

# Growth hormone during in vitro fertilization in older women modulates the density of receptors in granulosa cells, with improved pregnancy outcomes

Article

Accepted Version

Creative Commons: Attribution-Noncommercial-No Derivative Works 4.0

Regan, S. L. P., Knight, P. G. ORCID: https://orcid.org/0000-0003-0300-1554, Yovich, J. L., Arfuso, F. and Dharmarajan, A. (2018) Growth hormone during in vitro fertilization in older women modulates the density of receptors in granulosa cells, with improved pregnancy outcomes. Fertility and Sterility, 110 (7). pp. 1298-1310. ISSN 0015-0282 doi: 10.1016/j.fertnstert.2018.08.018 Available at https://centaur.reading.ac.uk/81147/

It is advisable to refer to the publisher's version if you intend to cite from the work. See <u>Guidance on citing</u>.

To link to this article DOI: http://dx.doi.org/10.1016/j.fertnstert.2018.08.018

Publisher: Elsevier

All outputs in CentAUR are protected by Intellectual Property Rights law, including copyright law. Copyright and IPR is retained by the creators or other copyright holders. Terms and conditions for use of this material are defined in



the End User Agreement.

# www.reading.ac.uk/centaur

## CentAUR

Central Archive at the University of Reading

Reading's research outputs online

1	Growth hormone during in vitro fertilization in older women modulates
2	the density of receptors in granulosa cells, with improved pregnancy
3	outcomes.
4	
5	
6 7 8 9	Sheena L.P. Regan <sup>a</sup> *, Phil G. Knight <sup>b</sup> , John L.Yovich <sup>c</sup> , 'Frank Arfuso <sup>a</sup> , Arun Dharmarajan <sup>a</sup> ,
10 11	<sup>a</sup> Stem Cell and Cancer Biology Laboratory, School of Biomedical Sciences, Curtin Health Innovation Research Institute, Curtin University, Perth, Australia. <sup>b</sup> School of Biological Sciences,
12 13 14 15 16 17	Hopkins Building, University of Reading, Whiteknights, Reading RG6 6UB, UK. <sup>c</sup> PIVET Medical Centre, Perth, Australia.
18	* Dr Sheena LP Regan,
19 20 21 22 23	<sup>a</sup> Stem Cell and Cancer Biology Laboratory, School of Biomedical Sciences, Curtin Health Innovation Research Institute, Curtin University (CHIRI), Perth, Australia, GPO Box U1987, Perth, WA 6845, Australia Email: <u>sheenaregan@aapt.net.au</u>
24	
25	
26	
27 28	
29	
30	
31	
32	
33	
34 35	
35 36	

### 37 Introduction

38 Ovarian depletion of primordial follicles is a continual natural process from gestation to adulthood,

39 which culminates in the loss of ovarian function and which eventuates in the state of menopause (1,

40 2). When activated, the primordial follicles grow and develop into small antral follicles, the majority

41 of which succumb to apoptosis (3, 4). At puberty, cyclic increases in circulating follicle stimulating

42 hormone (FSH) recruit a cohort of small antral follicles at the start of each menstrual cycle (3, 5).

43 The follicles grow under the influence of FSH, and express follicle stimulating hormone receptor

44 (FSHR) and luteinizing hormone receptor (LHR). The activation of FSH and the FSHRs stimulates

45 oestrogen synthesis, which subsequently stimulates proliferation of the granulosa cells and

46 development of the oocyte. (6-9).

47

The ovulation rate is determined by the stage-specific decrease in pituitary secreted FSH, and results in follicles with insufficient LHRs that succumb to apoptosis (10-12). The follicle continues to grow until pre-ovulatory maturation when proliferation ceases and granulosa cell differentiation occurs in preparation for ovulation of the oocyte.

52

As the ovarian reserve of primordial follicles is depleted over the reproductive lifespan, regulation of
 folliculogenesis is altered, which results in decreased fertility (13). Ovarian depletion can be

indirectly measured by the number of small antral follicles present at the beginning of a cycle, and is

56 highly correlated to chronological age (14). During IVF treatment, high doses of recombinant human

57 (r) FSH are administered to recruit more of the small antral follicles, and to maintain their growth

- 58 during pituitary FSH down-regulation (15).
- 59

60 Infertility patients with a poor ovarian reserve have fewer small antral follicles available for

61 recruitment, and higher doses of rFSH are used but with diminishing effectiveness in recruiting more

62 follicles during IVF cycles. In an attempt to improve the pregnancy rate, patients have been offered

63 co-treatment with growth hormone (GH) (15, 16). The patients with a poor response to rFSH

treatment represent a large group of patients with critically diminishing ovarian reserve (17, 18). The

65 challenge remains to identify the changes taking place as the ovarian reserve declines, and to find

66 alternative stimulation to provide high quality oocytes for fertilisation.

67

68 Earlier studies showed GH treatment *in vivo* and *in vitro*, in conjunction with rFSH increased oocyte

69 survival rate and pregnancy rate (19-22). The granulosa cells, including cumulus cells, as well as the

70 oocyte of antral follicles express growth hormone receptor (GHR) and are therefore able to react to

71 pituitary-derived or ovarian sources of GH (23, 24). With regard to the latter, granulosa cells and the

72 oocyte, but not cumulus and theca cells, have been shown to express GH mRNA (23-27). GHRs are

activated by GH, which changes the conformation of the receptor, promoting formation of a complex

- vith janus kinase (JAK)2 (28). The GHR-JAK2 complex can elicit numerous cellular responses in
- the body, such as cell differentiation and oocyte maturation in the ovary (29).
- 76

77 The cellular mechanism underpinning the GH-induced improvement in oocyte quality and reduced 78 miscarriage rate has not been reported in human studies. However, many attempts have been made to 79 delineate the indirect changes taking place to serum and follicular fluid hormone levels. Previously, 80 we have presented comprehensive results on the granulosal cell surface receptor density profiles of 81 patients during ovarian ageing (30, 31). Ovarian granulosa cell receptor expression was found to 82 fluctuate at the two critical times of dominant follicle selection and again at the terminal end of 83 folliculogenesis in preparation for ovulation. Lower levels of receptor density and a reversal of this 84 regulatory pattern was associated with reduced fertility and ovarian reserve in older patients. In the 85 present study, we report the granulosal GHR density in different sized follicles from IVF patients 86 undergoing conventional ovarian stimulation, with rFSH alone and with rFSH combined with GH co-87 treatment in young compared to older women with a reduced ovarian reserve. In addition, we report 88 the granulosal FSHR, LHR, and BMPR1B receptor density in older, poor ovarian reserve patients 89 treated with GH.

### 90 Materials and Methods

### 91 **Patients**

Patients (women) were selected randomly in a prospective regimen, and aged between 23 and 45 years, with a range of infertility factors, but limited to exclude unusual medical conditions, endocrine dysfunction, polycystic ovarian syndrome and endometriosis. Infertility issues were comprised of male factor, low ovarian reserve, donor sperm or unexplained fertility; and fertilisation was via intracytoplasmic sperm injection (ICSI). A total of 483 follicles were collected from 64 patients undergoing standard fertility treatment at PIVET Medical Centre Perth, Western Australia, (Table 1).

### 99 Human IVF: Ovarian stimulation, follicular fluid, and oocyte

100 Patient treatment consisted of two types of gonadotrophin releasing hormone-LH suppression

101 (Orgalutran; MSD and Cetrotide; Merck Serono) in conjunction with commercially prepared

102 recombinant human (r) FSH (Puregon; MSD and Gonal-f; Merck Serono), from cycle day 2 for ~10

103 days, as described by Regan et al. (2015). Ovulation was triggered with 10 000 IU human chorionic

104 gonadotrophin (hCG: Pregnyl; MSD), and oocyte retrieval was 36 hours later by transvaginal oocyte

aspiration (30). Patients classified as poor prognosis due to poor ovarian response or with three or

- 106 more failed attempts to conceive through IVF treatment with gonadotrophin alone, were co-treated
- 107 with a total of 60 IU GH (Saizen, Serono, Australia) over a period of 20-24 days in the lead-up to
- 108 IVF. GH was administered to 11 patients starting on day 21 of the previous cycle, and on day 2, 6, 8,

- 109 10, and 12 (10 IU per injection, a total of 60 IU) of the current cycle to women aged  $\geq$ 39 years who
- 110 had at least one failed IVF cycle (15).
- 111

### 112 Antral follicle count

113 Patients received daily rFSH according to a long established algorithm based on the patient's profile

114 of age and ovarian reserve in order to determine the rFSH dose required to stimulate 8-12

- 115 preovulatory follicles, (32). Ovarian reserve was measured indirectly by the antral follicle count, and
- 116 was defined as the number of follicles between 2-10 mm in diameter, combining the number
- 117 collected from both ovaries; that were present on ~day 5 of a preliminary assessment cycle, without
- 118 rFSH (14). The patients were divided by age and ovarian reserve into groups based on the algorithm,
- as described previously by Regan et al. (2016) and a well-established clinical practice of patient
- 120 treatment (32, 33): Group A+ = 30-39 small follicles; group A = 20-29 small follicles; group B = 13-
- 121 19 small follicles; group C = 9-12 small follicles, group D = 5-8 small follicles; group E =  $\leq$  4 small
- 122 follicles.

### 123 Immunolabelling of granulosa cells

- 124 The ovarian follicles studied ranged in diameter from 4 to 27 mm, and an average of ~8000
- 125 granulosa cells per individual follicle were analysed. Cell surface-expressed mature GHR protein
- density was measured by immunofluorescent labelling and flow cytometry. The diameter of the
- 127 follicle was calculated using ultrasonography as described previously (30, 31, 34). Flushing of the
- 128 follicle (Quinn's Advantage with Hepes, Sage Media, Pasadena, California) removed the loosely
- 129 attached layers of granulosa cells. Aliquots of suspended granulosa cells  $(1 \times 10^6 \text{ cells in } 100 \,\mu\text{l})$  were
- immunolabelled and incubated separately with an optimised concentration of 4 µg/ml affinity
- 131 purified polyclonal antibody to bone morphogenetic hormone receptor (BMPR1B), FSHR, LHR or
- 132 GHR for 25 min at 5 °C.
- 133 3D image analysis using immunofluorescence detection has established the specificity of the
- 134 antibodies in sheep, polyclonal goat anti-BMPR1B (sc-5679), goat anti-FSHR (sc-7798), and goat
- 135 anti-LHR (sc-26341) (Santa Cruz Biotechnology, Santa Cruz, CA, USA), (35); and GHR (AF1210;
- 136 Life Technologies, Victoria, Australia) (36). In addition, use of these antibodies has been previously
- 137 reported in human studies (37-44) and for use in flow cytometry (38). The cells were washed with
- 138 PBS and centrifuged at 300 g at 5°C for 5 min. To render a homogeneous population of granulosa
- 139 cells the monoclonal antibody CD45 was added to BMPR1B, GHR, and LHR tubes to enable the
- subtraction of the positive leukocyte common antigen ( $\sim 3\%$ ) not removed during isolation of the
- 141 granulosa cells with the ficoll gradient (555485; BD Biosciences, Perth, Australia), (Fig. 1A and 1B).

- 143 Unstained samples or the substitution of a primary antibody with pre-immune goat IgG (Millennium
- 144 Science, Surrey Hills, Victoria Australia) at the same concentration as the primary antibody served as
- 145 a negative control for auto-fluorescence (Fig. 1A). A blocking peptide for FSH receptor and bone
- 146 morphogenetic protein receptor 1B indicated nonspecific binding applied to human granulosa cells
- 147 (sc-7798P, sc-5679P; Millennium Science, Surrey Hills, Victoria Australia), (Fig. 1B), and as
- 148 previously published (37, 38). Pre-absorbed LH (Lutropin, Merck Serono, Frenchs Forest, NSW,
- Australia), and GH (Saizen, Merck Serono, Australia) also confirmed binding specificity. In the
- 150 current study, the 'normal' goat IgG and unstained control cells emitted an average mean fluorescent
- 151 intensity (MFI) that was classified as non-specific auto-fluorescence. The auto-fluorescence and the
- 152 nonspecific binding determined by the unstained control for each follicle was subtracted from each
- 153 follicle (Fig. 1B), and as described previously (30, 31).
- 154 Re-suspended 10µl aliquots of GHR immunolabelled, live granulosa cells were placed on slides and
- 155 visualized using an Olympus DP 70 camera fitted to an Olympus BX-51 upright fluorescent
- 156 microscope with a 40x UPlan N 0.4 N.A. objective (Olympus Imaging Australia, Macquarie Park,
- 157 Australia), (Fig.1C). Fluorescent microscopy revealed a positive staining of the cell membrane-
- bound GHR as an intermittent, bright, ring-like pattern around the cells (Fig .1C). Pre-absorbed GH
- 159 was used as a negative control. A pure granulosa cell population was identified by graphing forward
- 160 scatter to remove doublets (FSC-H verses FSC-A), as previously described (30, 31, 34). The uniform
- 161 granulosa cell population revealed positive staining for FSHR, which is unique to granulosa cells
- 162 (45). The data were analysed using FlowJo software (Tree Star Inc., Oregon, USA).
- 163

### 164 Serum and Follicular fluid assessment

165 The peak oestrogen concentration in serum was used to predict the follicular health of the follicle as

- 166 opposed to the serum levels collected at the time of follicle aspiration. Serum was analysed using
- 167 biochemical analysis on the days leading up to collection and on the day of collection. IVF patients
- 168 undergoing treatment were examined in a natural cycle and during exogenous rFSH stimulated
- 169 cycles. Follicular fluid collected from follicles 17 to 23 mm were analysed for testosterone, FSH, and
- 170 LH using a random access immunoassay system (Siemens Medical Solutions, Bayswater, Victoria,
- 171 Australia). Follicular fluid, testosterone, FSH, and LH were analysed undiluted, whereas oestrogen
- and progesterone were diluted manually 1:1000 with a multi-diluent and, when required, a further
- 173 manual dilution of progesterone 10 x and oestrogen; 5 x. Percentage coefficient of variance (CV) for
- a concentration range 137.4 pmol/L to 3257 pmol/L was oestrogen = 5.2; LH = 3.9; FSH = 2.9;
- 175 testosterone = 5.9; progesterone = 9.4.
- 176

### 177 Statistics

- 178 Mean fluorescent intensity was obtained using ~8000 granulosa cells per individual follicle for the
- 179 direct measurement of receptor protein expression. The data were subjected to statistical verification

- 180 using one-way ANOVA with an uncorrected Fisher's LSD for follicular size using GraphPad Prism
- 181 6. Values in graphs are means  $\pm$  S.E.M., and differences were considered significant if \*p<0.05,
- 182 \*\*p<0.01, \*\*\*p<0.005, and \*\*\*\*p<0.001. The letter, such as 'a', signifies a statistical significant
- 183 difference to the matching letter (e.g. 'a\*'). The attached asterisk (a\*) indicates the significance level
- 184 for the size follicle. A two tailed, student t-test and chi squared was also used.

### 185 Human Ethics

- 186 Patients undergoing standard fertility treatment at PIVET Medical Centre, Perth, Australia provided
- 187 informed consent according to Curtin University Human Research Committee (HR RD26-10:2010
- and 2016); and all methods were performed in accordance with the relevant guidelines and
- 189 regulations under State Legislation and National Accreditation processes.
- 190

### 191 **Results**

### 192 GHR density without growth hormone co-treatment and ovarian reserve depletion

- 193 In the youngest patients with good ovarian reserve, a constant level of granulosal GHR was
- 194 expressed during follicular growth in both the A+ and A groups, both of which are typical for a
- 195 patient in this age group (Fig. 2A).
- 196

197 GHR density was significantly decreased as the ovarian reserve was depleted in all of the three older

- age groups. In the 31-34 y patient group, GHR density on the granulosa cells from follicles of the
- same size was significantly reduced in the patients with a reduced ovarian reserve for the age group
- 200 (p=0.039, 14 mm follicles; 0.0037, 16 mm follicles, Fig. 2B). This trend was also found in the 35-38
- 201 y patient group; (p=0.029, 4 mm follicles;, Fig. 2C) and in the 39+ y patient group (p=0.0001, 4 mm
- follicles; p=0.0012 14 mm follicles, Fig. 2D). In the older patients (39+ y), with a comparatively
- 203 better ovarian reserve of B or C, the level of GHR was significantly reduced in the larger follicles to
- the level observed in the poorer D and E ovarian reserve group (p<0.001; Fig. 2D).
- 205

### 206 GHR receptor density profile independent of patient age

The patient data were analysed based on ovarian follicle reserve, independent of chronological age (Suplemetary Fig. 1). In patients with good ovarian reserve, an initial high level of GHR in the smaller follicles was followed by a decline as the follicles increased in size (14 to 23 mm follicles, p=0.0005). This pattern was reversed in the poorer ovarian reserve patient groups of D & E (p<0.05). Granulosal GHR receptor density was greater in the 10 mm (p<0.01) and 14 mm (p<0.005) follicles in the good ovarian reserve patient group compared to patients with the poorest ovarian reserve (Suplemetary Fig. 1).

214

### 215 Growth hormone co-treatment restores preovulatory down-regulation of FSHR

216 **BMPR1B and LHR** 

- 217 The level of GHR was significantly increased in IVF patients receiving GH co-treatment in follicles
- from 10 to 23 mm compared to the same age patients of 39+ y with an ovarian reserve of D & E
- 219 (p<0.01 to p<0.001, Fig.3A). The level of GHR expression in different sized follicles was not
- significantly different in patients treated with GH (Fig 4A).
- 221

222 The level of FSHR was significantly increased in IVF patients receiving GH in 16 mm follicles

compared to the same age patients of 39+y with an ovarian reserve of D & E without GH (p<0.001,

Fig.4B). The level of FSHR in GH treated patients was also increased in the larger follicles from 4

225 mm to 16 mm (Fig. 3B, p<0.005). This was followed by a significant down-regulation of the largest

226 preovulatory follicles (p<0.01, 19 mm).

227

- 228 The level of LHR was significantly increased in IVF patients receiving GH in 16 mm follicles
- compared to the same age patients of 39+y with an ovarian reserve of D & E without GH (p<0.005,

Fig.4C). The LHR density of the granulosa cells collected from patients who received GH co-

treatment during an IVF cycle was also significantly elevated in the 10 to 16 mm follicles (p<0.01,

Fig. 3C). In contrast to the untreated group, GH co-treated patients showed down-regulation of

- granulosal LHR density in follicles between 16 and 19 mm in diameter (p<0.005, Fig. 3C).
- 234

The level of BMPR1B was significantly increased in IVF patients receiving GH in 10 mm, 14 mm

and 16 mm follicles compared to the same age patients of 39+ y with an ovarian reserve of D & E

237 without GH (p<0.001, p<0.005, p<0.05, respectively; Fig.4D).

238 Granulosal BMPR1B density was significantly higher in 10 mm, follicles from the GH co-treated

patients compared to the larger pre-ovulatory follicles of either 16 mm or 19 mm (p < 0.05, Fig. 3D).

- 240 In contrast, to the untreated group, GH co-treated patients showed down-regulation of granulosal
- BMPR1B density in the largest follicles of 16 to 19 mm (p<0.05, p<0.05, respectively; Fig. 3D)
- 242

When the follicles sizes are combined, the average granulosal density for GHR, FSHR, LHR and BMPR1B was significantly higher in the GH treated group with the same ovarian reserve and age

- 245 (Fig. 3 A-D Inset, p<0.005).
- 246

### 247 Growth hormone co-treatment and pregnancy rate in IVF patients

The number of pregnancies was calculated based on the number of embryos that were transferred to the patients, which included subsequent FET cycles of cryo preserved embryos.

- 250 The number of FET cycles was not significantly different between groups of patient. There was a
- significant difference in the pregnancy rate in GH treated patients compared to the same age and
- ovarian reserve patients without GH co-treatment (p=0.003; Fig. 5.). The number of live births per

- 253 embryo transfer was also significantly greater in the GH co-treated older age group compared to the 254 equivalent age and ovarian reserve (p=0.0406, Fig. 6). The level of oestrogen and progesterone in
- 255
- serum and follicular fluid was not significantly different when comparing GH treatment in the
- 256 equivalent older patient group of 39+ y.
- 257

#### 258 Serum & Follicular Fluid & GH co-treatment

259 The results from the current study indicate that the GH co-treatment did not alter the oestrogen level 260 of the 39+ year group cohort with an ovarian reserve of D or E during an IVF cycle (Supplementary 261 Fig 2A). Furthermore, neither the ratio of oestrogen was not altered, nor the levels of oestrogen 262 secreted, based on either the total number of follicles or the number of follicles greater than 14 mm 263 present in the ovary at the time of collection, which were not significantly different. In addition, the 264 follicular fluid concentration of oestrogen, progesterone, FSH, or testosterone was not significantly 265 different to the age matched patients with a similar ovarian reserve that were co-treated with GH 266 (Supplementary Fig 2B).

#### Discussion 267

- 268 GHRs are predominantly found on the granulosa cell membrane surface and in the endoplasmic
- 269 reticulum, and to a lesser degree, but commonly, in the nuclear membrane of highly prolific cells (46,
- 270 47). The GHR is regulated by GH binding proteins, which are secreted from the GHR, and by other
- 271 growth factors indirectly such as FSH, BMPs and somatostatin (48, 49). In the current study, GH
- 272 treatment induced a direct change to the receptor expression of GHR itself and indirectly to the other
- 273 receptors FSHR, LHR and BMPR1B.
- 274
- 275 In support of the clinical data on ageing, human granulosa receptor density and dysregulation of 276 FSHR, LHR and BMPR1B has been associated with ovarian depletion and reduced fertility (30, 31). 277 We now provide additional data in support of a reversal of the dysregulated receptor expression 278 observed in older patients that occurs when they are treated with GH. In addition, depletion of the 279 ovarian reserve was accompanied by a reduction in GHR density, whereas GH co-treatment during 280 IVF increased the receptor density in older women who had a reduced ovarian reserve. These 281 findings provide a possible cellular regulatory mechanism involved in the poor pregnancy and live 282 birth rate in the older 39-45 y patients and reported by others (15, 16, 50-55) and reviewed by (56-283 58).
- 284

285 Evidence from our previously published work and the current study suggest that ovarian reserve and

286 age are associated with reduced and dysregulated levels of receptor expression on granulosa cells.

- 287 Therefore, the influence of age and ovarian reserve of subjects or animals needs to be considered as a
- 288 confounding variable in previous studies. In heifers, GH may not have resulted in any change to
- 289 FSH and LH receptor binding because the cows were young, with an uncompromised ovarian reserve

and a sufficient receptor density (59). The effect of GH co-treatment on receptor density in patients

291 with a good ovarian reserve for age remains at this time unknown.

292

293 While GH increased the receptor expression on granulosa cells from the larger follicles, it had no 294 effect on the FSHR and LHR density at the critical time of dominant follicle selection (smallest 295 follicles of 4 mm). Previously, a poor ovarian response to rFSH stimulation has been associated with 296 reduced granulosal FSHR expression (37). However, GH co-treatment was found not to alter the 297 FSHR density in small bovine follicles, which is consistent with our findings for small human 298 follicles (60). The lack of effect on FSHR and LHR expression of small pre-ovulatory follicles may 299 explain why the number of oocytes collected was not increased in the current human model and 300 others (56).

301

302 Conversely, animal studies have reported an increase in oocyte number (61-65). For example, even 303 though more small bovine antral follicles were produced after 45 days of GH treatment in a natural 304 cycle, the granulosal FSHRs and LHRs from pre-ovulatory bovine follicles were not affected (59, 305 60). This is surprising; however, the receptor binding studies were determined only for the three 306 largest follicles from each cow. Therefore, the expected pre-ovulatory down-regulation of these large 307 follicles would have reduced FSHR expression which would confound these results. Added to this 308 the receptor binding was not measured in any of the smaller follicles. In other studies, GH treatment 309 increased the receptors in the rat (66) however; in the pig the receptor expression was reduced (67). 310 In our human model, small antral follicles had high levels of FSHR followed by down-regulation 311 which coincides with dominant follicle selection. The high level of FSHRs induce LHR expression in 312 a natural cycle to ensure recruitment to the dominant cohort of follicles (68). In a natural cycle, 313 pituitary secreted FSH is reduced at this critical time, whereas in an IVF cycle; rFSH is abundant; 314 therefore the densities of the gonadotrophin receptors (FSHR and LHR) are pivotal in regulating 315 follicle growth and dominance.

316

Patients with a reduced ovarian reserve have a poorer response to rFSH treatment in IVF, and produce oocytes of poorer quality (37). The poor responder group of patients also have an associated high risk of foetal aneuploidy that has been correlated to ovarian reserve (69). Recently published data have shown that GH co-treatment increases the pregnancy rate by a suggested improvement in oocyte quality, rather than the quantity of follicles recruited (15, 16).

322

323 If the oocyte number is not significantly different in the GH treated older women, then the focus

324 shifts to the effect of GH on the quality of the oocyte. Regulation of proliferation, steroid production,

325 luteinisation, ovulation, and recommencement of meiosis fundamentally resides with the functional

- 326 expression of receptors in the follicle cells and oocyte.
- 327

- 328 A decline in granulosal BMPR1B and FSHR density occurred at the time of cyclic dominant follicle
- 329 selection, and again during the terminal stage of folliculogenesis, in young (23-30 years) IVF patients
- 330 with good ovarian reserve (30, 31). Older patients (39+ years) with poor ovarian reserve experienced
- a reversal of this pattern (30, 31). In addition, the LHR density failed to be down-regulated during
- 332 pre-ovulatory maturation in the 39+ year group, and was reduced with ovarian reserve (31).
- 333

In the present study, we report increased granulosa cell GHR density in different sized follicles from

- 335 IVF patients undergoing conventional ovarian stimulation in young compared to older women with a
- reduced ovarian reserve. In addition, we report increased granulosal GHR, FSHR, LHR, and
- BMPR1B receptor density in older, poor ovarian reserve patients treated with GH. Importantly, the
- 338 women treated with GH demonstrated receptor expression down-regulation in the largest follicles.
- 339 The down-regulation would be essential for maturation of the ovulatory follicles, luteinisation and a
- 340 shift to the luteal phase.
- 341

342 In addition, the increased granulosal LHR density observed in the GH co-treated patients would have

343 the potential to increase the sensitivity during the hCG/LH surge to trigger final maturation and

ovulation of the oocyte (70, 71). The improved sensitivity may give rise to improved oocyte quality

345 and live birth rate. In support of the link between receptor density and maturation, a previous

346 electron microscopy study revealed that oocytes that did not fertilise had reduced levels of granulosal

347 luteinisation and were less responsive to hCG, which binds to the LHR (72).

348

### 349 **Conclusion**

- The complexity and limitations of a largely observational, *in vivo* study in humans makes it difficult to define the cellular mechanism through which numerous growth factors and pathways contribute to the regulation of follicular growth and differentiation. However, the present study has generated evidence suggesting several cellular mechanisms that could contribute to the improved oocyte quality observed in GH co-treated IVF patients with a poor ovarian reserve.
- 355

356 GH co-treatment increased granulosal GHR density that would increase GHR-JAK-STAT activity,

and result in an increase in the intermediate products of transcription. This, in turn, could be

358 mechanistically linked to the corresponding increase in gonadotrophin receptors and BMPR1B

density observed in GH co-treated patients. GH co-treatment did not alter the gonadotrophin receptor

360 density of the small follicles, and would therefore account for the lack of improvement in the number

361 of follicles recruited during dominant follicle selection.

362

363 In contrast, GH co-treatment also restored the pre-ovulatory down-regulation of FSHR, BMPR1B

and LHR density, which may improve the maturation process of luteinisation in GH co-treated

365 patients with reduced ovarian reserve. Combined with the latter, an increase in LHR density may

- 366 improve follicle development and provide another possible cellular mechanism responsible for the
- 367 improved pregnancy and live birth rate. Objectively, we remain uncertain whether the beneficial
- action is mediated via improved oocyte quality or other responses such as endometrial receptivity.
- 369

### 370 Acknowledgements

The authors thank all the participants who generously donated their samples to this study, the clinicaldoctors, embryologists, and nursing staff.

### 373 Authors' roles

- 374 SLPR conceived the study, experimental design, conducted all experiments, the analysis and
- interpretation of data, wrote the first draft of the manuscript and the final version of the paper.
- 376 Obtained informed consent from patients and ethics approval. PK supervised, interpretation of data,
- 377 contributed to the draft of the manuscript, interpretation of data, and critically revised the manuscript.
- 378 JLY supervised, participated in the study design, participated in obtaining granulosa cells,
- interpretation of data, and critically revised the manuscript. FA supervised, contributed to the draft of
- the manuscript, interpretation of data, and critically revised the manuscript. AD supervised,
- 381 participated in the study design, interpretation of data, contributed to the draft of the manuscript, and
- 382 critically revised the manuscript.
- 383

### 384 Funding

- 385 S.L.P.R. was a recipient of an Australian Postgraduate Award. This work was supported by
- additional private external funding, which was gratefully accepted from Denby Macgregor.

### 387 Conflict of interest

- 388 The authors declare that there is no conflict of interest that could be perceived as prejudicing the
- impartiality of the research reported.
- 390

### 391 References

- Gougeon A. Dynamics of follicular growth in the human: a model from preliminary results.
   Hum Reprod 1986;1:81-7.
- Richards JS. Hormonal Control of Gene Expression in the OVary. Endocrine Reviews
   1994;15:725-51.
- 396 3. Ginther OJ, Beg MA, Gastal EL, Gastal MO, Baerwald AR, Pierson RA. Systemic
- concentrations of hormones during the development of follicular waves in mares and women: acomparative study. Reproduction 2005;130:379-88.
- Hillier SG. Gonadotropic control of ovarian follicular growth and development. Molecular
   and Cellular Endocrinology 2001;179:39-46.
- 401 5. Austin EJ, Mihm M, Evans ACO, Knight PG, Ireland JLH, Ireland JJ *et al.* Alterations in
  402 Intrafollicular Regulatory Factors and Apoptosis During Selection of Follicles in the First Follicular
  403 Wave of the Bovine Estrous Cycle. Biology of Reproduction 2001;64:839-48.
- Campbell BK, Dobson H, Baird DT, Scaramuzzi RJ. Examination of the relative role of FSH
  and LH in the mechanism of ovulatory follicle selection in sheep. Journal of reproduction and
  fertility 1999;117:355-67.
- 407 7. Ginther OJ, Khan FA, Hannan MA, Rodriguez MB, Pugliesi G, Beg MA. Role of LH in
- 408 luteolysis and growth of the ovulatory follicle and estradiol regulation of LH secretion in heifers.
  409 Theriogenology 2012;77:1442-52.
- 410 8. Luo W, Gumen A, Haughian J, Wiltbank M. The role of luteinizing hormone in regulating
- 411 gene expression during selection of a dominant follicle in cattle. Biology of Reproduction
- 412 2011;84:369-78.

- 413 9. Picton HM, McNeilly AS. Evidence to support a follicle-stimulating hormone threshold 414 theory for follicle selection in ewes chronically treated with gonadotrophin-releasing hormone 415 agonist. Journal of reproduction and fertility 1991;93:43-51. 416 10. Minegishi T, Tano M, Abe Y, Nakamura K, Ibuki Y, Miyamoto K. Expression of luteinizing 417 hormone/human chorionic gonadotrophin (LH/HCG) receptor mRNA in the human ovary. Molecular 418 Human Reproduction 1997;3:101-7. 419 11. Mihm M, Baker PJ, Ireland JLH, Smith GW, Coussens PM, Evans ACO et al. Molecular 420 Evidence That Growth of Dominant Follicles Involves a Reduction in Follicle-Stimulating Hormone 421 Dependence and an Increase in Luteinizing Hormone Dependence in Cattle. Biology of Reproduction 422 2006;74:1051-9. 423 Yung Y, Aviel-Ronen S, Maman E, Rubinstein N, Avivi C, Orvieto R et al. Localization of 12 424 luteinizing hormone receptor protein in the human ovary. Molecular Human Reproduction 425 2014;20:844-9. 426 Klein NA, Battaglia DE, Fujimoto VY, Davis GS, Bremner WJ, Soules MR. Reproductive 13. 427 aging: accelerated ovarian follicular development associated with a monotropic follicle-stimulating 428 hormone rise in normal older women. The Journal of Clinical Endocrinology & Metabolism 429 1996;81:1038-45. 430 Hansen KR, Hodnett GM, Knowlton N, Craig LB. Correlation of ovarian reserve tests with 14. 431 histologically determined primordial follicle number. Fertility and Sterility 2011;95:170-5. 432 Yovich JL, Stanger JD. Growth hormone supplementation improves implantation and 15. 433 pregnancy productivity rates for poor-prognosis patients undertaking IVF. Reproductive 434 BioMedicine Online 2010;21:37-49. 435 Tesarik J, Hazout A, Mendoza C. Improvement of delivery and live birth rates after ICSI in 16. 436 women aged >40 years by ovarian co-stimulation with growth hormone. Human Reproduction 437 2005;20:2536-41. 438 de Ziegler D, Streuli I, Meldrum D, Chapron C. The value of growth hormone supplements 17. 439 in ART for poor ovarian responders. Fertility and Sterility 2011;96:1069-76. 440 18. Kyrou D, Kolibianakis E, Venetis C, Papanikolaou E, Bontis J, Tarlatzis B. How to improve 441 the probability of pregnancy in poor responders undergoing in vitro fertilization: a systematic review 442 and meta-analysis. Fertility and Sterility 2009;91:749-66. 443 Folch J, Ramon JP, Cocero MJ, Alabart JL, Beckers JF. Exogenous growth hormone 19. 444 improves the number of transferable embryos in superovulated ewes. Theriogenology 2001;55:1777-445 85. 446 Barreca A, Artini PG, Del Monte P, Ponzani P, Pasquini P, Cariola G et al. In vivo and in 20. 447 vitro effect of growth hormone on estradiol secretion by human granulosa cells. Journal of Clinical 448 Endocrinology & Metabolism 1993;77:61-7. 449 Izadyar F, Colenbrander B, Bevers MM. In vitro maturation of bovine oocytes in the 21. 450 presence of growth hormone accelerates nuclear maturation and promotes subsequent embryonic 451 development. Molecular reproduction and development 1996;45:372-7. 452 Izadyar F, Zeinstra E, Bevers MM. Follicle-stimulating hormone and growth hormone act 22. 453 differently on nuclear maturation while both enhance developmental competence of in vitro matured 454 bovine oocytes. Molecular reproduction and development 1998;51:339-45. 455 Izadyar F, Zhao J, Van Tol HT, Colenbrander B, Bevers MM. Messenger RNA expression 23 456 and protein localization of growth hormone in bovine ovarian tissue and in cumulus oocyte 457 complexes (COCs) during in vitro maturation. Molecular reproduction and development 1999;53:398-406. 458 459 24. Abir R, Garor R, Felz C, Nitke S, Krissi H, Fisch B. Growth hormone and its receptor in 460 human ovaries from fetuses and adults. Fertility and Sterility 2008;90:1333-9. 461 Izadyar F, Hage WJ, Colenbrander B, Bevers MM. The promotory effect of growth hormone 25. 462 on the developmental competence of in vitro matured bovine oocytes is due to improved cytoplasmic 463 maturation. Molecular reproduction and development 1998;49:444-53. 464 Izadyar F, Van Tol HT, Colenbrander B, Bevers MM. Stimulatory effect of growth hormone 26. on in vitro maturation of bovine oocytes is exerted through cumulus cells and not mediated by IGF-I. 465 466 Molecular reproduction and development 1997;47:175-80. 467 27. Bevers MM, Izadyar F. Role of growth hormone and growth hormone receptor in oocyte
- 468 maturation. Molecular and Cellular Endocrinology 2002;197:173-8.

469 28. Lan H, Li W, Fu Z, Yang Y, Wu T, Liu Y et al. Differential intracellular signalling 470 properties of the growth hormone receptor induced by the activation of an anti-GHR antibody. 471 Molecular and Cellular Endocrinology 2014;390:54-64. Waters MJ, Hoang HN, Fairlie DP, Pelekanos RA, Brown RJ. New insights into growth 472 29. 473 hormone action. Journal of molecular endocrinology 2006;36:1-7. Regan SLP, Knight PG, Yovich J, Stanger J, Leung Y, Arfuso F et al. Dysregulation of 474 30. 475 granulosal bone morphogenetic protein receptor 1B density is associated with reduced ovarian reserve and the age-related decline in human fertility. Molecular and Cellular Endocrinology 476 477 2016;425:84-93. 478 Regan SLP, Knight PG, Yovich JL, Stanger JD, Leung Y, Arfuso F et al. Infertility and 31. 479 ovarian follicle reserve depletion are associated with dysregulation of the FSH and LH receptor 480 density in human antral follicles. Molecular and Cellular Endocrinology 2017;446:40-51. 481 Yovich J, Stanger J, Hinchliffe P. Targeted gonadotrophin stimulation using the PIVET 32. 482 algorithm markedly reduces the risk of OHSS. Reproductive BioMedicine Online 2012;24:281-92. 483 33. Yovich JL, Alsbjerg B, Conceicao JL, Hinchliffe PM, Keane KN. PIVET rFSH dosing 484 algorithms for individualized controlled ovarian stimulation enables optimized pregnancy 485 productivity rates and avoidance of ovarian hyperstimulation syndrome. Drug Design, Development 486 and Therapy 2016;10:2561-73. Regan SLP, Knight PG, Yovich JL, Stanger JD, Leung Y, Arfuso F et al. The effect of 487 34. 488 ovarian reserve and receptor signalling on granulosa cell apoptosis during human follicle 489 development. Molecular and Cellular Endocrinology 2017. 490 Al-Samerria S, Almahbobi G. Three-dimensional image analysis to quantify the temproro-35. 491 smacial expression of cellular receptors. Journal of Medical and Bioengineering 2014 3:179-82. 492 Weall BM, Al-Samerria S, Conceicao J, Yovich JL, Almahbobi G. A direct action for GH in 36. 493 improvement of oocyte quality in poor-responder patients. Reproduction 2015;149:147-54. 494 Cai J, Lou H, Dong M, Lu X, Zhu Y, Gao H et al. Poor ovarian response to gonadotropin 37. 495 stimulation is associated with low expression of follicle-stimulating hormone receptor in granulosa 496 cells. Fertility and Sterility 2007;87:1350-6. Gao S, De Geyter C, Kossowska K, Zhang H. FSH stimulates the expression of the 497 38. 498 ADAMTS-16 protease in mature human ovarian follicles. Molecular Human Reproduction 499 2007;13:465-71. 500 Pidoux G, Gerbaud P, Tsatsaris V, Marpeau O, Ferreira F, Meduri G et al. Biochemical 39. 501 characterization and modulation of LH/CG—receptor during human trophoblast differentiation. 502 Journal of cellular physiology 2007;212:26-35. 503 Abir R, Ben-Haroush A, Melamed N, Felz C, Krissi H, Fisch B. Expression of bone 40. morphogenetic proteins 4 and 7 and their receptors IA, IB, and II in human ovaries from fetuses and 504 505 adults. Fertility and Sterility 2008;89:1430-40. 506 Haÿ E, Lemonnier J, Fromigué O, Guénou H, Marie PJ. Bone morphogenetic protein 41. 507 receptor ib signaling mediates apoptosis independently of differentiation in osteoblastic cells. Journal 508 of Biological Chemistry 2004;279:1650-8. 509 42. Bozzola M, Zecca M, Locatelli F, Radetti G, Pagani S, Autelli M, Tatò L, Chatelain P. 510 Evaluation of growth hormone bioactivity using the Nb2 cell bioassay in children with growth 511 disorders. J Endocrinol Invest 1998 Dec;21(11):765-70. Weall BM, Al-Samerria S, Conceicao J, Yovich JL, Almahbobi G. A direct action for growth 512 43. 513 hormone in improving oocyte quality in poor responder patients. Reproduction 2014. 514 44. Regan SLP, McFarlane JR, O'Shea T, Andronicos N, Arfuso F, Dharmarajan A et al. Flow cytometric analysis of FSHR, BMPR1B, LHR and apoptosis in granulosa cells and ovulation rate in 515 516 merino sheep. Reproduction 2015; 150 151-63. Hermann BP, Heckert LL. Transcriptional regulation of the FSH receptor: New perspectives. 517 45. 518 Molecular and Cellular Endocrinology 2007;260-262:100-8. 519 Brooks AJ, Wooh JW, Tunny KA, Waters MJ. Growth hormone receptor; mechanism of 46. 520 action. The International Journal of Biochemistry & Cell Biology 2008;40:1984-9. 521 Zhu T, Goh ELK, Graichen R, Ling L, Lobie PE. Signal transduction via the growth 47. 522 hormone receptor. Cellular Signalling 2001;13:599-616. Le Roith D, Bondy C, Yakar S, Liu J-L, Butler A. The Somatomedin Hypothesis: 2001. 523 48. 524 Endocrine reviews 2001;22:53-74.

- 525 49. Nakamura E, Otsuka F, Inagaki K, Miyoshi T, Matsumoto Y, Ogura K et al. Mutual
- 526 regulation of growth hormone and bone morphogenetic protein system in steroidogenesis by rat 527 granulosa cells. Endocrinology 2012;153:469-80.
- 528 50. Kucuk T, Kozinoglu H, Kaba A. Growth hormone co-treatment within a GnRH agonist long 529 protocol in patients with poor ovarian response: a prospective, randomized, clinical trial. Journal of 530 Assisted Reproduction and Genetics 2008;25:123-7.
- Suikkari A, MacLachlan V, Koistinen R, Seppälä M, Healy D, Double-blind placebo 531 51.
- 532 controlled study: human biosynthetic growth hormone for assisted reproductive technology. Fertil 533 Steril 1996; Apr 65:800-5.
- 534 Keane KN, Yovich JL, Hamidi A, Hinchliffe PM, Dhaliwal SS. Single-centre retrospective 52. 535 analysis of growth hormone supplementation in IVF patients classified as poor-prognosis. BMJ Open 536 2017;7.
- 53. Eftekhar M, Aflatoonian A, Mohammadian F, Eftekhar T, Adjuvant growth hormone therapy 537 538 in antagonist protocol in poor responders undergoing assisted reproductive technology. Arch 539 Gynecol Obstet 2013;287:1017-21.
- 540 54.
- Levy T, Limor R, Villa Y, Eshel A, Eckstein N, Vagman I et al. Another look at co-541 treatment with growth hormone and human menopausal gonadotrophins in poor ovarian responders.
- 542 Human Reproduction 1993;8:834-9.
- 543 Volpe A, Coukos G, Barreca A, Artini PG, Minuto F, Giordano G et al. Ovarian response to 55.
- 544 combined growth hormone-gonadotropin treatment in patients resistant to induction of 545 superovulation. Gynecological Endocrinology 1989;3:125-33.
- 546 56. Homburg R, Singh A, Bhide P, Shah A, Gudi A. The re-growth of growth hormone in 547 fertility treatment: a critical review. Human fertility 2012;15:190-3.
- 548 Kolibianakis EM, Venetis CA, Diedrich K, Tarlatzis BC, Griesinger G. Addition of growth 57. 549 hormone to gonadotrophins in ovarian stimulation of poor responders treated by in-vitro fertilization: 550 a systematic review and meta-analysis. Human Reproduction Update 2009;15:613-22.
- 551 Kyrou D, Kolibianakis EM, Venetis CA, Papanikolaou EG, Bontis J, Tarlatzis BC. How to 58. 552 improve the probability of pregnancy in poor responders undergoing in vitro fertilization: a 553 systematic review and meta-analysis. Fertility and Sterility 2009;91:749-66.
- 554 59. Gong JG, Bramley T, Webb R. The effect of recombinant bovine somatotropin on ovarian 555 function in heifers: follicular populations and peripheral hormones. Biology of Reproduction 556 1991;45:941-9.
- 557 60. Garverick H, Baxter G, Gong J, Armstrong D, Campbell B, Gutierrez C et al. Regulation of 558 expression of ovarian mRNA encoding steroidogenic enzymes and gonadotrophin receptors by FSH 559 and GH in hypogonadotrophic cattle. Reproduction 2002;123:651-61.
- 560 Bachelot A, Monget P, Imbert-Bolloré P, Coshigano K, Kopchick JJ, Kelly PA et al. Growth 61. 561 Hormone Is Required for Ovarian Follicular Growth. Endocrinology 2002;143:4104-12.
- 562 Scaramuzzi RJ, Murray JF, Downing JA, Campbell BK. The effects of exogenous growth 62. 563 hormone on follicular steroid secretion and ovulation rate in sheep. Domestic animal endocrinology 564 1999;17:269-77.
- 565 63. Slot KA, Kastelijn J, Bachelot A, Kelly PA, Binart N, Teerds KJ. Reduced recruitment and 566 survival of primordial and growing follicles in GH receptor-deficient mice. Reproduction 567 2006:131:525-32.
- 568 Gong JG, Baxter G, Bramley TA, Webb R. Enhancement of ovarian follicle development in 64. 569 heifers by treatment with recombinant bovine somatotrophin: a dose-response study. Journal of 570 reproduction and fertility 1997;110:91-7.
- Saccon TD, Moreira F, Cruz LA, Mondadori RG, Fang Y, Barros CC et al. Ovarian aging 571 65. 572 and the activation of the primordial follicle reserve in the long-lived Ames dwarf and the short-lived 573 bGH transgenic mice. Molecular and Cellular Endocrinology 2017;455:23-32.
- 574 Jia X-C, Kalmijn J, Hsueh AJW. Growth Hormone Enhances Follicle-Stimulating Hormone-66. 575 Induced Differentiation of Cultured Rat Granulosa Cells. Endocrinology 1986;118:1401-9.
- Spicer LJ, Klindt J, Buonomo FC, Maurer R, Yen JT, Echternkamp SE. Effect of porcine 576 67.
- somatotropin on number of granulosa cell luteinizing hormone/human chorionic gonadotropin 577
- 578 receptors, oocyte viability, and concentrations of steroids and insulin-like growth factors I and II in 579 follicular fluid of lean and obese gilts. Journal of animal science 1992;70:3149-57.
- 580 Rice S, Ojha K, Whitehead S, Mason H. Stage-specific expression of androgen receptor, 68.
- 581 follicle-stimulating hormone receptor, and anti-müllerian hormone type ii receptor in single, isolated,

- human preantral follicles: Relevance to polycystic ovaries. The Journal of Clinical Endocrinology &
- 583 Metabolism 2007;92:1034-40.
- 584 69. Grande M, Borobio V, Jimenez JM, Bennasar M, Stergiotou I, Peñarrubia J et al. Antral
- follicle count as a marker of ovarian biological age to reflect the background risk of fetal aneuploidy.
  Human Reproduction 2014;29:1337-43.
- 587 70. Greisen S, Ledet T, Ovesen P. Effects of androstenedione, insulin and luteinizing hormone 588 on steroidogenesis in human granulosa luteal cells. Human Reproduction 2001;16:2061-5.
- 589 71. Donadeu F, Ascoli M. The Differential Effects of the Gonadotropin Receptors on Aromatase
- 590 Expression in Primary Cultures of Immature Rat Granulosa Cells Are Highly Dependent on the
- 591 Density of Receptors Expressed and the Activation of the Inositol Phosphate Cascade. Endocrinology 592 2005;146:3907-16.
- 593 72. Rotmensch S, Dor J, Furman A, Rudak E, Mashiach S, Amsterdam A. Ultrastructural
- characterization of human granulosa cells in stimulated cycles: correlation with oocyte fertilizability.
   Fertility and Sterility 1986;45:671-9.

```
596
```

597

### 599 Fig. 1 Validation of immunofluorescent labelling

- 600 A. Subtraction of nonspecific binding (red) and auto-fluorescence (green) at ~10<sup>3</sup>; granulosa cell. B. Live
- 601 granulosa cells, unstained control for GHR auto-fluorescence (blue) compared to positive fluorescent signal
- 602 measurement (box). Gated and removed CD45 positive cells (circle) also confirmed binding specificity
- 603 (Saizen, Merck Serono, Australia). C. Live human granulosa cells with positive fluorescence for GH receptor
- 604 (a & b), and pre-absorbed GH for negative control and binding specificity of GHR (c & d). Bar 10 μm.
- 605

### 606 Fig. 2 Granulosal GHR density and ovarian reserve depletion.

- 607 GHR expression density on granulosa cells collected from patients during IVF treatment with a range of
- 608 ovarian reserves of follicles, A. 23-30 y patient group, B. 31-34 y patient group, C. 35-38 y patient group, D.
- 609 39+ y patient group. Ovarian reserve was measured indirectly by the antral follicle count (AFC). AFC is the
- 610 number of follicles between 2-10 mm on day 2-5 of a cycle. Mean fluorescent intensity (MFI) was obtained
- 611 using an average of ~8000 granulosa cells per follicle for the direct measurement of receptor protein
- 612 expression. The number within the column represents the number of follicles analysed for that group. The data
- 613 were subjected to statistical verification using one-way ANOVA with an uncorrected Fisher's LSD for
- $614 \qquad follicular size. Values in graphs are means \pm S.E.M., and differences were considered significant if *p<0.05,$
- $615 \qquad {**p}{<}0.01 \text{ and } {***p}{<}0.005.$
- 616

# Fig. 3 Follicle size and the granulosa cell density of GHR, FSHR, BMPR1B and LHR in poor response 39+ v patients co-treated with GH

619 Follicles of different sizes were individually collected and analysed. Granulosa receptor density during an IVF

- 620 cycle with or without GH co-treatment was measured by flow cytometry. A. GHR, B. FSHR, C. LHR and D.
- 621 BMPR1B. The number within the column represents the number of follicles analysed for that follicle size. Inset
- 622 A Combined follicles of different sizes-GHR. Inset B FSHR, Inset C LHR, and Inset D. The number within the
- 623 column represents the number of follicles analysed. Ovarian reserve measured indirectly by the antral follicle
- 624 count (AFC). AFC is the number of follicles between 2-10 mm on day 2-5 of a cycle. Mean fluorescent
- 625 intensity (MFI) was obtained using an average of ~8000 granulosa cells per follicle for the direct measurement
- 626 of receptor protein expression. The data were subjected to statistical verification using one-way ANOVA with
- 627 an uncorrected Fisher's LSD for follicular size. Values in graphs are means  $\pm$  S.E.M., and differences were
- 628 considered significant if \*p<0.05, \*\*p<0.01, \*\*\*p<0.005, and \*\*\*\*p<0.001.
- 629

### 630 Fig. 4 GH associated pregnancy and live birth outcome

- A. The effect of GH treatment on pregnancy rate during IVF treatment. B. The effect of GH treatment on
- 632 pregnancy rate during IVF treatment. The data were subjected to statistical verification using chi square. The
- 633 chi-square statistic *p*-value is p=0.0033. The data were based on the number of embryos transferred per patient
- age group or treatment, including subsequent frozen embryo cycles (FET). One patient with an ectopic
- 635 pregnancy (classed as miscarriage) was present in the 23-30 y and the 35-38 y groups. Patients were selected
- 636 randomly in a prospective regimen.
- 637

638

639

### 640

### 641 Table 1 Patient ovarian reserve, based on antral follicle count (AFC)

- 642 <sup>1</sup> Typical Ovarian Reserve for age group
- <sup>643</sup> <sup>2</sup> Ovarian reserve measured indirectly by the Antral Follicle Count (AFC). AFC is the number of follicles

between 2-10 mm on day 2-5 of a cycle: group A + = 30-39 follicles; group A = 20-29

- follicles; group B = 13-19 follicles; group C = 9-12 follicles, group D = 5-8 follicles; group E =  $\leq 4$  follicles.
- 646 Follicle count is based on the combined total from both ovaries to determine AFC. The number of follicles
- 647 aspirated from patients from the specified ovarian reserve group.
- 648 <sup>3</sup> CCF-Number of patients with complete failed fertilisation compared to same age group without GH
- <sup>4</sup> Percentage per total number of embryos transferred
- <sup>5</sup> The average number of oocytes collected at TVOA for the age group
- 651 a One patient with an ectopic pregnancy (classed as miscarriage)
- All subsequent frozen embryo cycles (FET) cycles were included in the analysis therefore the data was based
- on number of embryos transferred.
- \*\*p=0.003, \*p=0.041 Chi square test (d) =+GH 39+ y compared to (c) = 39+ y patient groups.

Age years	IVF patient	Total follicle	Ovarian reserve <sup>1</sup>	0\	Ovarian Reserve Group <sup>2</sup> Number of follicles collected per group					Oocyte quality			Fertility N (%) <sup>4</sup>			
				A+	А	В	С	D	Е	# <sup>5</sup>	CCF <sup>3</sup>	ET	Not Pregnant	Pregnant	Miscarriage	Live Birth
21-30 <sub>a</sub>	10	68	20-40	26	42	-	-	-	-	10	0	12	4(33)	8(67)	3(37)	5(42)
31-34	12	96	13-29	-	48	23	16	9	-	8	0	15	9(60)	7(47)	1(14)	6(40)
35-38 <sub>a</sub>	12	108	9-19	-	6	46	17	34	-	9	0	16	5(31)	11(68)	5(46)	4(25)
39-45	19	131	3-8	-	-	42	5	64	19	7	3	25	22(88)	3(12 <sup>c</sup> )	2(68)	1(4 <sup>°</sup> )
+GH39-45	11	48	3-8	-	-	-	-	25	23	4.5	3	10	4(40)	6(60 <sup>d</sup> )**	<sup>k</sup> 4(68)	2(20 <sup>d</sup> )*