+18,-22	46,XX,+18,-22
70 3BA	45,XX,-11,+12,-21	45,XX,-11,+12,-21	45,XX,-11,+12,-21
71 4CB	47,XY,+1,-13,+21	47,XY,+1,-13,+21	47,XY,+1,-13,+21
72 3BB	47,XY,+11,+18,-20	47,XY,+11,+18,-20	47,XY,+11,+18,-20
73 5BC	47,XY,+14,-15,+19	47,XY,+14,-15,+19	47,XY,+14,-15,+19
74 5BC	49,XX,+16,+18,+19	49,XX,+16,+18,+19	48,XX,+16,+18,+19*
75 4AB	48,XX,-2,+3,+9,+22	48,XX,-2,+3,+9,+22	48,XX,-2,+3,+9,+22
76 5BA	46,XY,+3,+4,-18,-21	46,XY,+3,+4,-18,-21	46,XY,+3,+4,-18,-21
77 5CB	47,XY,+16,del(20)(q13.2q13.33)	47,XY,+16,del(20)(q13.2q13.33)	47,XY,-1*,+16,-20*,del(5)(q23.1q35.3)*
78 4AB	46,XY,del(6)(q16.1q27)	46,XY,del(6)(q16.1q27)	46,XY,del(6)(q16.1q27)*
79 4BC	46,XX,del(1)(q43q44)	46,XX,del(1)(q43q44)	46,XX,del(1)(q43q44)

Blastocyst Study id# Gardner Grade	Clinical TE Biopsy	ICM Biopsy	Second TE Biopsy
80 4AC	45,XX,-14	45,XX,-14,del(1)(p36.32p36.12)*	45,XX,-14
81 4BB	45,XY,-16	45,XY,-16,dup(2)(p25.3p23.1)*	45,XY,-16
82 4AA	45,XY,-18	45,XY,-18,dup(2)(q23.3q37.3)*	45,XY,-18
83 5BA	45,XY,-21	45,XY,-21,dup(18)(p11.32q12.1)*	45,XY,-21
84 3BC	45,XX,-22	45,XX,-15*,-22	45,XX,-22
85 4BC	45,X	45,X,+20*,+21*	45,X,+2*,+14*
86 4BB	47,XX,+16	47,XX,+16,dup(5)(q12.1q35.1)*	47,XX,+16
87 5BB	47,XY,+20	47,XY,+20,del(1)(p36.33p33)*	47,XY,+20
88 5CC	46,XX,-10,+15	48,XX,-10*,+15,+17	46,XX,-10,+15
89 4BC	44,XX,-15,-22	44,XX, -15,+18*,-22,dup(1)(q12q44)*	44,XX,-15,-22
90 5BB	44,XY,-21,-22	44,XY,-21,-22,del(1)(p36.33p33)*	44,XY,-21,-22
91 5BC	47,XX,+4,+9,-13	47,XX,+3*,+4,+7*,+9,-13	47,XX,+3*,+4,+7*,+9,-13
92 5CC	47,XX,+20,del(2)(p25.3p24.1)	47,XX,+20,del(2)(p25.3p24.1)*	47,XX,+20,del(2)(p25.3p24.1)
93 5BB	46,XX,del(16)(p13.3p11.2)	45 XX,-16	46,XX,del(16)(p13.3p11.2),dup(16)(p11.2p24.3)

#### Clinical TE-ICM Aneuploid-Mosaic Discordan

	onnical re four Ancapiola mobale piecordant		
Blastocyst Study id#	Clinical TE Biopsy	ICM Biopsy	Second TE Biopsy
94 4BC	45,XY,-19	46,XY,-19*	45,XY,-19
95 5AB	47,XY,+6	46,XY,del(4)(q32.1q35.2)*	46,XY

#### Clinical TE-ICM Aneuploid-Euploid Discordant

Blastocyst Study id#	Clinical TE Biopsy ICM Biopsy		Second TE Biopsy	
96 4AB	47,XX,+12	46,XX	46,XX	
97 4BB	46,XX,del(2)(q22.1q31.1)	46,XX	46,XX,del(2)(q22.1q31.1)	
98 4BB	46,XY,dup(3)(q26.2q29)(x2)	46,XY	46,XY,dup(3)(q26.2q29)*,dup(22)(q11.1q13.31)*	
99 4AB	46,XX,del(9)(q12q34.3)	46,XX	46,XX,del(9)(q12q34.3)*	
100 3BB	46,XX,dup(11)(q23.2q25)	46,XX	n/a	

\* MOSAIC

Page 7 A of 87



## В



FIG2

#### All Blastocysts



Paired clinical TE-ICM Biopsy Status

- 79 Aneuploid-Aneuploid Perfect Concordant
- 14 Aneuploid-Aneuploid Imperfect Concordant -



2 Aneuploid-Mosaic Discordant

5 Aneuploid-Euploid Discordant

Total = 100



http://humrep.oupjournals.org

Dra**Clinical TEtBiopsiy**ted to Human Repro**dCivioBiopsy**er Review







**Biopsy Comparisons** 



Unrelated Blastocysts Controls
 Full-Sib Controls
 ICM/TE from Same Blastocysts

Draft Manuscript Submitted to Human Reproduction for Peer Review

	Nuclei (Hoechst)	Trophecto TE (GATA3)	derm Biopsy	Merge	Nuclei (Hoechst)	ICM	Biopsy	Merge
Embryo # 1	8	8/8	0/8 High	200	5	0/5	5/5 High	-
Embryo # 2	a	9/9	0/9 High	No.	9	0/9	9/9 High	
Embryo # 3	8	8/8	0/8 High	361	10	0/10	10/10 High	ŝ
Embryo # 4	s	9/9	0/9 High	<b>*</b>	7	0/7	7/7 High	50
Embryo # 5	8	8/8	0/8 High	800 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	5	0/5	5/5 High	\$
Embryo # 6	5	5/5	0/5 High	100	5	0/5	5/5 High	*
Embryo # 7	6	6/6	0/6 High	***** 8	a blo	0/7	7/7 High	Sec.
Embryo # 8	6	6/6	0/6 High	<b>E</b>	10	0/10	10/10 High	A.
Embryo # 9	8	8/8	0/8 High	a)	7	017	7/7 High	
Embryo # 10	8	8/8	0/8 High	60 60 60	9	0/9	9/9 High	Ć#
Embryo # 11		717	0/7 High	100	9	0/9	9/9 High	09

http://humrep.oupjournals.org





		400 3300 2280 2.40 2.40 2.40		# 72 Copy Number
		2200		# 71 Copy Number
	* · · · · · · · · · · · · · · · · · · ·	1.20 , , , , , , , , , , , , , , , , , , ,		# ~~
		0.80 0.40 0.40 2.2000 2.200 2.20000 2.200000000		t 70 py Number
		400 3380 2280 2280 2280 2280 2280 2280 22		# 69 Copy Number
		2.280 2.240 2.240 2.240 1.20 1.20 1.20 0.40		# 68 Copy Number
		1.20 0.40		<b>;</b>
		2200 × 100 ×		¢ 67
		2.80 2.40 2.40 2.40 1.50 1.50 1.50 1.50 0.80 0.40		# 66 Copy Number
		0.40 , , , , , , , , , , , , , , , , , , ,		<b>‡</b>
		400 0.40		65 Number
		3.80 3.20 2.80 2.40 2.40 2.40 2.20 2.00 4.80 4.80 4.80 4.80 4.80 4.80 4.80 4		# 64 Copy Number
		400 v		# 6; <sup>COPY NUI</sup>
		2.400 · 400		53 Iumber
		3 300 - 3 200 - 2 200 - 2 240		# 62 Copy Number
	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	400 0.80		# co
n/a		400	Number         No.         No.<	61 Jopy Number
		2.200 2.400 2.400 4.200 4.200 4.200 0.400		# 60 Copy Number
	や 0 v 0 0 小 0 0 0 小 0 0 0 0 0 0 0 0 0 0 0	320		
		3.80 3.20 2.40 2.40 2.40 2.40 1.20		# 59 Copy Number
n/a		400 0.40 0.40 0.40 0.40 0.40 0.40 0.40		# 5 <sub>Сору N</sub>
		2.40 0.40 2.20 7		58 Iumber
	Copy Number	3.20		# 57 Copy Number
	0         0	1400 0.400 7		# cor
		400 3.20 2.240 2.200		¥ 56 opy Number
the state of the s		2.80 2.40 2.40 1.400 1.400 1.400 1.400 0.40		# 55 Copy Number
	マ マ マ マ マ マ マ マ マ マ マ マ マ マ	1,20 0,40 7 3,20 1,20		
		400 320 220 220 220 220 400		# 54 Copy Number
		2200 - 22		# 53 Copy Number
		3380 0.40 0.80		
		4.00 3.20 2.20 2.20 2.20 2.20 2.20 2.20		# 52 Copy Number
		1120 0.40 -		# 5 <sub>Copy Nur</sub>
		2400		51 Number
		3.20 2.80 2.240 2.240 2.240 1.00 1.00 1.00 1.20 1.20 0.80 1.20 0.40		# 50 Copy Number
· · · · · · · · · · · · · · · · · · ·				
		2.00		# 49 Copy Number
		2200 - 22		#48 Copy Number
		1.20 		
		400 3.20 2.20 2.40 2.40 2.40 1.00		# 47
		2.40 2.20 1.20 0.40 0.40 7		# 4(
· · · · · · · · · · · · · · · · · · ·	* 0 × 0 0 人 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	200 040 0 200 7		
		3.80		# 45 Copy Number
		220 1120 0.40 7		# <b>4</b>
	ウ ウ ト ウ ウ ハ 中 ウ ク 小 ク ク へ ク ク ク ク ク ク ク ク ク ク ク ク ク ク ク	2280 7		er
		380		# 43 Copy Number
	1 0 1 0 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0	400 0.400		# 4 <sub>Copy</sub>
		2200 × 2200		42 <sup>py Number</sup>
		3.20 2.20 2.40 2.40 2.40 1.20 1.20 1.20 1.20 1.20		# 41 Copy Number
	マ ロ Y ロ ロ ハ ロ ロ 谷 小 谷 谷 谷 谷 久 谷 谷 谷 八 谷 谷 谷 ケ 谷 谷 八 谷 谷 谷 ケ 谷 谷 八 谷 谷 谷 ケ 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 八 谷 谷 り 谷 子 谷 谷 小 谷 谷 り 谷 日 日 日 日 日 日 日 日 日 日 日 日 日 日 日 日	400 0.80 1100 1 1 100 1 1 100	100 100 000 100 100 100 100 100 100 100	
		400 3.80 2.20 2.40 2.20		<b># 40</b>
		2.280		# 39 Copy Number
		400 3.20 2.20 2.20 2.20 2.20 2.20 2.20 2.		# 38 Copy Number
		2.80 2.40 2.40 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1		# 37 Copy Number
	<ul> <li>→ → → → → → → → → → → → → → → → → → →</li></ul>			# 
		22400 v0		36 y Number
		2.20 2.20 2.40 2.40 2.40 4.100 4.120 4.120 4.120		# 35 Copy Number
		400		# : Copy N
		0.40 2.200 2.2		34 umber
		3.20 3.20 2.20 2.40 2.40 1.20 1.20		# 33 Copy Number
		2200		# 3′ <sub>Copy</sub>
		3 200 0.40 2 200		2 Aumber
		3.80 3.20 2.40 2.240 1.20 1.20		# 31 Copy Number
		400 0.80 0.80		# 30 <sub>Copy</sub> ,
		3 3 3 2 0 0 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		łumber
		3.20 2.20 2.20 2.20 2.20 1.20 1.20 1.20 1		# 29 Copy Number
		0.40 		#
		0.40 3.80 2.400 2.200 2.200 2.200 2.200 2.200 2.200		28 <sup>yy</sup> Number
		3.20 2.80 2.40 2.40 1.20 1.20 1.20 1.20 1.20 1.20 1.20 1.2		# 27 Copy Number
	・ ・ ・ ・ ・ ・ ・ ・ ・ ・ ・ ・ ・ ・	400 		ca
		2200 × 100 ×		# 26
		4466 3380 2280 2.40 2.40 1.20 1.20 1.20		# 25 Copy Number
		2:40 2:00 1:20 0:40 7		# 2 Copy Ni
	2.0	28		<b>↓</b> øber

#8	#7	#6	#5	#4	#3	#2	#1	
Copy Number	Copy Number	Copy Number	Copy Number	Copy Number	Copy Number	Copy Number	Copy Number	
400 400 400 400 400 400 400 400							Clinical TE Biopsy	Draft Manuscript Su Clinical TE-I
							ICM Biopsy	bmitted to Human Reprodu CM Aneuploid-Aneuploi
	40 300 100 100 100 100 100 100 10	4 00 00 00000 + 1 4 3 3 0 2 3 0 2 3 0 1 4 1 3 0 1 4 1 5 1 5 1 5 1 5 1 5 1 5 1 5 1 5	40 300 200 100 100 000 000 000 000 0	400 330 2.20 2.20 1.20 0.00 1.20 0.00 1.20 0.00	400 300 100 100 100 100 100 100 1	400 300 200 100 100 100 100 100 100 1	Second	ction for Peer Review d Perfect Concordant



#23 #22



## Supplemental FIG3

Blastocyst Stage



Grade









## Supplemental FIG. 4

#### Supplemental Table I.

Blastocyst	Primary And Compositio	cestry n of	1000 Genomes Reference Panel Super-Population		
ID	K = 10 Near	est Neighbors			
	ICM	TE	-		
#94	C/S Asia	C/S Asia	SAS		
#95	C/S Asia	C/S Asia	SAS		
#96	East Asia	East Asia	EAS		
#97	Europe	Middle East	EUR		
#98	East Asia	East Asia	EAS		
#99	East Asia	East Asia	EAS		
#100	East Asia	East Asia	EAS		

#### Supplemental Data 1

#### Analysis of tissue relatedness

'Tilde' (Vohr et al., 2015) was used to rule out sample cross-contamination or mislabeling and infer, based on low-coverage sequencing data, whether ICM and TE biopsies were derived from the same blastocyst. This method facilitates indirect comparison of low-coverage samples based on the principle that sparse observed genotypes are informative of genotypes at nearby unobserved markers due to patterns of linkage disequilibrium (LD) in the population.

Reads were mapped to the hg19 reference using the BWA (version 0.7.17) backtrack algorithm with default parameters (Li and Durbin, 2009). We then used the LASER method (version 2.04; (Wang et al., 2015)) to select the appropriate ethnically matched 1000 Genomes Project super-population (Genomes Project et al., 2015) for each blastocyst, as required by tilde. LASER combines genotype imputation with principal components analysis to infer individual ancestry based on low-coverage sequence data. Blastocyst genotypes were visualized in reference ancestry space defined by principal components analysis of the HGDP reference panel (Supplemental Fig. 4; (Li et al., 2008)). Blastocysts were then assigned to corresponding 1000 Genomes super-populations based on ancestries of the K=10 nearest neighbor reference samples in principal components space (Supplemental Table I). In the case of

blastocyst #97, whose ICM and TE biopsies were assigned to European and Middle Eastern reference populations, respectively, we selected the European super-population as the reference panel. We note that these populations fall close to one another in space defined by the top three principal components. Furthermore, Vohr et al. (2015) demonstrated that tilde is relatively robust to misspecification of the reference panel.

Tilde computes a log-likelihood ratio comparing a model in which two samples are derived from the same individual (i.e., same embryo) to a model in which two samples are derived from unrelated individuals (i.e., different embryos). Positive log-likelihood ratios indicate that the data support the former model, while negative log-likelihood ratios indicate that the data support the latter model. Bootstrapping was performed to generate distributions and assess uncertainty in log-likelihood ratio estimates.

As controls, we applied this method to twenty-four comparisons of presumed unrelated embryos as well as four comparisons of full sibling (full-sib) embryos obtained from the same patient. Results from the unrelated negative controls supported the capacity of tilde to distinguish these samples, reflected by negative distributions of log-likelihood ratios (Fig. 4). For all seven embryos producing discordant TE-ICM results, the data supported a model in which the samples were derived from the same corresponding embryo, reflected by positive distributions of log-likelihood ratios. Meanwhile, the full-sib samples from the same patient also produced positive distributions of log-likelihood ratios, but intermediate between the unrelated and same-embryo comparisons, supporting

the power of tilde to distinguish varying levels of relatedness. Together, our data suggest no evidence of cross-contamination or sample mislabeling and substantiate the conclusion that the TE and ICM biopsies of discordant karyotype were derived from the same respective embryos. More generally, our analysis demonstrates that methods such as tilde, which take advantage of patterns of LD in the population, can be useful for research on low-coverage sequencing-based PGT-A datasets.

#### Supplemental Data 1 Bibliography

Genomes Project, C., Auton, A., Brooks, L.D., Durbin, R.M., Garrison, E.P.,

Kang, H.M., Korbel, J.O., Marchini, J.L., McCarthy, S., McVean, G.A., et al.

(2015). A global reference for human genetic variation. Nature 526, 68-74.

Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with

Burrows-Wheeler transform. Bioinformatics 25, 1754-1760.

Li, J.Z., Absher, D.M., Tang, H., Southwick, A.M., Casto, A.M., Ramachandran,

S., Cann, H.M., Barsh, G.S., Feldman, M., Cavalli-Sforza, L.L., et al. (2008).

Worldwide human relationships inferred from genome-wide patterns of variation. Science *319*, 1100-1104.

Vohr, S.H., Buen Abad Najar, C.F., Shapiro, B., and Green, R.E. (2015). A method for positive forensic identification of samples from extremely low-coverage sequence data. BMC Genomics *16*, 1034.

Wang, C., Zhan, X., Liang, L., Abecasis, G.R., and Lin, X. (2015). Improved ancestry estimation for both genotyping and sequencing data using projection procrustes analysis and genotype imputation. Am J Hum Genet *96*, 926-937.