



Ovarian stimulation for IVF/ICSI

UPDATE 2025

Guideline of European Society of Human Reproduction and Embryology

The ESHRE Ovarian Stimulation Guideline Group



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How to cite the guideline

The ESHRE Guideline Group on Ovarian Stimulation, Ata B., Bosch E., Broer S., Griesinger G., Grynberg M., Kolibianakis E., Kunicki M., La Marca A., Lainas G., Le Clef N., Massin N., Polyzos N.P., Sunkara S.K., Timeva T., Töyli M., Urbancsek J., Broekmans F.

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199		

Draft for review



Introduction to the guideline

200

201

202 Ovarian stimulation for IVF/ICSI has not been addressed by existing evidence-based guidelines. Ovarian
203 stimulation for IVF/ICSI has been discussed briefly in the NICE guideline on Fertility problems
204 (<https://www.nice.org.uk/guidance/cg156>) and the Royal Australian and New Zealand College of
205 Obstetricians and Gynaecologist has published a statement on ovarian stimulation in assisted
206 reproduction

207 ([https://www.ranzcog.edu.au/RANZCOG_SITE/media/RANZCOG-
208 MEDIA/Women%27s%20Health/Statement%20and%20guidelines/Clinical%20-
209 %20Gynaecology/Ovarian-Stimulation-in-infertility-\(C-Gyn-2\)-Review-Mar-14.pdf?ext=.pdf](https://www.ranzcog.edu.au/RANZCOG_SITE/media/RANZCOG-MEDIA/Women%27s%20Health/Statement%20and%20guidelines/Clinical%20-%20Gynaecology/Ovarian-Stimulation-in-infertility-(C-Gyn-2)-Review-Mar-14.pdf?ext=.pdf)).

210 A narrative review of evidence provided for WHO guidance on management of ovarian stimulation for
211 IVF was published in 2017, but this document did not include recommendations (Farquhar et al., 2017).

212 Based on the lack of guidelines, the ESHRE SIG Reproductive Endocrinology initiated the development
213 of an ESHRE guideline focussing on all aspects of ovarian stimulation, which was published in 2019
214 (ESHRE Ovarian Stimulation guideline group, 2020).

215 The current guideline is an update of the version from 2019, with amendments to the
216 recommendations based on recently published data. Where amendments were made, this is labelled
217 as such [updated]. If the GDG felt rewording of a recommendation was necessary without new evidence
218 on the topic, this was indicated with [reworded].

219 The 2019 guideline and the update are developed according to a well-documented methodology,
220 universal to ESHRE guidelines and described in the Manual for ESHRE guideline development
221 (www.eshre.eu). Details on the methodology of the current guideline are outlined in Annex 4.

222 The guideline development group (GDG) for the current update consisted of the previous guideline
223 group with minor changes. One member of the GDG (2019) decided to step down and was replaced.
224 The members of the guideline development group are listed in Annex 1.

225 GUIDELINE SCOPE

226 The aim of this guideline is to provide clinicians with evidence-based information on the different
227 options for the performance of ovarian stimulation for IVF/ICSI, taking into account issues such as the
228 'optimal' ovarian response, live birth rates, safety, patient compliance, and individualisation.
229 Knowledge gaps were identified and prioritized.

230 The following issues were outside the scope of the current document: patients with specific medical
231 conditions (except for PCOS), and treatment of the ovarian hyper-stimulation syndrome (OHSS).

232 TARGET USERS OF THE GUIDELINE

233 Infertility specialists and specialty nurses performing the daily care for patients undergoing ovarian
234 stimulation for the purpose of IVF/ICSI.



235 TERMINOLOGY

236 Ovarian stimulation is defined as pharmacological treatment with the intention of inducing the
237 development of ovarian follicles and trigger the ovulation process of these follicles. It can be used for
238 two purposes: 1) for timed intercourse or insemination; 2) for IVF/ICSI, to obtain multiple oocytes at
239 follicular aspiration (Zegers-Hochschild et al., 2017). The GDG decided to use the term ovarian
240 stimulation (OS) confined to ovarian stimulation for IVF/ICSI.

241 The GDG would also like to point to the importance of ‘simplicity of ovarian stimulation’. When
242 comparing compounds, dosages or add-on treatments for ovarian stimulation in this guideline
243 document, preference was always given to the more basic option, unless a clear benefit of more
244 complex treatments was shown.

245 Response after ovarian stimulation is usually classified as poor, normal and excessive. However, this
246 terminology can be potentially stigmatising/traumatising towards patients. Therefore, the GDG would
247 like to propose to use the terminology low, normal and high response to categorize (the observed as
248 well as the expected/predicted) response to OS for future referencing. However, the definition of low
249 response proposed in this guideline is the same as the definition of the Bologna poor responder and
250 the poor responder as defined by ICMART (Ferraretti et al., 2011, Zegers-Hochschild et al., 2017).

251 Due to the lack of universally accepted definitions of high and low ovarian response, the definitions and
252 terminology in the studies included in the evidence synthesis were varied. However, for future practice
253 and research, the GDG suggests using the following definitions:

- 254 - High ovarian response is an exaggerated response to conventional ovarian stimulation (150-
255 225 IU FSH), characterized by the presence of more follicles and/or oocytes than intended
256 (Griesinger et al., 2016). Generally, more than 18 follicles ≥ 11 mm in size on day of oocyte
257 maturation trigger and/or 18 oocytes collected characterize a high response (Griesinger et al.,
258 2016), defined by a risk increase for OHSS occurrence.
- 259 - Low ovarian response is a diminished response to conventional ovarian stimulation,
260 characterized by the presence of a low number of follicles and/or oocytes (Ferraretti et al.,
261 2011). Generally, ≤ 3 follicles on day of oocyte maturation trigger and/or ≤ 3 oocytes obtained
262 characterize a low response.

263 In this guideline, in line with the research, terminology and discussion on ovarian stimulation is focused
264 on women. The guideline group recognises that there are individuals who do not identify with the terms
265 used in the literature. For the purposes of this guideline, we use the terms “women”, “patients”,
266 “low/poor responder”, “normal responder” and “high responder”, however, it is not intended to
267 isolate, exclude, or diminish any individual’s experience nor to discriminate against any group.

268 Outcomes for this guideline

269 The guideline focuses on outcomes of efficacy, safety and patient-related outcomes.

270 The critical outcomes for this guideline are **efficacy** in terms of cumulative live birth rate (CLBR) per
271 started cycle and live birth rate (LBR) per started cycle; and **safety** in terms of the risk of moderate
272 and/or severe OHSS.



273 Other outcomes used for efficacy were (in order of importance) cumulative ongoing pregnancy rate per
274 started cycle, clinical pregnancy rate per started cycle, number of MII oocyte retrieved (yield), number
275 of oocytes retrieved.

276 Other outcomes used for safety include incidence of different grades of OHSS, cycle cancellation for
277 hyper-response, intra-abdominal or vaginal bleeding, infection, ovarian torsion, long-term effects on
278 maternal/child health, and other treatment-related adverse events.

279 Patient-related outcomes are compliance, drop-out rates, patient burden, quality of life (QoL), and
280 patient preferences.

281 All outcomes were defined, where possible, as per started cycle.

282

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299



Introduction

300

301

302 **IVF: the purpose and significance.**

303 Infertility is a disease state with potential profound consequences for the quality of life of both men
304 and women. Reproduction is one of the key elements of life and failing to achieve the creation of
305 offspring may lead to lifelong mental and physical health problems. Also, couples faced with infertility
306 are frequently subjected to long-lasting, time consuming and agonizing treatment schedules, living
307 often between hope, fear and frustration (Brandes et al., 2010, Brandes et al., 2009, Gameiro and
308 Finnigan, 2017). The development of IVF as a tool for treating infertility as a result of tubal disease,
309 severe male factor causes, anovulation and even, although not convincingly proven, conditions like
310 unexplained infertility, has brought enormous potential to the infertility treatment armamentarium.
311 Still, of all couples visiting infertility centres, roughly 35-40% will not achieve the so desired goal, in spite
312 of lengthy efforts, including IVF, and most of these couples will remain permanently childless
313 (McLernon et al., 2016, Olivius et al., 2002). This indicates that currently we still have areas of low-level
314 knowledge on the key factors of success, such as gamete quality, embryo quality and endometrial
315 receptivity. Improving the IVF technology may well depend on progress in these fields of research.

316 **Stimulation: how important is it.**

317 Very soon after the development of the IVF technology, performing IVF in a natural menstrual cycle
318 was superseded by the use of ovarian stimulation in order to obtain multiple oocytes. This was aimed
319 at solving two problems: one was the elimination of the risk of having no oocyte at all. The other was
320 the urge to improve efficiency by obtaining several embryos and replacing the best quality embryo(s)
321 to improve the probability of pregnancy. Ovarian stimulation has thereby become one of the
322 cornerstones of the IVF treatment, next to the *in vitro* handling of gametes and embryos, and the
323 embryo replacement procedure. The relative contribution to the overall success of IVF by the ovarian
324 stimulation phase is difficult to assess. Many years of research have aimed at optimizing this specific
325 phase. Many issues have been addressed, ranging from using urinary FSH products or recombinants,
326 using high or low FSH dosages, final oocyte maturation with urinary or recombinant, high or low dosage
327 of hCG, adding LH or LH like activity to the FSH as principal drug, management of high and low
328 responders, to the use of adjuvant medications to improve follicle availability and quality, etcetera. At
329 the same time, debates have been there on strong beliefs, like “the more (oocytes) the better”, less
330 (mild stimulation) is more (quality), “normal (8-17 oocytes) is the best”, and “we need eggs, not ALL the
331 eggs”. It seems that agreement on the optimal ovarian stimulation approach, aimed at getting more
332 than 1 oocyte, as in the normal menstrual cycle, is far from settled.

333 **Basics: FSH elevation.**

334 Complex as it seems, the endocrine background for ovarian stimulation is quite straightforward. FSH
335 levels must become elevated above the level that normally will help to select and grow ONE follicle out
336 of a group of antral follicles presenting in the FSH ‘window’. During this window, levels of FSH surpass
337 a certain threshold above which follicle granulosa cells become responsive for proliferative actions,
338 leading to expansion of the granulosa cell mass and the follicle fluid volume, typically of only one follicle,
339 while other potential responsive follicles continue to enter and proceed the stages of atresia. In
340 surpassing the threshold to a greater extent, and for a much longer period of time with the use of



341 ovarian stimulation, more than one follicle will become capable of entering this dominant follicle
342 development stage. The tools available for increasing FSH exposure are several, but basically most
343 comprise preparations containing FSH. The source of FSH can be urinary (purified or highly purified) or
344 recombinant (the FSH molecule is produced by programmed cells from hamster, mouse or human).
345 Some preparations combine FSH with LH, or LH like activity (hCG). The vast majority of FSH compounds
346 are distributed for dosing in International Units, a standardisation based on an oestradiol output bio-
347 assay. Only one compound is delivered in micrograms, and dosing here is based on a dosing algorithm.

348 Apart from administering FSH as an exogenous drug, compounds such as selective oestradiol receptor
349 blockers or oestradiol biosynthesis inhibitors may yield the same effect: increased and prolonged FSH
350 exposure.

351 **Source: Ovarian Antral Follicles, continuous versus cyclic recruitment.**

352 The follicles presenting in the window of elevated FSH levels are part of a continuous recruitment
353 process. Starting from the resting pool of primordial follicles, follicles develop through several phases,
354 reaching the antral stages after approximately 200 days (McGee and Hsueh, 2000). At that time point
355 they attain relevant FSH sensitivity. Without FSH exposure, such as in the prepubertal years, these
356 follicles will reach maximum sizes of 2-3 mm and vanish into the process of atresia. Without any FSH
357 exposure, this wastage process would continue until around the age of 50 years, when the ovarian
358 primordial follicle pools will have become depleted. It is the presence of FSH in varying levels that allows
359 the ovaries to pick up follicles in the antral stages, which become more prominent at ultrasound, and
360 from there deliver the ovulating follicle of the month, or, as in ovarian stimulation, recruit several to
361 many follicles from those that present in the window of opportunity to respond to FSH. This ovarian
362 activity is referred to as cyclic recruitment. The number of follicles that present in the opportunity
363 window of cyclic recruitment is highly variable between women and between age groups. As a general
364 rule, the number of antral follicles that can be stimulated will decline gradually with increasing age, as
365 an expression of the shrinking pool of primordial follicles.

366 **Store of Antral Follicles: can we manipulate it?**

367 Obtaining only few oocytes is an agonizing condition, as it may affect the prospects for a live birth in
368 IVF, albeit that this prospect is also very much determined by the age of the woman. Still, there is a
369 continuous search for methods to improve the egg number in low responders, and from the
370 aforementioned, it can be deduced that such method should interfere with early stages of follicle
371 development, where initial recruitment and/or later survival during continuous recruitment is
372 promoted. Numerous strategies and interventions have been suggested to enhance this sequence of
373 events; however, clinical useful strategies are still awaited.

374 **Oocyte number and Dosage: what is the relation like?**

375 The cohort of antral follicles being the finite source for oocytes, the level of exposure to FSH may add
376 to the total number of oocytes obtained. With the need of a minimum exposure to grow more than 1
377 follicle, there seems to be a positive relation between FSH dosage and oocyte yield, ranging from about
378 50 IU daily for a minimal response of 2 oocytes up to about 225 IU to obtain a maximal response (Lensen
379 et al., 2017, Sterrenburg et al., 2011). For the optimal response level in terms of oocytes a daily dosage
380 of 150 to 225 IU is mostly considered as standard. This implies that when using a stimulation dosage of
381 150 IU per day and creating a low follicle response, the range of opportunities in dose adjustments is
382 likely to be limited. This is certainly much dependant on the of Antral Follicle Count or AMH result. With
383 test results below a certain level, the so called predicted low responder may not produce more oocytes



384 with a higher FSH dosage. With AFC and/or AMH levels within the normal range, an unexpected low
385 responder may well obtain more oocytes with a higher FSH dosage. The question then remains whether
386 more oocytes will improve the prospects for a live birth? We still need to see evidence that a few
387 oocytes more or less will make the desired or feared difference in terms of live birth rates. At this point
388 it may be emphasized that the various cross-sectional cohort data on the relation between oocyte
389 number and cumulative live birth rates have suggested that 'more is better' and 'less is bad'. These
390 observations are correlation data, without the possibility to conclude that there is a causal relationship.
391 With respect to the latter, we may reflect on the implications of many randomised comparative trials
392 demonstrating that a few more or less oocytes within the individual couple will fail to make an obvious
393 difference in the live birth prospects.

394

395 At the other side of the spectrum, a high response to a standard dosage of 150 IU may be undesirable
396 as it is a potential source for the development of the Ovarian Hyperstimulation Syndrome (OHSS), even
397 today a potential life-threatening condition. Reduction of the FSH stimulation dosage may bring a more
398 mitigated response, with better safety, without jeopardizing overall live birth prospects. However, it is
399 to be understood that the driver of the syndrome occurring in high responder cases in fact is the
400 exposure of the granulosa cells to human chorion gonadotropin (hCG). Necessary as this may be for the
401 final oocyte competence attainment, circumventing administration of this drug by creation of an
402 endogenous LH surge by applying a GnRH agonist trigger is certainly a powerful way to decrease the
403 risk of OHSS. Finally, prevention of pregnancy-derived hCG to occur by freezing all embryos is another
404 important and logical step.

405 **Control on ovulation: agonists and antagonist.**

406 When stimulating the ovaries to create multifollicular development, the fast-rising oestradiol levels may
407 elicit an untimely LH surge. Untimely, as follicles may not have grown sufficiently large to ensure the
408 best quality oocytes, and when passed unnoticed, oocyte pick up may become a failed procedure. The
409 use of agents that block the signalling by the GnRH pulse generator towards the pituitary, such as GnRH
410 agonists, GnRH antagonists and progestins, have almost completely ruled such mishaps and have
411 greatly contributed to the efficiency of ovarian stimulation for IVF/ICSI.

412 **Oocytes, and then?**

413 Although the primary goal of ovarian stimulation is obtaining several oocytes, the timed replacement
414 of the embryo necessitates parallel and physiologically correct development of the endometrium.
415 Implantation is dependent on proper endocrine conditions, such as oestradiol exposure, in order to
416 ensure proliferation, and progesterone exposure commencing around ovulation in order to have the
417 endometrium differentiated into a receptive state. Stimulation per se is a guarantee for oestradiol
418 synthesis and release from the many developing follicles. The LH peak, or as in many cases, hCG
419 exposure, will enable granulosa cell differentiation into a progesterone producing system, that, in
420 normal condition, will be driven by continued endogenous LH pulses. In the GnRH agonist suppression
421 and GnRH antagonist approach, the interference with the GnRH receptor will lead to LH levels dropping
422 to low levels, and the hCG exposure here takes over the role of LH in maintaining luteal function up till
423 maximally 7-9 days after the ovulation trigger. On top of that, supraphysiological exposure to
424 endogenous estradiol and progesterone, driven by the exogenous administration of FSH and later hCG,
425 will further add in the insufficiency of the pituitary to produce the amounts of LH needed for continued
426 support of the corpora lutea. As such, luteal support is almost exclusively applied in the form of
427 exogenous natural progesterone, which is initiated often already at the day of follicle aspiration.



428 However, pharmacokinetics may not always be very stable for these compounds, and when
429 endogenous LH exposure by using an GnRH agonist trigger is applied, instead of the hCG signal, luteal
430 phase becomes insufficient in many cases even with the current exogenous progesterone
431 administration. The luteal phase support approach therefore remains an important area of research for
432 improvement of the quality of the embryo implantation phase.

433 Many years of basic and clinical research have delivered us tools for ovarian stimulation that make this
434 procedure effective, efficient, safe and an essential contribution to the total process of Assisted
435 Reproduction. In this guideline, important knowledge is brought together using a set of relevant
436 questions, for which searches and selections of the literature, grading of the knowledge base regards
437 quality, and well-balanced recommendations will provide the best possible answers to the question.
438 These recommendations will help clinicians to decide on what best to do or better not to do in clinical
439 conditions where we wish to provide optimal care to our patients.

440

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List of all recommendations

Recommendation	Strength	Quality of evidence	Remarks
Part A: Pre-stimulation evaluation			
Ovarian response prediction			
1	For predicting high and low response to ovarian stimulation, use of either antral follicle count (AFC) or anti-Müllerian hormone (AMH) is recommended. [updated]	Strong	⊕○○○
2	Age, BMI, basal FSH, inhibin B basal oestradiol, basal progesterone and basal LH are not recommended for the prediction of ovarian response. [2025]	Strong	⊕○○○
Pregnancy prediction			
3	AFC, AMH, basal FSH, basal LH, basal oestradiol, basal progesterone and inhibin B are not recommended for the prediction of pregnancy and live birth. [updated]	Strong	⊕○○○
4	Female age and BMI are predictors of pregnancy and live birth. [2025]	Strong	⊕○○○
Part B: Pre-treatment therapies			
Pre-treatment therapies			
5	Pre-treatment with oestrogen before ovarian stimulation using the GnRH antagonist protocol is not recommended for improving efficacy. [updated]	Strong	⊕⊕○○ SoF table 1 a,b
6	Pre-treatment with progesterone before ovarian stimulation is probably not recommended for improving efficacy. [reworded]	Conditional	⊕⊕○○ SoF table 2 a,b
7	Oestrogen or progesterone pre-treatment can be used for scheduling purposes given the data on efficacy and safety. [reworded]	GPP	
8	COCP pre-treatment (12-28 days) is not recommended in the GnRH antagonist protocol with FSH alone stimulation, because of reduced efficacy. [updated]	Strong	⊕⊕○○ SoF table 3 a,b,c
9	A minimal wash out period of 5 days should be applied if COCP is used for programming cycle in the case of a fresh transfer. [2025]	GPP	
10	GnRH antagonist pre-treatment before ovarian stimulation in a delayed-start gonadotrophin protocol is probably not recommended. [2019]	Conditional	⊕○○○ SoF table 4 a,b
11	hCG pre-treatment can only be used in the context of a clinical trial. [2025]	Research only	



Pituitary suppression and ovarian stimulation

Stimulation protocols

12	Delayed-start ovarian stimulation is probably not recommended routinely in predicted high responders to decrease the risk of OHSS. [2025]	Conditional	⊕○○○	SoF table 5
/	There is no evidence to justify the use of NC or MNC for OS in high responders.	/	/	Conclusion
13	A reduced gonadotropin dose is probably recommended to decrease the risk of OHSS in predicted high responders. [2025]	Conditional	⊕○○○	SoF table 6
14	The GnRH antagonist protocol is recommended for predicted high responders. However, if GnRH agonist protocols are used, a reduced gonadotropin dose is recommended to decrease the risk of OHSS. [updated]	Strong	⊕○○○	
15	Delayed-start ovarian stimulation is probably not recommended over a conventional gonadotrophin dose for predicted normal responders. [2025]	Conditional	⊕○○○	SoF table 7
16	Neither a reduced nor increased gonadotrophin dose is probably recommended over a conventional gonadotrophin dose (equivalent to 150-225 IU) for predicted normal responders. [updated]	Conditional	⊕○○○	SoF table 8
17	Delayed start ovarian stimulation is probably not recommended for predicted low responders. [2025]	Conditional	⊕○○○	SoF table 9
18	The use of modified natural cycle is probably not routinely recommended over conventional stimulation for low responders. [updated]	Conditional	⊕○○○	
19	The GDG recognises that low responders are a heterogeneous group and in women with very low ovarian reserve, clinicians could choose to use a modified natural cycle. [2025]	GPP		
20	A higher gonadotropin dose is probably not recommended over conventional (equivalent to 150-225 IU) for predicted low responders. [updated]	Conditional	⊕○○○	SoF table 10
21	A gonadotropin dose higher than 300 IU is not recommended for predicted low responders. [2019]	Strong	⊕○○○	

Pituitary suppression regimes

22	If GnRH agonists are used, the long GnRH agonist protocol is recommended over the short or ultrashort GnRH agonist protocol. [updated]	Strong	⊕⊕○○	SoF table 11 a,b
23	The GnRH antagonist protocol is recommended over the GnRH agonist protocols given the comparable efficacy and higher safety in the general IVF/ICSI population. [2019]	Strong	⊕⊕⊕○	SoF table 12 a,b
24	The flexible and fixed GnRH antagonist protocol is probably equally recommended. [2025]	Conditional	⊕⊕○○	
25	If freeze-all is planned, the use of progestin for pituitary suppression is probably equally recommended to GnRH analogues. [updated]	Conditional	⊕○○○	SoF table 13 a,b,c,d



Types of gonadotropins and other ovarian stimulation drugs

26	The use of recombinant FSH (rFSH) and human menopausal gonadotropin (hMG) for ovarian stimulation is equally recommended. [2019]	Strong	⊕⊕⊕○	SoF table 14
27	The use of recombinant FSH (rFSH) and purified FSH (p-FSH) for ovarian stimulation in GnRH agonist protocol is equally recommended. [2019]	Strong	⊕⊕○○	SoF table 15
28	The use of either recombinant FSH (rFSH) and highly purified FSH (hp-FSH) for ovarian stimulation in GnRH agonist protocol is equally recommended. [2019]	Strong	⊕⊕○○	SoF table 16
29	The combination of rFSH with rLH and rFSH alone are probably equally recommended for the general IVF population. [2025]	Conditional	⊕⊕○○	SoF table 17a
30	The combination of rFSH with rLH and rFSH alone are probably equally recommended for low responders. [2025]	Conditional	⊕⊕○○	SoF table 17b
31	The combination of rFSH with rLH and rFSH alone are probably equally recommended for women of advanced age (≥35 year). [2025]	Conditional	⊕⊕○○	SoF table 17c
32	The combined use of recombinant FSH with Human Menopausal Gonadotropin, either from the start or mid-phase of ovarian stimulation, is probably not recommended over the use of either recombinant FSH or hMG alone in normal and low responders. [2025]	Conditional	⊕⊕○○	SoF table 18 a,b
33	The use of long-acting and daily recombinant FSH (rFSH) is equally recommended in GnRH antagonist cycles for normal responders. [2019]	Strong	⊕○○○	SoF table 19
34	Follitropin delta and follitropin alpha/beta are probably equally recommended for ovarian stimulation. [2025]	Conditional	⊕○○○	
35	The use of highly purified FSH (hp-FSH) and human menopausal gonadotropin (hMG) for ovarian stimulation in GnRH agonist protocols is equally recommended. [2019]	Conditional	⊕⊕○○	
36	The use of recombinant LH + recombinant FSH (rFSH+rLH) for ovarian stimulation is probably not recommended over hMG in GnRH agonist protocols with regards to safety. [2019]	Conditional	⊕○○○	
37	Adding low dosages of hCG to the FSH stimulation is probably not recommended. [2025]	Conditional	⊕○○○	SoF table 20 a,b,c
38	A stimulation scheme starting with gonadotropins followed by letrozole is probably not recommended over gonadotropins alone in low responders. [reworded]	Conditional	⊕○○○	
39	The addition of letrozole to gonadotropins in stimulation protocols for predicted high responders is probably not recommended. [updated]	Conditional	⊕○○○	SoF table 21 a
40	The addition of letrozole to gonadotropins in stimulation protocols is probably not recommended for predicted normal responders. [2019]	Conditional	⊕○○○	SoF table 21 b
41	The addition of letrozole to gonadotropins in stimulation protocols is probably not recommended for predicted low responders. [2019]	Conditional	⊕⊕○○	
/	There is no evidence available to recommend the substitution of FSH by Clomiphene Citrate in ovarian stimulation.	/	/	Conclusion



42	The addition of Clomiphene Citrate to gonadotropins in stimulation protocols is probably not recommended for predicted high responders. [2019]	Conditional	⊕⊕○○	SoF table 22 a
43	The addition of Clomiphene Citrate to gonadotropins in stimulation protocols is probably not recommended for predicted normal responders. [2025]	Conditional	⊕⊕⊕○	SoF table 22 b
44	Clomiphene citrate alone or in combination with gonadotrophins, and gonadotropin stimulation alone are probably equally recommended for predicted low responders. [updated]	Conditional	⊕⊕○○	

Adjustment of gonadotropin dose

45	Adjustment (increase or decrease) of the gonadotrophin dose in the mid-stimulation phase during ovarian stimulation is probably not recommended. [2019]	Conditional	⊕○○○	
46	Given the lack of evidence of the value of dose adjustments during ovarian stimulation, it is important that the gonadotropin starting dose is appropriate based on patient characteristics and desired outcome. [2025]	GPP		

Adjunct therapies

47	Routine use of adjuvant metformin before and/or during ovarian stimulation is probably not recommended when using the GnRH antagonist protocol for women with PCOS. [Updated]	Conditional	⊕⊕○○	SoF table 23
48	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for normal responders. [2025]	Strong	⊕○○○	SoF table 24 a
49	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for low responders. [Updated]	Strong	⊕○○○	SoF table 24 b
50	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for women with PCOS. [2025]	Strong	⊕⊕○○	
51	Use of testosterone before ovarian stimulation is not recommended for low responders. [updated]	Strong	⊕⊕⊕○	SoF table 25
52	Use of DHEA before and/or during ovarian stimulation is not recommended for low responders. [2019]	Strong	⊕⊕○○	SoF table 26
53	Use of DHEA before and/or during ovarian stimulation is not recommended for normal responders. [2025]	Strong	⊕⊕○○	
54	Use of aspirin before and/or during ovarian stimulation is not recommended in the general IVF/ICSI population and for low responders. [2019]	Strong	⊕⊕⊕○	SoF table 27
55	Use of sildenafil before and/or during ovarian stimulation is not recommended for poor responders. [2019]	Strong	⊕○○○	
56	Use of myo-inositol before and/or during ovarian stimulation is probably not recommended for women with PCOS undergoing IVF. [2025]	Conditional	⊕○○○	SoF table 28 a
57	Use of myo-inositol before and/or during ovarian stimulation is not recommended in low responders. [2025]	Strong	⊕⊕○○	
58	Use of myo-inositol before and/or during ovarian stimulation is not recommended in non-PCOS women undergoing IVF. [2025]	Strong	⊕⊕○○	SoF table 28 b



Non-conventional start of ovarian stimulation

59	Random-start ovarian stimulation could be used when a fresh transfer is not intended and there is no possibility of natural conception. [Reworded]	GPP	
60	Luteal start ovarian stimulation could be used when a fresh transfer is not intended and there is no possibility of natural conception. [Updated]	Conditional	⊕○○○
61	Late luteal phase start of gonadotropins with fresh transfer is probably not recommended for low responders. [Updated]	Conditional	⊕○○○
62	Double stimulation can be considered for urgent fertility preservation cycles. [2019]	GPP	
63	Double stimulation can be used with the intention to accumulate oocytes or embryos when fresh transfer is not planned. [Updated]	Strong	⊕⊕○○

Part D: Fertility preservation and oocyte donation

Fertility preservation for patients facing gonadotoxic treatment

64	For patients facing gonadotoxic treatment, ovarian stimulation for fertility preservation should be started irrespective of the menstrual cycle phase. [updated]	Strong	⊕○○○
65	For ovarian stimulation in women seeking fertility preservation for medical reasons the GnRH antagonist protocol is probably recommended. [2019]	Conditional	⊕○○○
66	In ovarian stimulation for fertility preservation in oestrogen sensitive diseases the concomitant use of anti-oestrogen therapy, such as letrozole or tamoxifen, can be considered. [2019]	GPP	
67	For final oocyte maturation, hCG is preferred, unless the patient is at risk of early OHSS, in which case GnRH agonist trigger is advised. [2025]	GPP	

Elective oocyte cryopreservation

68	Ovarian stimulation for elective oocyte preservation can be started irrespective of the menstrual cycle phase. [2025]	Conditional	⊕○○○
69	GnRH antagonist or progestin protocol are probably recommended over GnRH agonist protocols for pituitary suppression in elective oocyte cryopreservation. [2025]	Conditional	⊕○○○
70	For final oocyte maturation in elective oocyte cryopreservation, hCG is preferred, unless the patient is at risk of early OHSS, in which case GnRH agonist trigger is advised. [2025]	GPP	

Oocyte donation

71	Conventional follicular start or random-start ovarian stimulation are equally recommended for oocyte donation cycles. [2025]	Strong	⊕○○○
72	If random-start ovarian stimulation is used, oocyte donors need to adopt contraceptive measures to prevent the possibility of a natural pregnancy. [2025]	GPP	



73	The use of any type of contraception (hormonal, non-hormonal, oral, vaginal or intrauterine) before or during ovarian stimulation is not a contraindication in oocyte donors. [2025]	GPP	
74	For pituitary suppression in oocyte donors the GnRH antagonist and progestin protocol are probably equally recommended. [2025]	Strong	⊕⊕○○
75	A GnRH agonist protocol is not recommended in oocyte donors. [2025]	GPP	
76	The use of recombinant FSH, purified FSH, long-acting rFSH or hMG is probably equally recommended in oocyte donors undergoing ovarian stimulation. [2025]	Conditional	⊕○○○
77	Gonadotropin dose should be individualised based on ovarian reserve with the goal to maintain donors' safety and also obtain an optimal number of oocytes. [2025]	GPP	
78	The routine use of a GnRH agonist trigger is recommended in oocyte donors using the GnRH antagonist or progestin protocols for pituitary suppression. [2025]	Strong	⊕⊕○○
79	The use of a hCG trigger is not routinely recommended in oocyte donation cycles. [2025]	Strong	⊕⊕○○

Part E: Monitoring

Hormonal assessment during ovarian stimulation

80	The addition of oestradiol measurements to ultrasound monitoring is probably not recommended. [2019]	Conditional	⊕⊕○○
81	The addition of a hormonal panel consisting of a combination of oestradiol, progesterone and LH measurements to ultrasound monitoring is probably not recommended. [2019]	Conditional	⊕○○○

Endometrial thickness

82	Routine monitoring of endometrial thickness during controlled ovarian stimulation is probably not recommended. [2019]	Conditional	⊕○○○
83	The guideline group suggests performing a single measurement of the endometrium during ultrasound assessment on the day of triggering or oocyte pick-up to counsel patients on potential lower pregnancy chance. [2019]	GPP	

Criteria for triggering

84	The association of follicle size as a triggering criterion with outcome has not been sufficiently studied. Physicians may choose the follicle size upon which final oocyte maturation is triggered on a case to case basis. [2019]	Conditional	⊕⊕○○
85	The decision on timing of triggering in relation to follicle size is multi-factorial, taking into account the size of the growing follicle cohort, the hormonal data on the day of pursued trigger, duration of stimulation, patient burden, financial costs, experience of previous cycles and organizational factors for the centre. Most often, final oocyte maturation is triggered at sizes of several of the leading follicles between 16-22 mm. [2019]	GPP	



86	The GDG does not recommend to base timing of final oocyte maturation triggering on oestradiol levels alone. [2019]	GPP
87	The GDG does not recommended to base timing of final oocyte maturation on oestradiol/follicle ratio alone. [2019]	GPP

Hormonal assessment on the day of final oocyte maturation

88	It is probably recommended to measure serum progesterone levels on the day of final oocyte maturation in cycles aimed for a fresh embryo transfer. [2025]	Conditional	⊕○○○
89	If serum progesterone levels are high, the patient should be counselled about potentially lower ongoing pregnancy/live birth rates. The decision to defer embryo transfer should include other factors (number of oocytes, number of embryos, and embryo quality). [2025]	GPP	
90	It is not recommended to routinely measure serum oestradiol levels on the day of HCG trigger in ovarian stimulation cycles with an intent for a fresh embryo transfer. [2025]	Strong	⊕○○○
91	It is not recommended to measure serum LH levels on the day of HCG trigger in ovarian stimulation cycles aimed for a fresh embryo transfer. [2025]	Strong	⊕○○○
92	It is not recommended to measure serum oestradiol, progesterone or luteinizing hormone levels on the day of a GnRH agonist trigger in freeze-all cycles. [2025]	Strong	⊕○○○

Criteria for cycle cancellation

93	A low response to ovarian stimulation alone is not a reason to cancel a cycle. [2019]	Strong	⊕○○○
94	The physician should counsel the individual unexpected low responder regarding pregnancy prospects and decide individually whether to continue this cycle. [Updated]	GPP	
95	In GnRH agonist cycles with an ovarian response of ≥19 follicles of ≥11 mm, there is an increased risk of OHSS and preventative measures are recommended, which should include primarily cancelling final oocyte maturation trigger. [Updated]	Strong	⊕○○○
96	In GnRH antagonist cycles, withholding GnRH agonist triggering may still be considered in women with extremely high ovarian response. [2025]	GPP	

Part F: Triggering ovulation and luteal support

Triggering of final oocyte maturation

97	The use of recombinant hCG and urinary hCG is equally recommended for triggering final oocyte maturation during ovarian stimulation protocols. [2019]	Strong	⊕⊕○○	SoF table 29
98	A reduced-dose of 5000 IU urinary hCG for final oocyte maturation is probably recommended over a 10.000 IU dose in GnRH agonist protocols, as it may improve safety. [2019]	Conditional	⊕○○○	



99	It is not recommended to administer recombinant LH for triggering final oocyte maturation. [2019]	Strong	⊕○○○	SoF table 30
100	The use of GnRH agonist for final oocyte maturation is not recommended in the general IVF/ICSI population with fresh transfer, regardless of luteal phase support (with or without LH-activity). [updated]	Strong	⊕⊕○○	SoF table 31
101	If the GnRH agonist trigger with triptorelin is applied, dosages ranging of 0.1-0.4mg can be chosen. [2019]	GPP		
102	The addition of a GnRH agonist to hCG as a dual trigger for final oocyte maturation is probably not recommended for predicted normal responders. [2019]	Conditional	⊕⊕○○	SoF table 32 a,b
103	The addition of a GnRH agonist to hCG as a dual trigger for final oocyte maturation is probably not recommended for low responders. [2025]	Conditional	⊕⊕○○	SoF table 32 c
/	There is too limited evidence to draw conclusions on the use of double trigger for final oocyte maturation for IVF/ICSI.	/	/	SoF table 33

Luteal phase support

104	Progesterone is recommended for luteal phase support after IVF/ICSI. [2019]	Strong	⊕○○○	SoF table 34
105	Any of the previously mentioned administration routes (non-oral) for natural progesterone as luteal phase support can be used. [2019]	GPP		
106	The dosing of natural progesterone has evolved empirically, usually dosages used include: 50 mg once daily for intramuscular progesterone 25 mg once daily for subcutaneous progesterone 90 mg once daily for vaginal progesterone gel 200 mg three times daily for micronized vaginal progesterone in-oil capsules 100 mg two or three times daily for micronized vaginal progesterone in starch suppositories 400 mg two times daily for vaginal pessary. [2019]	GPP		SoF table 35 a,b,c,d
107	Starting of progesterone for luteal phase support should be in the window between the evening of the day of oocyte retrieval and day 3 post oocyte retrieval. [2019]	GPP		SoF table 36 a,b,c
108	Progesterone support should be administered until at least the day of the pregnancy test. [2019]	GPP		SoF table 37
109	Dydrogesterone is probably recommended for luteal phase support. [2019] <i>There are pharmacovigilance reports of association between dydrogesterone exposure and increased risk of congenital malformations, although the observed relations cannot necessarily be translated into a conclusion on causality.</i>	Conditional	⊕⊕⊕○	SoF table 38
110	The addition of oestradiol to progesterone for luteal phase support is probably not recommended. [2019]	Conditional	⊕⊕○○	SoF table 39
111	In hCG triggered ovarian stimulation cycles, hCG as luteal phase support in standard dosages of 1500 IU is not recommended. [updated]	Strong	⊕⊕○○	SoF table 40 a,b



112	A GnRH agonist bolus, in addition to progesterone for luteal phase support in hCG triggered cycles is probably not recommended. [updated]	Conditional	⊕⊕○○	SoF table 41
113	Repeated GnRH agonist injections, alone or in addition to progesterone for luteal phase support in hCG triggered cycles is probably not recommended. [updated]	Conditional	⊕○○○	SoF table 42
114	Addition of LH to progesterone for luteal phase support can only be used in the context of a clinical trial. [2019]	Research only		SoF table 43

Part G: Prevention of OHSS

115	A GnRH agonist trigger is recommended for final oocyte maturation in women at risk of OHSS combined with a freeze-all strategy to minimise the risk of severe OHSS. [updated]	Strong	⊕○○○	SoF table 44 a,b,c
116	If a GnRH agonist protocol with hCG trigger is used in high responders, a freeze-all strategy is recommended to decrease the risk of late-onset OHSS. [updated]	GPP		
117	The addition of hCG to GnRH agonist as a dual trigger for final oocyte maturation is probably not recommended for high responders. [2025]	Conditional	⊕○○○	
118	In patients at risk of OHSS, the use of a GnRH agonist for final oocyte maturation is probably recommended over hCG in cases where no fresh transfer is performed. [2019]	Conditional	⊕○○○	
119	A GnRH agonist trigger for final oocyte maturation with or without a freeze-all strategy is preferred over a coasting strategy in patients at risk of OHSS. [2019]	GPP		
120	Dopamine agonists are recommended to decrease the risk of early OHSS, particularly in patients receiving hCG for final oocyte maturation. [updated]	Strong	⊕⊕○○	SoF table 45
121	A freeze-all strategy is recommended to minimise the risk of late-onset OHSS. [updated]	Strong	⊕⊕○○	SoF table 46
122	Prior to start of ovarian stimulation, a risk assessment for high response is advised with the purpose of applying personalised treatment choices on pituitary suppression protocol, FSH dosage, final oocyte maturation trigger and embryo transfer strategy. [updated]	GPP		

466



PART A: Pre-stimulation evaluation

467

468

1. Ovarian response prediction

469

PICO QUESTION: IS THE ASSESSMENT OF THE PREDICTED RESPONSE TO OVARIAN STIMULATION SUFFICIENTLY RELIABLE?

470 Implications following the prediction of an extremely ovarian response is relevant for both the clinicians
471 and patients. Clinicians may suggest personalizing the treatment based on that prediction, and such
472 strategies will be discussed elsewhere in this guideline. For the patients, ovarian response prediction
473 provides information about the chances of success, the safety risks and complications.

474 ANTRAL FOLLICLE COUNT (AFC)

475 Evidence

476 *A high number of studies have investigated the role of AFC in the prediction of ovarian response to*
477 *ovarian stimulation. Most of these studies have a limited number of patients, and the definition of low*
478 *and high response has not been uniform. AFC has been studied in GnRH agonist and antagonist cycles*
479 *and in patients stimulated with different dosages and protocols of FSH. Also, several narrative reviews*
480 *and meta-analyses have been conducted on the subject.*

481 A systematic review and meta-analysis¹ investigated the performance of the AFC to predict a high (6
482 studies) and low (15 studies) response to ovarian stimulation (Liu et al., 2023). To predict high response,
483 the overall pooled sensitivity of AFC was 0.83 (95% CI 0.77-0.87) and pooled specificity 0.78 (95% CI
484 0.64-0.88). High heterogeneity was present. The AUC for the predictive value of AFC for a high response
485 to ovarian stimulation was 0.87 (95% CI 0.84-0.89). To predict low ovarian response, the overall pooled
486 sensitivity was 0.75 (95% CI 0.67-0.81) and pooled specificity was 0.82 (95% CI 0.76-0.87). Again, high
487 heterogeneity was found for both. The AUC for the predictive value of AFC for a low response to ovarian
488 stimulation was 0.85 (95% CI 0.82-0.88).

489 Several studies were identified assessing the predictive accuracy for AFC in ovarian response prediction
490 which were not included in the meta-analysis or were published afterwards, which show similar results
491 to the meta-analysis (Arce et al., 2013, Bancsi et al., 2002, Elgindy et al., 2008, Hochberg et al., 2024,
492 Jayaprakasan et al., 2009, Khairy et al., 2008, Kwee et al., 2007, Lan et al., 2013, Lee et al., 2020,
493 Oehninger et al., 2015, Penarrubia et al., 2010, Sun et al., 2022, Tsakos et al., 2014, Wang et al., 2021).

494

¹ The IPD meta-analyses by Broer et al 2013a and b are replaced by a more recent meta-analysis. The cohort studies by Bancsi et al., 2004, Jayaprakasan et al., 2010, Mutlu et al., 2013, Soldevila et al., 2007, Tolikas et al., 2011 are included in the meta-analysis.



495 Table 1: Accuracy of AFC in predicting ovarian response.

AFC Study	Cohort (n)	High ovarian response		Low ovarian response		Remark
		Criterion	ROC-AUC	Criterion	ROC-AUC	
Liu 2023	7190		0.90		0.87	
Other studies:						
Hochberg 2024	4220	≥15 oocytes	0.80			
Sun 2022	2585	>15 oocytes	0.73	≤3 oocytes	0.92	
Wang 2021	84884			≤3 oocytes	0.84	
Lee 2020	263	≥ 20 oocytes	0.81	< 4 oocytes	0.82	
Oehninger 2015	686	>18 oocytes	0.88	<6 oocytes	0.88	
Tsakos 2014	105	>12 oocytes	0.86	<4 oocytes	0.86	
Lan 2013	382	>20 oocytes	0.81	≤3 oocytes	0.80	
Arce 2013	375	≥15 oocytes	0.64	≤3 oocytes	0.74	rFSH stimulation
Arce 2013	374	≥15 oocytes	0.65	≤3 oocytes	0.67	hMG stimulation
Penarrubia 2010	98			≤3 oocytes	0.90	
Jayaprakasan 2009	141			<4 oocytes	0.89	
Khairy 2008	148			<4 oocytes	0.79	
Elgindy 2008	33			<4 oocytes	0.94	
Kwee 2007	110	>20 oocytes	0.92	<6 oocytes	0.83	
Bancsi 2002	120			<4 oocytes	0.87	

496 Conclusion

497 The prediction of ovarian response categories by AFC alone is reliable.

498 **ANTI-MÜLLERIAN HORMONE (AMH)**

499 Evidence

500 A high number of studies have investigated the role of AMH in the prediction of ovarian response to
 501 ovarian stimulation. Most of these studies have a limited number of patients, and studies have used
 502 different assays for the measurement of the AMH values. AMH has been studied in GnRH agonist and
 503 antagonist cycles and in patients stimulated with different dosages and protocols of FSH. Moreover, the
 504 definition of a low and high response has not been uniform, which nevertheless showed AMH to be a
 505 good predictor of ovarian response. Several narrative reviews have been written next to different meta-
 506 analyses on the subject.

507 A systematic review and meta-analysis² investigated the performance of AMH to predict a high (13
 508 studies) and low (29 studies) response to ovarian stimulation (Liu, et al., 2023). To predict high
 509 response, the overall pooled sensitivity of AMH was 0.79 (95% CI 0.74-0.83) and pooled specificity 0.79
 510 (95% CI 0.74-0.83). The AUC for the predictive value of AMH for a high response to ovarian stimulation
 511 was 0.86 (95% CI 0.82-0.89). To predict low ovarian response, the overall pooled sensitivity was 0.78

² The IPD meta-analyses by Broer et al 2013a and b are replaced by a more recent meta-analysis. The cohort studies by Heidar et al., 2015, Jayaprakasan et al., 2010, Li et al., 2016, Mutlu et al., 2013, Tolikas et al., 2011 are included in the meta-analysis.



512 (95% CI 0.74-0.80) and pooled specificity was 0.79 (95% CI 0.76-0.83). High heterogeneity was found
 513 for both. The AUC for the predictive value of AMH for a low response to ovarian stimulation was 0.85
 514 (95% CI 0.81-0.88).

515 Several studies were identified assessing the predictive accuracy for AMH in ovarian response
 516 prediction which were not included in the meta-analysis or were published afterwards, which show
 517 similar results (Andersen et al., 2011, Arce et al., 2013, Bosch et al., 2023, Elgindy et al., 2008, Hochberg
 518 et al., 2024, Huang et al., 2019, Lan et al., 2013, Lee et al., 2020, Oehninger et al., 2015, Sun et al., 2022,
 519 Tsakos et al., 2014).

520 *Table 2: Accuracy of AMH in predicting ovarian response.*

AMH Study	Cohort (n)	High ovarian response		Low ovarian response		Remark
		Criterion	ROC-AUC	Criterion	ROC-AUC	
Liu 2023	7190		0.89		0.87	
Other studies:						
Hochberg 2024	4220	≥15 oocytes	0.71			
Bosch 2023	1248	>15 oocytes	0.89	≤3 oocytes	0.85	
Sun 2022	2585	>15 oocytes	0.73	≤3 oocytes	0.79	
Wang 2021	41702			≤3 oocytes	0.86	
Lee 2020	263	≥ 20 oocytes	0.80	< 4 oocytes	0.85	
Huang 2019	523	>15 oocytes	0.77	< 4 oocytes	0.86	
Oehninger 2015	686	>18 oocytes	0.86	<6 oocytes	0.87	
Tsakos 2014	105	>12 oocytes	0.66	<4 oocytes	0.63	
Arce 2013	374	≥15 oocytes	0.77	≤3 oocytes	0.78	hMG stimulation
Arce 2013	375	≥15 oocytes	0.81	≤3 oocytes	0.90	rFSH stimulation
Lan 2013	382	>20 oocytes	0.76	≤3 oocytes	0.88	
Andersen 2011	442	>18 oocytes	0.77	<6 oocytes	0.84	
Elgindy 2008	33			<4 oocytes	0.90	

521 Conclusion

522 The prediction of ovarian response categories by AMH alone is reliable.

523 BASAL FOLLICLE STIMULATING HORMONE (FSH)

524 Evidence

525 *A high number of studies have investigated the role of basal FSH levels in the prediction of ovarian*
 526 *response to ovarian stimulation. Most of these studies have a limited number of patients, and the*
 527 *definition of a low and high response has not been uniform. Also, several narrative reviews and meta-*
 528 *analyses have been conducted on the subject.*

529 An IPD meta-analysis assessed the accuracy of basal FSH and reported moderate accuracy of basal FSH
 530 in predicting both a low response (ROC-AUC of 0.66 (95% CI 0.62-0.69) and an excessive response (ROC-
 531 AUC of 0.64 (95% CI 0.61-0.67)) (Broer et al., 2013a, Broer et al., 2013b).

532 Several studies were identified assessing the predictive accuracy for basal FSH in ovarian response
 533 prediction which were not included in the IPD meta-analysis or were published afterwards, which show



534 similar results to the IPD meta-analyses (Arce et al., 2013, Bancsi et al., 2002, Elgindy et al., 2008,
 535 Jayaprakasan et al., 2009, Khairy et al., 2008, Kwee et al., 2007, Lee et al., 2020, Mutlu et al., 2013,
 536 Oehninger et al., 2015, Penarrubia et al., 2010, Soldevila et al., 2007, Tolikas et al., 2011, Tsakos et al.,
 537 2014, Wang et al., 2021).

538 *Table 3: Accuracy of basal FSH in predicting ovarian response.*

Study	Cohort (n)	High ovarian response		Low ovarian response		Remark
		Criterion	ROC-AUC	Criterion	ROC-AUC	
Broer 2013a/b	4786/5705	>15 oocytes	0.64	≤4 oocytes	0.66	
Other studies:						
Wang 2021	85052			≤3 oocytes	0.69	
Lee 2020	263	≥ 20 oocytes	0.63	< 4 oocytes	0.73	
Oehninger 2015	686	>18 oocytes	0.88			
Tsakos 2014	105	>12 oocytes	0.72	<4 oocytes	0.67	
Arce 2013	374	≥15 oocytes	0.71	≤3 oocytes	0.73	hMG stimulation
Arce 2013	375	≥15 oocytes	0.73	≤3 oocytes	0.72	rFSH stimulation
Mutlu 2013	192			<4 oocytes	0.75	
Tolikas 2011	90			<4 oocytes	0.65	
Penarrubia 2010	98			≤3 oocytes	0.62	
Jayaprakasan 2009	141			<4 oocytes	0.69	
Elgindy 2008	33			<4 oocytes	0.85	
Khairy 2008	148			<4 oocytes	0.69	
Kwee 2007	110	>20 oocytes	0.80	<6 oocytes	0.83	
Soldevila 2007	327			≤5 oocytes	0.63	
Bancsi 2002	120			<4 oocytes	0.84	

539 **Conclusion**

540 The prediction of ovarian response categories by basal FSH alone is not sufficiently reliable, compared
 541 to the predictive accuracy by the AFC and AMH.

542 **INHIBIN B**

543 **Evidence**

544 A high number of studies has investigated the role of inhibin B in the prediction of ovarian response to
 545 ovarian stimulation (OS). In 2006, a systematic review and meta-analysis (9 studies, 788 cycles) has
 546 been performed including inhibin B (Broekmans et al., 2006). Although variations between studies
 547 regarding definition of poor response, study quality and study characteristics existed, statistical analysis
 548 showed these not related to the predictive performance of inhibin B. The sensitivity of inhibin B in the
 549 prediction of a poor response ranged from 32 to 89%, the specificity ranged from 29 to 95%. The
 550 spearman correlation coefficient for sensitivity and specificity was -0.93. From logistic regression the
 551 pre- and post-test probabilities of a poor response were calculated. These demonstrated that inhibin B
 552 has a modest accuracy in the prediction of a poor response (Broekmans et al., 2006).



553 Since the publication of this meta-analysis a few more studies have been published assessing the
 554 predictive accuracy for inhibin B in ovarian response prediction (Arce et al., 2013, Fawzy et al., 2002,
 555 Hendriks et al., 2005, Kwee et al., 2007, Penarrubia et al., 2010, van Rooij et al., 2002).

556 *Table 4: Accuracy of Inhibin B in predicting ovarian response.*

Inhibin B Study	Cohort (n)	High ovarian response		Low ovarian response		Remark
		Criterion	ROC-AUC	Criterion	ROC-AUC	
Arce 2013	374	≥15 oocytes	0.60	≤3 oocytes	0.62	hMG stimulation
Arce 2013	375	≥15 oocytes	0.53	≤3 oocytes	0.64	rFSH stimulation
Penarrubia 2010	98			≤3 oocytes	0.61	
Kwee 2007	110	>20 oocytes	0.93	<6 oocytes	0.86	for the increment of inhibin B in the EFORT
Hendriks 2005	63			<4 oocytes	0.76	
Fawzy 2002	54			<8 MII oocytes	0.96	
Van Rooij 2002	119			<4 oocytes	0.76	

557

558 Conclusion

559 The prediction of ovarian response categories by inhibin B alone is not sufficiently reliable.

560 BASAL OESTRADIOL

561 Evidence

562 Basal oestradiol has also been studied as a predictor of ovarian response to ovarian stimulation. The
 563 systematic review by Broekmans et al., mentioned before, also investigated the performance of basal
 564 oestradiol in predicting ovarian response (10 studies, 3911 women) (Broekmans et al., 2006). The
 565 sensitivity of basal oestradiol in the prediction of a poor response ranged from 3 to 83%, the specificity
 566 ranged from 13 to 98%. The spearman correlation coefficient for sensitivity and specificity was -0.50.
 567 From LR the pre- and post-test probability of a poor response was calculated. This demonstrated that
 568 basal oestradiol has a low accuracy in the prediction of a poor response (Broekmans et al., 2006).

569 Since the publication of this meta-analysis, a few more studies have been published assessing the
 570 predictive accuracy for basal oestradiol in ovarian response prediction (Hendriks et al., 2005, Khairy et
 571 al., 2008, Kwee et al., 2007, Lee et al., 2020, Penarrubia et al., 2010, van Rooij et al., 2002). These have
 572 confirmed the low accuracy of basal oestradiol.

573 *Table 5: Accuracy of basal oestradiol in predicting ovarian response.*

basal estradiol Study	Cohort (n)	High ovarian response		Low ovarian response		Remark
		Criterion	ROC-AUC	Criterion	ROC-AUC	
Lee 2020	263	≥ 20 oocytes	0.52	< 4 oocytes	0.66	
Penarrubia 2010	98			≤3 oocytes	0.55	
Khairy 2008	148			<4 oocytes	0.51	
Kwee 2007	110	>20 oocytes	0.83	<6 oocytes	0.75	for the increment of basal oestradiol in the EFORT
Hendriks 2005	63			<4 oocytes	0.54	
Van Rooij 2002	119			<4 oocytes	0.52	



574 **Conclusion**

575 Basal oestradiol alone is not a predictor of ovarian response.

576 **BASAL PROGESTERONE**

577 **Evidence**

578 *No studies were retrieved investigating the role of basal progesterone in the prediction of ovarian response to ovarian stimulation in terms of sensitivity, specificity and AUC.*

580 **BASAL LH**

581 **Evidence**

582 *No studies were retrieved investigating the role of basal progesterone in the prediction of ovarian response to ovarian stimulation in terms of sensitivity, specificity and AUC.*

584 **AGE**

585 **Evidence**

586 *A high number of studies have investigated the role of age in the prediction of ovarian response to ovarian stimulation. Most of these studies have a limited number of patients, and the definition of low and high response has not been uniform. However, all these studies show an unsatisfactory ROC curve for age as predictor of ovarian response. Several meta-analyses have been conducted on the subject.*

590 The IPD meta-analyses mentioned earlier also assessed the accuracy of age and reported a limited accuracy of age alone in predicting both a poor response (ROC-AUC of 0.60 (95% CI 0.57-0.64)) and an excessive response (ROC-AUC of 0.61 (95% CI 0.58-0.64)) (Broer, et al., 2013a, Broer, et al., 2013b).

593 Several studies were identified assessing the predictive accuracy for age in ovarian response prediction which were not included in the IPD meta-analysis or were published afterwards (Bancsi et al., 2002, Jayaprakasan et al., 2009, Khairy et al., 2008, Kwee et al., 2007, Lee et al., 2020, Mutlu et al., 2013, Oehninger et al., 2015, Penarrubia et al., 2010, Wang et al., 2021).

597 *Table 6: Accuracy of age in predicting ovarian response.*

Age	High ovarian response		Low ovarian response		
	Study	Cohort (n)	Criterion	ROC-AUC	Criterion
Broer 2013a/b	4786/5705	>15 oocytes	0.61	≤4 oocytes	0.60
Other studies:					
Sun 2022	2585	>15 oocytes	0.65	≤3 oocytes	0.75
Wang 2021	88987			≤3 oocytes	0.72
Lee 2020	263	≥ 20 oocytes	0.65	< 4 oocytes	0.68
Oehninger 2015	686	>18 oocytes	0.55	<6 oocytes	0.55
Mutlu 2013	192			<4 oocytes	0.76
Penarrubia 2010	98			≤3 oocytes	0.75
Jayaprakasan 2009	141			<4 oocytes	0.74
Khairy 2008	148			<4 oocytes	0.71
Kwee 2007	110	>20 oocytes	0.71	<6 oocytes	0.63
Bancsi 2002	120			<4 oocytes	0.61



598

599 **Conclusion**

600 The prediction of ovarian response categories by age alone is not sufficiently reliable.

601 **BODY MASS INDEX (BMI)**

602 **Evidence**

603 With the growing interest for ovarian response prediction, the role of BMI in ovarian response has been
604 questioned. However, there are only a few studies actually assessing the accuracy of BMI as a predictor
605 of ovarian response. In these studies, BMI was found to have a small to no predictive accuracy for
606 ovarian response to ovarian stimulation.

607 The IPD meta-analyses mentioned earlier also assessed the accuracy of BMI and concluded that BMI
608 was not a significant predictor of ovarian response, neither for poor nor a high response (Broer, et al.,
609 2013a, Broer, et al., 2013b).

610 *Table 7: Accuracy of BMI in predicting ovarian response.*

BMI	Cohort (n)	High ovarian response		Low ovarian response	
		Criterion	ROC-AUC	Criterion	ROC-AUC
Broer 2013a/b	4786/5705	>15 oocytes		≤4 oocytes	
Other studies:					
Sun 2022	2585	>15 oocytes	0.51	≤3 oocytes	0.58
Lee 2020	263	≥ 20 oocytes	0.52	< 4 oocytes	0.54
Khairy 2008	148			<4 oocytes	0.68

611

612 **Conclusion**

613 BMI alone is not a predictor of ovarian response.

614 **OVERALL RECOMMENDATION**

615 **Evidence**

616 Based on the available evidence both AFC and AMH show a high accuracy in the predication of a low
617 and high response (Table 1 and 2). The accuracy of Basal FSH and Inhibin B levels is moderate (Table 3
618 and 4). Basal oestradiol, age and BMI are not good predictors of ovarian response to hyperstimulation
619 (Table 5, 6 and 7).

620 **Recommendation**

For predicting high and low response to ovarian stimulation, use of either antral follicle count (AFC) or anti-Müllerian hormone (AMH) is recommended. [updated]	Strong ⊕○○○
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621



Age, BMI, basal FSH, inhibin B and basal oestradiol are not recommended for the prediction of ovarian response. [2025]

Strong ⊕○○○

622

623 Justification

624 AFC and AMH both have a high accuracy in the prediction of ovarian response category (high or low).
625 Taking into account false positive and negative rate of the test it may be recommended for clinical
626 application. The clinician can decide which test is most appropriate for their clinical setting.

627 In this guideline, we did not compare AMH and AFC with each other nor studied the added effect of
628 using both tests for ovarian response prediction. However, the IPD meta-analysis did demonstrate that
629 these tests do have added value to female age alone. Moreover, there was no difference in the
630 performance of these tests and combining them did not improve the prediction of ovarian response
631 (Broer, et al., 2013a, Broer, et al., 2013b).

632 Basal FSH and inhibin B do have some predictive value for ovarian response, however for an accurate
633 prediction very high cut-off levels need to be used. This implies that only very few women will have
634 such an abnormal FSH or Inhibin B test results. This results in hardly any clinical value, especially since
635 there are other tests available with a higher accuracy. Age also has some predictive value, however
636 assessment of ovarian response category by age alone is not sufficiently reliable. Basal oestradiol and
637 BMI alone are not predictors of ovarian response. Therefore, we recommend not using basal FSH,
638 inhibin B, basal oestradiol, age or BMI for the prediction of ovarian response.

639 As all original studies have been performed using different assays or ranges for AFC and AMH, it is not
640 possible to combine these data to calculate cut-offs for the prediction of a low or high response.
641 Regarding the use of AMH and AFC for individualised gonadotropin dose selection, the reader is
642 referred to the Cochrane review by Lensen et al. since this was not investigated in this guideline (Lensen
643 et al., 2017).

644

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732



733 2. Pregnancy prediction

PICO QUESTION: WHAT IS THE PROGNOSTIC VALUE OF HORMONAL ASSESSMENT AT BASELINE?

734 ANTRAL FOLLICLE COUNT (AFC)

735 Evidence

736 In an IPD meta-analysis, including 55 study reports, AFC had no predictive effect for ongoing pregnancy
737 after IVF (AUC 0.50, 95% CI 0.40-0.59) (Broer et al., 2013).

738 Conclusion

739 AFC alone is not a predictor for the outcome pregnancy.

740 ANTI-MÜLLERIAN HORMONE (AMH)

741 Evidence

742 In an IPD meta-analysis, including 55 study reports, AMH had only a very low predictive value for
743 ongoing pregnancy after IVF (AUC 0.55, 95% CI 0.45-0.64) (Broer et al., 2013).

744 In a prospective cohort study, the relationship between AMH levels and pregnancy outcomes was
745 investigated in 50 patients undergoing ovarian stimulation for IVF/ICSI (Umarsingh et al., 2020). The
746 studied population was divided into low to normal AMH (0.3-0.9 ng/mL; n=3), normal AMH (<1 ng/mL;
747 n=17) and high AMH (<3 ng/mL; n=22). Pregnancy rates were 27.3% (6/22) in the high AMH group,
748 35.3% (6/17) in the normal AMH group and 0% (0/3) in the low to normal AMH group. AUC to predict
749 pregnancy outcomes of AMH was 0.497.

750 In a prospective cohort study, the possible association between AMH and clinical outcomes in IVF cycles
751 was investigated in 124 patients undergoing their first ovarian stimulation cycle (Li et al., 2015). No
752 direct correlation was observed between serum AMH and inhibin B levels on day 2/3 and clinical
753 pregnancy.

754 In a prospective cohort study, it was investigated if AMH level on day 3 could predict reproductive
755 outcomes in 164 women with PCOS undergoing their first IVF treatment cycle (Xi et al., 2012). The
756 studies population was divided into low AMH (≤ 4.85 ng/mL; n=41), average AMH (4.85-8.82 ng/mL;
757 n=82) and high AMH (≥ 8.82 ng/mL; n=41). No significant difference in clinical pregnancy rate was
758 observed with low, average, or high AMH (65% (26/40) vs. (66.7% (50/75) vs. 45.9% (17/37)).

759 In a prospective cohort study, the possible relationship between AMH levels on day 3 and reproductive
760 outcomes was investigated in 60 women with PCOS (80 cycles) (Kaya et al., 2010). The studied
761 population was divided according to the <25th (21 cycles), 25-75 (39 cycles) and >75th percentile (20
762 cycles) of serum AMH on day 3. The clinical pregnancy rate increased significantly with AMH levels
763 (33.3% (7/21) vs. 46.1% (19.39) vs. 60% (12/20)). For predicting clinical pregnancy rates, using a cutoff
764 value of 3.2 ng/mL, the sensitivity was 72.7% and the specificity 77.3%.



765 **Conclusion**

766 AMH alone is not a predictor of the outcome pregnancy.

767 **BASAL FOLLICLE STIMULATING HORMONE (FSH)**

768 **Evidence**

769 In an IPD meta-analysis, including 55 study reports, basal FSH had only a very low predictive value for
770 ongoing pregnancy after IVF (AUC 0.53, 95% CI 0.43-0.62) (Broer, et al., 2013).

771 In a large retrospective cohort study, including 19682 cycles, the relationship between early follicular
772 FSH levels and oestradiol levels and reproductive outcomes was investigated (Frazier et al., 2004). In
773 the final model for live birth delivery, statistically significant negative predictors included increasing
774 age, elevated FSH ratio, elevated oestradiol ratio.

775 **Conclusion**

776 Basal FSH alone is not a predictor of the outcome pregnancy.

777 **INHIBIN B**

778 **Evidence**

779 In a systematic review and meta-analysis, including 3 study reports, ROC curves were estimated for the
780 predictive accuracy of inhibin B for non-pregnancy (Broekmans et al., 2006). Extreme threshold levels
781 were necessary to obtain a modest positive likelihood ratio of ~4–5, resulting in a post-test pregnancy
782 rate of approximately 5%. Such abnormal test results occur only in a very limited number of patients.

783 In a prospective cohort study, the possible association between AMH and clinical outcomes in IVF cycles
784 was investigated in 124 patients undergoing their first ovarian stimulation cycle (Li et al., 2015). No
785 direct correlation was observed between inhibin B levels on day 2/3 and clinical pregnancy.

786 **Conclusion**

787 Inhibin B alone is not a predictor of the outcome pregnancy.

788 **BASAL OESTRADIOL**

789 Assessment of oestradiol at initiation of stimulation is frequently performed in IVF/ICSI and an elevated
790 level usually signifies the presence of a simple follicular cyst, which is then confirmed at ultrasound.
791 However, prediction of the outcome of stimulation has also been attempted using E2 level at initiation
792 of stimulation.

793 **Evidence**

794 In a systematic review and meta-analysis, including 9 study reports, ROC curves were estimated for
795 the predictive accuracy of oestradiol for non-pregnancy (Broekmans, et al., 2006). For prediction of
796 non-pregnancy no clear threshold levels could be identified for that would lead to an adequate
797 combination of LR, post-test probability and abnormal test rate.



798 In a large retrospective cohort study, including 19682 cycles, the relationship between early follicular
799 FSH levels and oestradiol levels and reproductive outcomes was investigated (Frazier, et al., 2004). In
800 the final model for live birth delivery, statistically significant negative predictors included the
801 combination of increasing age, elevated FSH ratio, elevated oestradiol ratio.

802 One retrospective study in patients with unexplained infertility undergoing ovarian stimulation and
803 intercourse shows a significantly lower chance of pregnancy in women with higher oestradiol levels at
804 initiation of stimulation (Costello et al., 2001).

805 Conclusion

806 Oestradiol alone is not a predictor of the outcome pregnancy.

807 BASAL PROGESTERONE

808 In a proportion of cycles, progesterone remains elevated at menstruation. Elevated progesterone levels
809 at the intended starting date of ovarian stimulation could be associated with reduced pregnancy rates.
810 The proportion of patients with progesterone levels >1.6 ng/ml on cycle day 2 was 4.9% (95% CI 3.2-
811 7.4) in a cohort study by Kolibianakis et al. (2004) and 6.2% (95% CI 4-9) in a cohort study by Blockeel
812 et al. (Blockeel et al., 2011, Kolibianakis et al., 2004). A more recent study by Hamdine et al. reported
813 13.3% (95% CI 8-20) of patients with progesterone levels >1.5 ng/ml. Faulisi et al. reported 0.3% (95%
814 CI 0.01-1.15) of patients with progesterone levels >1.6 ng/ml on cycle day 3 (Faulisi et al., 2017,
815 Hamdine et al., 2014). Due to the low incidence it seems unnecessary to evaluate this research question
816 for progesterone levels >1.6 ng/ml on cycle day 3.

817 Evidence

818 A meta-analysis³, including 3 cohort studies and 773 women, investigated the effect of elevated
819 progesterone levels at baseline on reproductive outcomes (Lim et al., 2024). No significant difference
820 was found for live birth rate with elevated progesterone levels at baseline at threshold level >1.5
821 ng/mL (OR 0.76, 95% CI 0.39–1.49, 2 studies, N=309). Similarly, no significant difference was found for
822 clinical pregnancy rate at threshold level >0.65 ng/mL (OR 1.41, 95% CI 0.93–2.13, 1 study, n=464) or
823 threshold level >1.5 ng/mL (OR 0.81, 95% CI 0.38-1.71, 2 studies, n=309).

824 A retrospective cohort study (418 women, 461 cycles) reported lower live birth rates of 18.2% (2/11)
825 and 16.7% (1/6) with progesterone < or >1.5 ng/mL on hCG day respectively, in patients with elevated
826 (>1.5 ng/mL) levels at the start of ovarian stimulation, compared to 33.8% in controls (progesterone
827 <1.5 ng/mL both at the start of OS and on hCG day) (Panaino et al., 2017).

828 Fausili et al. showed that progesterone assessment on day 3 of stimulation is inaccurate in predicting
829 clinical pregnancy (ROC-AUC 0.54, 95%CI 0.47-0.61) (Faulisi et al., 2017).

830 Conclusion

831 Assessment of progesterone prior to initiation of stimulation on cycle day 2 in women undergoing
832 ovarian stimulation with GnRH antagonist and gonadotrophins may be beneficial to identify cases

³ The meta-analysis by Hamdine et al., 2014 cited here in the 2019 version of the guideline was replaced by a more recent meta-analysis.



833 with a lower than normal probability of pregnancy. The currently available evidence, however, is not
834 solid, and the clinical value of this test was not assessed.

835 **BASELINE LH**

836 **Evidence**

837 In a retrospective cohort study, the effect of elevated basal LH levels on reproductive outcomes was
838 investigated in poor, normal and high responders (Zhang et al., 2024). Women were divided in two
839 groups based on their baseline LH levels: <5 IU/L and ≥ 5 IU/L. OHSS rate was significantly lower in poor
840 responders with low baseline LH levels (0% (0/270) vs. 2.6% (4/157). No significant difference in OHSS
841 was observed for normal and high responders with LH levels below or above the threshold of 5 IU/L.
842 No significant difference in clinical pregnancy rates were observed in poor responders (50.0% (134/270)
843 vs. 47.8% (75/157), normal responders (58.0% (196/338) vs. 53.5% (124/232) or high responders
844 (59.6% (87/146) vs. 68.9% (173/251)) with LH levels below or above the threshold.

845 In a retrospective cohort study, the effect of elevated basal LH levels on reproductive outcomes after
846 IVF/ICSI was assessed in women with PCOS (Liu and Wang, 2023). Women were divided into two group
847 based on basal LH levels, i.e. high basal LH ($LH \geq 12.455$ IU/L; $n=59$) and low basal LH ($LH < 12.455$ IU/L,
848 $n=176$). Comparing the results of women with high and low basal LH, no significant difference was
849 observed in cumulative live birth rate (61.82% (34/55) vs. 60% (99/165) or incidence of OHSS (3.39%
850 (2/59) vs. 1.14% (2/176).

851 In a retrospective cohort study, the effect of elevated basal LH levels on reproductive outcomes after
852 IVF/ICSI was assessed in women with PCOS (Wang et al., 2022). Women were divided into three groups
853 based on basal LH, i.e. ≤ 5 mIU/mL ($n=65$), 5-10 mIU/mL ($n=54$) and ≥ 10 mIU/mL ($n=23$). Comparing the
854 results of women with ≤ 5 mIU/mL, 5-10 mIU/mL and ≥ 10 mIU/mL, no significant differences were found
855 for cumulative live birth rate (23.08 (15/65) vs. 31.48% (17/54) vs. 17.39% (4/23)).

856 In another retrospective cohort study, the effect of elevated basal LH levels on reproductive outcomes
857 after IVF/ICSI was assessed in women with PCOS (Sun et al., 2018). Women were divided into categories
858 based on basal LH, i.e; < 5 mIU/mL ($n=575$), between 5 and 7.5 mIU/mL ($n=216$), between 7.5 and 10
859 mIU/mL ($n=115$), and ≥ 10 mIU/mL ($n=105$). The number of metaphase II oocytes was significantly
860 higher in the group with basal $LH \geq 10$ mIU/mL than the groups with basal LH between 7.5 and
861 10mIU/mL, basal LH between 5 and 7.5 mIU/mL, and basal LH < 5 mIU/mL (17.18 ± 9.60 vs. 13.47 ± 9.38
862 vs. 13.97 ± 8.65 vs. 11.10 ± 7.24). The number of MII oocytes retrieved was positively correlated with the
863 basal LH level ($r=0.261$). However, no significant difference was seen in clinical pregnancy rates
864 between the different groups of basal LH (47.7% (288/604) vs. 46.5% (112/241) vs. 58.8% (70/119) vs.
865 55.5% (61/110).

866 In a retrospective study, the possible influence of endogenous LH concentrations on ongoing pregnancy
867 rates were investigated (Doody et al., 2010). Patients were stratified into the 25th, 25-75, and 75th
868 percentiles of serum LH concentrations. The ongoing pregnancy rates were not significantly different
869 in women with low, normal or high LH levels on day 1 (36.8% (29.6-44.4) vs. 36.8% (31.7-42.1) vs. 37.9%
870 (30.7-45.6)).



871 **Conclusion**

872 Most studies divided patients into categories based on basal LH levels. However, none of the included
873 studies reported significant differences in the outcome pregnancy across LH level categories.

874 **AGE**

875 **Evidence**

876 In an IPD meta-analysis, of all patient characteristics, female age alone was the strongest predictor of
877 ongoing pregnancy (OR 0.94, 95% CI 0.89-0.99) (Broer et al., 2013).

878 **Conclusion**

879 Female age alone is a predictor of the outcome pregnancy.

880 **BODY MASS INDEX (BMI)**

881 **Evidence**

882 In an IPD meta-analysis, among patient characteristics, BMI was significantly associated with ongoing
883 pregnancy (OR 0.91, 95% CI 0.85-0.97). In a multivariable model, only BMI added any predictive value
884 to age (Broer et al., 2013).

885 **Conclusion**

886 BMI alone is a predictor of the outcome pregnancy.

887 **OVERALL RECOMMENDATION**

888 **Evidence**

889 Based on the available evidence only female age and BMI are predictors of pregnancy and live birth.
890 The accuracy of AFC, AMH, basal FSH, basal LH, basal oestradiol, basal progesterone and inhibin B levels
891 are slight to not predictive for pregnancy and live birth.

892 **Recommendation**

AFC, AMH, basal FSH, basal LH, basal oestradiol, basal progesterone and inhibin B are not recommended for the prediction of pregnancy and live birth. [updated]	Strong ⊕○○○
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893

Female age and BMI are predictors of pregnancy and live birth. [2025]	Strong ⊕○○○
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894



895 Justification

896 The IPD meta-analysis and the systematic review show that only female age and BMI have predictive
897 value for pregnancy and live birth (Broekmans et al., 2006, Broer et al., 2013).

898 Assessment of progesterone prior to initiation of stimulation on cycle day 2 in women undergoing
899 ovarian stimulation with GnRH antagonist and gonadotrophins may be beneficial to identify cases with
900 a lower than normal probability of pregnancy. The currently available evidence, however, is not solid,
901 and the clinical value of this test was not assessed. The necessity of progesterone testing is dubious due
902 to the very low incidence of abnormal test results. Moreover, as a diagnostic test it has no meaningful
903 and evidence-based link to a change of the treatment strategy, in order to undo the potential negative
904 effect on prognosis. Also, cycle cancellation or delaying stimulation initiation has not been shown to
905 improve clinical outcomes. However, since a blood test is required at initiation of stimulation (cycle day
906 2), progesterone assessment can be incorporated in the patient evaluation prior to FSH administration.
907 The recommendation is not applicable to patients >39 years of age.
908

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968



PART B. Pre-treatment therapies

969

970 3. Pre-treatment therapies

PICO QUESTION: DOES HORMONE PRE-TREATMENT IMPROVE EFFICACY AND SAFETY OF OVARIAN STIMULATION?

971 Pre-treatment therapies aim to suppress or to reduce LH and/or FSH secretion prior to gonadotrophin
972 stimulation in IVF cycles. They are used by clinicians for different purposes such as synchronisation of
973 follicular development, prevention of occurrence of early large follicle or spontaneous LH-surge,
974 reduction of cyst formation. Pre-treatment is also used for scheduling IVF cycles for the benefit of
975 clinicians and people in the laboratory as well as patients. It allows to plan IVF activity within weeks and
976 months and to avoid work on weekends and holidays. The use of pre-treatment for scheduling purpose
977 is not addressed in this guideline.

978 OESTROGEN PRE-TREATMENT

979 Evidence

980 A systematic review and meta-analysis⁴ compared reproductive outcomes for IVF/ICSI with oestrogen
981 pre-treatment compared to no pre-treatment in the GnRH antagonist protocol (Zhu et al., 2022). No
982 significant difference was found between oestrogen pre-treatment and no pre-treatment in women
983 with a normal response to ovarian stimulation for live birth rate (4 RCTs; OR 0.98; 95% CI 0.74-1.30;
984 919 women) or ongoing pregnancy rate (7 RCTs; OR 0.92; 95% CI 0.69-1.21; 1236 women).

985 A recent RCT investigated the use of steroid pre-treatment in IVF/ICSI in the GnRH antagonist protocol
986 in 52 women (Fernández-Prada et al., 2022). Estradiol valerate treatment was started on day 25 of the
987 cycle preceding at a daily dose of 2 mg/12 hours until the day before the start and not beyond 7th day
988 of cycle. No significant difference was found between oestrogen pre-treatment and no pre-treatment
989 for cumulative live birth rate (27.3% (6/22) vs. 47.6% (10/21)), live birth rate (28.6% (4/14) vs. 46.7%
990 (7/15)). There was also no significant difference in the number of MII oocytes between the study and
991 control group (5.76±3.67 vs. 6.15±4.68).

992 Two RCTs compared oestrogen pre-treatment to no pre-treatment in the GnRH antagonist protocol in
993 women experiencing a low ovarian response to stimulation (Ghasemzadeh et al., 2020, Zhang et al.,
994 2022). In the RCT by Ghasemzadeh *et al.*, oral estradiol valerate (4 mg) was initiated from the 21st day
995 of the previous IVF cycle, and continued to the second day of the cycle, the day of starting gonadotropin
996 stimulation. No significant differences were found in the number of MII oocytes between oestradiol
997 pre-treatment and no pre-treatment (3.6±0.3 vs. 2.8±0.3) (Ghasemzadeh, et al., 2020). In the RCT by
998 Zhang *et al.*, estrogen valerate was started on day 7 after ovulation at a dose of 2mg twice a day until
999 day 2 of their next menstruation. No significant difference was found between oestrogen pre-treatment

⁴ The Cochrane meta-analysis by Farquhar et al., 2017 was replaced by a more recent systematic review. The RCT by Shahrokh Tehrani Nejad et al., 2018 is included in the meta-analysis and therefore no longer included separately.



1000 and no pre-treatment for clinical pregnancy rate per first transfer (19.3 (23/276) vs. 28.7% (43/276)) or
1001 number of MII oocytes (2.9±2.5 vs. 3.1±2.4) (Zhang, et al., 2022).

1002 In an RCT, oestrogen pre-treatment was compared to no pre-treatment in the fixed GnRH antagonist
1003 protocol for women of advanced age (38-42 years) (Cédrin-Durnerin et al., 2024). Oestradiol pre-
1004 treatment started between day 20 and 24 of the previous cycle, until Wednesday evening following the
1005 onset of the menses, followed by ovarian stimulation on Friday. No significant differences were found
1006 between the pre-treatment and no pre-treatment group for cumulative live birth rate (17.7% (26/147)
1007 vs. 22.9% (33/144)), live birth rate per transfer (16.2% (16/147) vs. 18.5% (17/144)) or number of
1008 mature oocytes retrieved (7.0±5.5 vs. 7.3±5.2).

1009 Recommendation

Pre-treatment with oestrogen before ovarian stimulation using the GnRH antagonist protocol is not recommended for improving efficacy. [updated]

Strong ⊕⊕○○

1010

1011 Justification

1012 There is no evidence of a beneficial effect on live birth rate/ongoing pregnancy rate using oestrogen as
1013 pre-treatment in GnRH antagonist protocol, compared to no pre-treatment. The evidence regarding
1014 the effect of oestradiol pre-treatment on the number of oocytes retrieved is conflicting.

1015 This recommendation is not restricted to a specific group of women.

1016 PROGESTOGEN PRE-TREATMENT

1017 Evidence

1018 A Cochrane systematic review and meta-analysis investigated the effect of progesterone pre-treatment
1019 for OS in 4 RCTs including 421 women. When progestogen pre-treatment was compared with no
1020 intervention, there was no difference between the groups in live birth/ongoing pregnancy rate in GnRH
1021 agonist protocols (2 RCT, OR 1.35, 95% CI 0.69-2.65, 222 women). There was insufficient evidence to
1022 determine whether there was a difference in live birth/ongoing pregnancy rate in the GnRH antagonist
1023 protocol (1 RCT, OR 0.67, 95% CI 0.18-2.54, 47 women) (Farquhar et al., 2017).

1024 There was insufficient evidence to determine whether pre-treatment with progestogen resulted in a
1025 difference between the groups in the mean number of oocytes retrieved, both in GnRH agonist (MD -
1026 0.52, 95%CI -2.07 to 1.02, 2 RCT; and GnRH antagonist protocols (MD 2.70, 95% CI -0.98 to 6.38, 1 RCT)
1027 (Farquhar et al., 2017).

1028



1029 Recommendation

Pre-treatment with progesterone before ovarian stimulation is probably not recommended for improving efficacy. [reworded]

Conditional ⊕⊕○○

1030

Oestrogen or progesterone pre-treatment can be used for scheduling purposes given the data on efficacy and safety. [reworded]

GPP

1031

1032 Justification

1033 The available evidence indicates no beneficial effect on live birth/ongoing pregnancy rate, using
1034 progestogen as pre-treatment in GnRH agonist nor GnRH antagonist protocols. There is low quality
1035 evidence of an increased clinical pregnancy rate with progestogen pre-treatment in GnRH agonist
1036 protocols.

1037 This recommendation is not restricted to a specific group of women, although women with PCOS were
1038 excluded from the meta-analysis by Farquhar et al. (Farquhar, et al., 2017).

1039 **COMBINED ORAL CONTRACEPTIVE PILL PRE-TREATMENT**

1040 Evidence

1041 A Cochrane systematic and meta-analysis reported that in the GnRH antagonist protocol with COCP
1042 pre-treatment (12-28 days), the rate of live birth/ongoing pregnancy was lower than with no pre-
1043 treatment (OR 0.74, 95% CI 0.58-0.95, 6 RCT, 1335 women). There was no evidence of a difference
1044 between the groups in OHSS rates (OR 0.98, 95% CI 0.28-3.40, 2 RCT, 642 women) or number of oocytes
1045 (MD 0.44, 95% CI -0.11 to 0.99, 6 RCT) (Farquhar et al., 2017). In a subgroup of poor responders (80
1046 women) there was no difference for live birth/ongoing pregnancy rate (OR 1.71, 95% CI 0.61-4.79, 1
1047 RCT) or number of oocytes (MD 0.70, 95% CI -0.11 to 1.51, 1 RCT) (Farquhar, et al., 2017, Kim et al.,
1048 2011).

1049 A recent RCT investigated the use of steroid pre-treatment in IVF/ICSI in a GnRH antagonist protocol in
1050 52 women (Fernández-Prada et al., 2022). No significant difference was found between COCP pre-
1051 treatment and no pre-treatment for cumulative live birth rate (38.7% (12/31) vs. 47.6% (10/21)), live
1052 birth rate (31.8% (7/22) vs. 46.7% (7/15)). There was also no significant difference in the number of MII
1053 oocytes between the study and control group (6.32±5.16 vs. 6.15±4.68).

1054 An RCT, more recent than the meta-analysis, also investigated the effect of COCP pre-treatment
1055 compared to no pre-treatment in a GnRH antagonist protocol in women with PCOS (Gao et al., 2024).
1056 The COCP consisted of ethinyl estradiol (0.03 mg) and drospirenone (3 mg) and were administered daily
1057 for 21 days to induce menstruation, followed by 7 days of washout. No significant differences were
1058 observed between COCP pre-treatment and no pre-treatment for cumulative live birth rate (ITT, 74.4%



1059 (90/121) vs. 77.7% (94/121)), live birth rate (per protocol, 52.8% (56/106) vs. 55.1% (60/109)) or
1060 incidence of moderate to severe OHSS (ITT: 6.6% (8/121) vs. 10.7% (13/121)).

1061 Recommendations

COCP pre-treatment (12-28 days) is not recommended in the GnRH antagonist protocol with FSH alone stimulation, because of reduced efficacy. [updated]

Strong ⊕⊕○○

1062

A minimal wash out period of 5 days should be applied if COCP is used for programming cycle in the case of a fresh transfer. [2025]

GPP

1063

1064 Justification

1065 There is low-quality evidence of a lower live birth/ongoing pregnancy rate using COCP pre-treatment in
1066 GnRH antagonist protocols compared with no pre-treatment. There is low-quality evidence regarding
1067 OHSS incidence.

1068 The type of COCP pre-treatment used in the studies was heterogenous regarding the oestrogen and
1069 progestogen components, as well as the starting days or duration of COCP. The duration varied from
1070 12 to 28 days, and 3 consecutive cycles in one study. In some studies, the duration was fixed and
1071 variable in others, depending on the purpose of scheduling or not (Farquhar et al., 2017). Another
1072 important condition with heterogeneity between studies is the wash-out period between the stop of
1073 COCP pre-treatment and the start of stimulation. This may have an important impact on hormonal
1074 environment (Cedrin-Durnerin et al., 2007).

1075 Lastly, it is important to note however that the available evidence comes predominantly from rFSH
1076 stimulation in GnRH-antagonist protocols and the usage of ethinyl oestradiol combined with either
1077 levonorgestrel or desogestrel as COCP. Whether a negative COCP effect exists in other treatment
1078 protocols or when using other COCPs is unknown.

1079 **GNRH ANTAGONIST PRE-TREATMENT**

1080 Evidence

1081 In an RCT, GnRH antagonist pre-treatment in a GnRH antagonist protocol was investigated in 136
1082 normal ovulatory women (Zhang et al., 2021). In the study group, ovarian stimulation was initiated after
1083 3 days of GnRH antagonist pretreatment. No significant differences were found between GnRH
1084 antagonist pre-treatment and no pre-treatment for live birth rate per embryo transfer (33.9% (20/59)
1085 vs. 43.1% (25/58)) or incidence of moderate to severe OHSS (1.5% (1/68) vs. 2.9% (2/68)). Furthermore,
1086 neither the ongoing pregnancy rate (33.9% (20/59) vs. 45.6% (26/58)) or the number of MII oocytes (7
1087 (6.0-11.0) vs. 9.0 (5.3-12.0)) was different between the study and the control group.



1088 One small RCT in 69 normogonadotropic women (not PCOS, not-poor responder) reported no
1089 difference in ongoing pregnancy rate (42% vs. 33%, 95% CI -13-3) and number of oocytes (12.8±7.8 vs.
1090 9.9±4.9) comparing early follicular pre-treatment with GnRH antagonist (delayed start protocol)
1091 compared to no pre-treatment in fixed antagonist protocol (Blockeel et al., 2011).

1092 In an RCT, including 110 women with PCOS (study group n=50, control group n=60), the effect of three
1093 days of GnRH antagonist pretreatment on the pregnancy outcomes in GnRH antagonist protocols for
1094 IVF/ICSI was evaluated (Eftekhar et al., 2018). The GnRH antagonist was administered for 3 days,
1095 starting on day 2 before the start of a GnRH flexible antagonist protocol with rFSH 150 IU on cycle day
1096 5. The incidence of moderate to severe risk of OHSS was not significantly different between GnRH
1097 antagonist pre-treatment and no pre-treatment (39% (15/38) vs. 36% (18/50). Furthermore, neither
1098 the ongoing pregnancy rate (28% (6/38) vs. 9% (2/50) or the number of MII oocytes (14.65±8.30 vs.
1099 14.10±8.79) was different between the study and the control group.

1100 Recommendation

GnRH antagonist pre-treatment before ovarian stimulation in a delayed-start gonadotrophin protocol is probably not recommended. [2019]	Conditional ⊕○○○
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1101

1102 Justification

1103 There is very low-quality evidence that ongoing pregnancy rate per embryo transfer and number of
1104 oocytes are not statistically different with GnRH antagonist pre-treatment in young normogonadotropic
1105 women (Blockeel, et al., 2011). Only one RCT reported on women with PCOS and reported no significant
1106 differences in efficacy and safety (Eftekhar et al., 2018).

1107 HCG PRE-TREATMENT

1108 Evidence

1109 In an RCT, the effect of short term pre-gonadotropin administration of hCG (n=27) was assessed in
1110 women entering an ICSI cycle and compared to no pre-treatment (n=19) (Beretsos et al., 2009). The
1111 long luteal GnRH agonist protocol with rFSH and 7 days hCG 200 IU/day before rFSH fixed dose of 200
1112 IU daily was used in the study group. Clinical pregnancy rate was significantly higher in the hCG pre-
1113 treatment group (46.2% vs. 31.8%).

1114 Recommendation

hCG pre-treatment can only be used in the context of a clinical trial. [2025]	Research only
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1115



1116 Justification

1117 Even though published results show a benefit of hCG pre-treatment before ovarian stimulation, current
1118 evidence is a single, very small RCT. Insufficient data are available to support or refute the use of hCG
1119 pre-treatment.

1120

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1165



PART C: Pituitary suppression and ovarian stimulation

4. Ovarian stimulation protocols

PICO QUESTION: ACCORDING TO PREDICTED RESPONSE-BASED STRATIFICATION, WHICH STIMULATION PROTOCOL IS MOST EFFICIENT AND SAFE?

Ovarian stimulation for IVF/ICSI aims at obtaining several oocytes that will be turned into embryo's through the laboratory process of in vitro fertilisation. These embryos can then be placed in the uterine cavity within the window of implantation in order to achieve a pregnancy leading to live birth. The contribution of ovarian stimulation to the maximisation of success is under debate for many years. The key issues here are 'how many oocytes do we need to ensure at least one good quality embryo for transfer', 'do more oocytes imply a better chance of obtaining a pregnancy', 'how can we limit the risk of OHSS by the way we stimulate the ovaries' and 'how the level of FSH exposure contribute to creating optimal live birth rates and safety'. In this chapter, the role of the individual predicted ovarian response and the various FSH dosing regimens will be discussed. The policy of getting only a few oocytes more than the one oocyte that will occur in a natural cycle is known under the term MILD stimulation. This is however, a non-standardised term. ICMART describes mild stimulation as a protocol in which the ovaries are stimulated with gonadotropins, and/or other pharmacological compounds, with the intention of limiting the number of oocytes following stimulation for IVF. The definition is often based on the number of follicles developed. It is seen as the intended approach. However, it is difficult to decide on a gonadotropin starting dose to obtain a set number of follicles. In literature, this results in high heterogeneity within study protocols. Therefore, data on this approach will therefore not be presented in this guideline.

A. HIGH RESPONDER

DELAYED-START STIMULATION

Evidence

In an RCT, delayed start of rFSH (day 4; n=22) was studied and compared to conventional start of rFSH (day 2; n=21) in expected high responders in a GnRH antagonist protocol (Revelli et al., 2020). Comparing delayed start stimulation to conventional start stimulation in expected high responders, both the cumulative live birth per oocyte pick-up (52.4% (11/21) vs. 57.1% (12/21)) and the clinical pregnancy rate per started cycle (50.0% (11/22) vs. 47.6% (10/21)) were comparable.

In an RCT, delayed start stimulation with 150 IU rFSH from day 4 in a GnRH antagonist protocol (n=203) was compared to a conventional long GnRH agonist protocol with rFSH (150 IU; n=207) in women with an expected high response to ovarian stimulation (non-PCOS) (Casano et al., 2012). No significant



1198 differences were reported between the delayed start and the conventional protocol for live birth rate
1199 per started cycle (24.9% (51/205) vs. 26.6% (55/207)) or OHSS rate (1.6% vs. 2.0%).

1200 Recommendation

Delayed-start ovarian stimulation is probably not recommended routinely in predicted high responders to decrease the risk of OHSS. [2025]	Conditional ⊕○○○
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1201

1202 Justification

1203 There is insufficient evidence to support or refute the use of delayed start ovarian stimulation for high
1204 responders, compared to conventional ovarian stimulation.

1205 The rationale for delayed-start stimulation is to decrease the risk of OHSS, however, no benefit for
1206 safety in terms of OHSS has been observed in current studies. Other studies on random-start
1207 stimulation have not shown a disadvantage in efficacy in terms of live birth rate.

1208 MODIFIED NATURAL CYCLE

1209 Modified natural cycle (MNC) for IVF is defined as a procedure in which one or more oocytes are
1210 collected from the ovaries during a spontaneous menstrual cycle. Pharmacological compounds are
1211 administered with the sole purpose of blocking the spontaneous LH surge and/or inducing final oocyte
1212 maturation (GLOSSARY).

1213 There is no evidence to justify the use of NC or MNC for OS in high responders.

1214 DOSE COMPARISONS

1215 Evidence

1216 *A Cochrane meta-analysis⁵ including 3 RCTs, including women with a high ovarian response to*
1217 *stimulation, investigated direct gonadotropin dose comparisons (Ngwenya et al., 2024). Since Arce et*
1218 *al. 2014 and Ishihara et al., 2021 were dose-response studies of a novel gonadotropin, the dosages were*
1219 *reported in µg and translation to IU is was not possible, therefore, pooling of the results was also not*
1220 *possible.*

1221 The RCT by Ishihara et al., compared ovarian stimulation with either 6 (n=25), 9 (n=25), 12 µg (n=25)
1222 rFSH in a GnRH antagonist protocol in women with a high ovarian response to stimulation (Ishihara et
1223 al., 2021). Live birth rates were not significantly different between dosages of rFSH (16% (4/25), 24%
1224 (6/25), 24% (6/25)). The rates of moderate or severe OHSS across the three dose groups were 16%, 8%,
1225 and 16%. The number of oocytes retrieved were 8±4.1 vs. 11±5.6 and 13±6.4.

1226 The RCT by Oudshoorn et al., including 521 predicted high responders, compared ovarian stimulation
1227 with 100 IU FSH (n=255) to ovarian stimulation with 150 IU FSH (n=266) either in a GnRH agonist or
1228 GnRH antagonist protocol (Oudshoorn et al., 2017). Comparable rates of ongoing pregnancy within 18
1229 months of FU resulting in live birth were reported (66.3% vs. 69.5%; RR 0.953, 95% CI 0.85–1.07) and

⁵ The Cochrane review by Lensen et al. 2017 was replaced by the updated Cochrane review.



1230 1st cycle live birth (fresh and cryopreserved embryos) (36.0% vs. 39.1%). Lower-dose stimulation
1231 resulted in significantly lower OHSS rate (5.2% vs. 11.8%) as compared with conventional ovarian
1232 stimulation (Oudshoorn, et al., 2017).

1233 The RCT by Arce et al., compared ovarian stimulation with either 5.2 (=23), 6.9 (n=26), 8.6 (n=24), 10.3
1234 (n=24), or 12.1 µg (n=26) of rFSH, or 11 µg (150 IU, n=25)) of follitropin alfa in a GnRH antagonist cycle
1235 in women with a high ovarian response to stimulation (AMH 15.0-44.9 pmol/L) (Arce et al., 2014). There
1236 was no significant difference between the different dosages and the conventional dose of follitropin
1237 alfa for cumulative live birth rate (43% (10/23), 54% (14/26), 46% (11/24), 38% (9/24), 50% (13/26) vs.
1238 56% (14/25)) or live birth rate (39% (9/23), 42% (11/26), 38% (9/24), 25% (6/24), 46% (12/26) vs. 48%
1239 (12/25). A statistically significant dose–response relationship with respect to number of oocytes
1240 retrieved was established for rFSH (5.9±3.9, 9.1±6.4, 10.6±4.8, 13.6±7.8, 14.4±5.8 vs. 12.4±5.4). Two
1241 cases of early OHSS were reported in the highest rFSH dose groups (10.3 and 12.1 µg, respectively), and
1242 three late OHSS (one in the 8.6 µg group and two in the 12.1 µg group).

1243 Recommendation

A reduced gonadotropin dose is probably recommended to decrease the risk of OHSS in predicted high responders. [2025]	Conditional ⊕○○○
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1244

The GnRH antagonist protocol is recommended for predicted high responders. However, if GnRH agonist protocols are used, a reduced gonadotropin dose is recommended to decrease the risk of OHSS. [updated]	Strong ⊕○○○
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1245

1246 Justification

1247 The recommendation is extrapolated from a stratified group analysis of three RCTs in women with high
1248 levels of AMH. Two RCTs were dose-finding studies for a new follitropin in the GnRH antagonist protocol
1249 (Arce et al., 2014, Ishihara et al., 2021) and in the third RCT, the majority of the patients were treated
1250 with the long GnRH agonist protocol. The data from the Oudshoorn trial shows that lowering
1251 gonadotropin dosage may increase safety in GnRH agonist protocol. However, the mix of GnRH agonist
1252 and antagonist protocols, the per protocol allowance of dose adjustments in 2nd cycle and the very high
1253 cycle cancellation rate in high responders should be carefully considered when interpreting the
1254 available evidence. Furthermore, the fact that a freeze-all policy was not adopted in the trial, a strategy
1255 which may reflects current clinical practice, questions the potential negative effects of conventional
1256 dosage stimulation in terms of cumulative pregnancy rate and OHSS rates. The two dose-finding trials
1257 were not powered to show a difference in OHSS incidence.



1258 B. NORMAL RESPONDER

1259 DELAYED-START STIMULATION

1260 Evidence

1261 In an RCT, delayed start of rFSH (day 4; n=19) was studied and compared to conventional start of rFSH
1262 (day 2; n=20) in expected normal responders (Revelli et al., 2020). Comparing delayed start stimulation
1263 to conventional start stimulation in expected normal responders, both the cumulative live birth per
1264 oocyte pick-up (16.7% (3/18) vs. 26.3% (5/19)) and the clinical pregnancy rate per started cycle (16.7%
1265 (3/18) vs. 26.3% (5/19)) were comparable.

1266 In an RCT, women with an expected normal response to ovarian stimulation, starting their first IVF cycle
1267 and younger than 35 years were randomised to receive either ovarian stimulation with hMG (150 IU
1268 daily) without pituitary suppression (n=30) or a long GnRH agonist protocol with rFSH (150-300 IU;
1269 n=30) (Lou and Huang, 2010). No significant difference was reported when comparing the study group
1270 to the control group for mild OHSS (0 vs. 6.7% (2/30)), ongoing pregnancy rate per started cycle (26.7%
1271 (8/30) vs. 23.3% (7/30)) or clinical pregnancy rate per started cycle (30.0% (9/30) vs. 30.0% (9/30)).

1272 Three older RCTs compared the late-start FSH (fixed dose of 150 IU starting on cycle day 5) with
1273 conventional-start FSH (Baart et al., 2007, Blockeel et al., 2011, Hohmann et al., 2003). The RCT by Baart
1274 et al. compared late-start FSH in the GnRH antagonist protocol with conventional FSH stimulation in
1275 the long GnRH agonist protocol in 111 women and reported no significant difference in ongoing
1276 pregnancy rate (19% (12/63) vs. 17% (7/41)). However, significantly less oocytes retrieved with the late-
1277 start FSH protocol (8.3±4.7 vs. 12.1±5.7) (Baart et al., 2007). The RCT by Hohmann et al. including 104
1278 predicted normal responders, compared late-start with conventional-start FSH in the GnRH antagonist
1279 protocol and reported no difference in ongoing pregnancy rate (16% (8/49) vs. 17% (8/48)) or number
1280 of oocytes retrieved (7 (1-27) vs. 8 (2-31)) (Hohmann et al., 2003). The RCT by Blockeel et al. including
1281 76 predicted normal responders also compared late-start with conventional-start FSH in the GnRH
1282 antagonist protocol and also reported no significant difference in ongoing pregnancy rate (25% 10/40
1283 vs. 28% (10/36) (Blockeel et al., 2011)).

1284 Recommendation

Delayed-start ovarian stimulation is probably not recommended over a conventional gonadotrophin dose for predicted normal responders. [2025]	Conditional ⊕○○○
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1285

1286 Justification

1287 The rationale to delay the start of ovarian stimulation would be the prevention of OHSS. However, this
1288 topic has not been researched well and current RCTs were not powered to show a difference in OHSS
1289 rate. As a result, no benefit for safety in terms of OHSS has been observed in current studies.

1290 There is insufficient evidence to support or refute the use of delayed start ovarian stimulation for
1291 normal responders, compared to conventional ovarian stimulation.



1292 DOSE COMPARISONS

1293 Evidence

1294 A Cochrane meta-analysis⁶ including 12 RCTs, including women with a normal ovarian response to
1295 stimulation, investigated direct gonadotropin dose comparisons (Ngwenya et al., 2024). For moderate
1296 or severe OHSS, the estimates of difference between the dose comparisons were very imprecise, there
1297 is little information about the true treatment effect.

1298 200 IU vs. 100 UI

1299 No significant difference in live birth/ongoing pregnancy rate was observed of the different doses (OR
1300 0.88, 95% CI 0.57-1.36, 2 RCTs, 522 women) (Ngwenya et al., 2024). No significant difference in the
1301 incidence of severe OHSS was found with the different gonadotropin doses (peto OR 0.14, 95% CI 0.00-
1302 6.96, 2 RCT, 522 women) or in the incidence of moderate to severe OHSS (peto OR 0.62, 95% CI 0.21-
1303 1.87, 2 RCTs, 522 women). The pooled estimate suggests a higher number of oocytes were obtained
1304 with the higher dose of gonadotropin (ratio of mean oocytes 1.58, 95% CI 1.43-1.77, 2 RCTs, 330
1305 women). However, the statistical heterogeneity was high.

1306 225/200 IU vs. 150 UI

1307 No significant difference in live birth rate was observed of the different doses (OR 0.98, 95% CI 0.70-
1308 1.36, 2 RCTs, 211 women) (Ngwenya, et al., 2024). Two RCTs reported on cumulative live birth rate,
1309 using two different definitions. However, these data could neither confirm nor rule out dose effects on
1310 cumulative live birth. No significant difference in the incidence of severe OHSS was found with the
1311 different gonadotropin doses (peto OR 1.00, 95% CI 0.20-5.02, 4 RCT, 740 women) or in the incidence
1312 of moderate to severe OHSS (peto OR 1.21, 95% CI 0.51-2.85, 4 RCTs, 740 women). The pooled estimate
1313 suggests a higher number of oocytes were obtained with the higher dose of gonadotropin (ratio of
1314 mean oocytes 1.16, 95% CI 1.08-1.25, 6 RCTs, 872 women).

1315 300 IU vs. 150 UI

1316 No clear impact of different doses on the probability of live birth were found (OR 0.80, 95% 0.19-3.42,
1317 1 RCT, 37 women (Ngwenya, et al., 2024, Shyamsunder et al., 2021). The ratio of mean oocytes was
1318 1.23 (95% CI 0.89-1.72, 57 women).

1319 300 IU vs. 225 UI

1320 No clear impact of different doses on the probability of live birth were found (OR 0.65, 95% 0.32-1.32,
1321 1 RCT, 47 women (Jayaprakasan et al., 2010, Ngwenya, et al., 2024). No significant difference in the
1322 incidence of severe OHSS was found with the different gonadotropin doses (peto OR 0.14, 95% CI 0.00-
1323 6.92, 1 RCT, 135 women) or in the incidence of moderate to severe OHSS (peto OR 0.67, 95% CI 0.11-
1324 3.99, 1 RCT, 135 women). The available evidence could not rule out or confirm an effect of
1325 gonadotropin dosing on the number of retrieved oocytes (ratio of mean oocytes 1.03, 95% CI 0.84-1.26,
1326 1 RCT, 135 women).

⁶ The Cochrane review by Lensen et al. 2017 on dose comparison and the meta-analysis on mild gonadotropin dosing by Sterrenburg et al., 2011 were replaced by the updated Cochrane review.



1327 Recommendation

Neither a reduced nor increased gonadotrophin dose is probably recommended over a conventional gonadotrophin dose (equivalent to 150-225 IU) for predicted normal responders. [updated]

Conditional ⊕○○○

1328

1329 Justification

1330 In published metaanalysis the chance of live birth in normal responders is not affected by modifications
1331 in the FSH starting dose. The heterogeneity of the studies is too high to be conclusive on the impact of
1332 dose and type of FSH on the number of retrieved oocytes as well as on the risk of OHSS.

1333 The meta-analysis suggests that the optimal daily rFSH stimulation dose is 150 IU/day in predicted
1334 normal responders. Although available studies suggest similar efficacy in terms of clinical pregnancy
1335 rate between reduced-dose and conventional-dose stimulation, the lower number of oocytes retrieved
1336 could potentially compromise cumulative live birth rate in predicted normal responders.

1337 The recommendation is based on studies conducted in GnRH agonist protocols, however, the guideline
1338 group thinks that the recommendation may also apply to GnRH antagonist protocol due to the
1339 increased safety with the option of the GnRH agonist trigger.

1340 C. LOW RESPONDER

1341 DELAYED-START STIMULATION

1342 Evidence

1343 In an RCT, delayed start of rFSH (day 4; n=15) was studied and compared to conventional start of rFSH
1344 (day 2; n=16) in expected poor responders (Revelli et al., 2020). Comparing delayed start stimulation to
1345 conventional start stimulation in expected poor responders, both the cumulative live birth per oocyte
1346 pick-up (0% (0/9) vs. 23.1% (3/13)) and the clinical pregnancy rate per started cycle (0% (0/15) vs. 18.7%
1347 (3/16)) were significantly lower.

1348 Recommendation

Delayed start ovarian stimulation is probably not recommended for predicted low responders. [2025]

Conditional ⊕○○○

1349

1350 Justification

1351 There is insufficient evidence to support or refute the use of delayed start ovarian stimulation for low
1352 responders, compared to conventional ovarian stimulation.



1353 **MODIFIED NATURAL CYCLE**

1354 **Evidence**

1355 In an RCT, 90 women with a low response to ovarian stimulation were randomised to receive either
1356 minimal ovarian stimulation (150 IU from day 7/8) or conventional stimulation (225 IU) in a GnRH
1357 antagonist protocol (Kim et al., 2009). No significant difference in clinical pregnancy per cycle was
1358 reported (13.3% (6/45) vs. 17.8% (8/45)). The number of MII oocytes retrieved was significantly lower
1359 in the lower dose gonadotropins group (1.3±0.8 vs. 2.5±1.4).

1360 One RCT compared MNC-IVF with a microdose GnRH agonist flare protocol in 125 poor responder
1361 women (215 cycles) and reported no significant difference in pregnancy rate (6.1% vs. 6.9%) (Morgia
1362 et al., 2004).

1363 In a retrospective cohort study, natural cycle IVF (n=230) was compared to conventional ovarian
1364 stimulation in GnRH antagonist protocol (n=355) in poor ovarian responders and aged ≥40 years. (De
1365 Marco et al., 2021). In the natural cycle IVF group, no treatment was administered for the selection and
1366 recruitment of follicles, however, ovulation was triggered with 10,000 IU of hCG. Comparing natural
1367 cycle IVF to conventional stimulation, no significant difference was seen in cumulative live birth rate
1368 (9.6% (22/230) vs. 14.4% (51/355)), however, the cumulative pregnancy rate per cycle was significantly
1369 higher with conventional stimulation (6.3% (36/576) vs. 12.9% (70/543)).

1370 **Recommendation**

The use of modified natural cycle is probably not routinely recommended over conventional stimulation for low responders.	Conditional ⊕○○○
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1371

The GDG recognises that low responders are a heterogeneous group and in women with very low ovarian reserve, clinicians could choose to use a modified natural cycle.	GPP
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1372

1373 **Justification**

1374 There are no good-quality, controlled studies available to support the use of modified natural cycle or
1375 natural cycle IVF in low responders. Furthermore, the number of oocytes were lower with modified
1376 natural cycle compared to conventional stimulation. Although there are no good quality studies looking
1377 at modified natural cycle in women with very low number of follicles, who would not benefit
1378 significantly from conventional stimulation, a modified natural cycle could be considered.



1379 **DOSE COMPARISONS**

1380 **Evidence**

1381 A Cochrane meta-analysis⁷ including 6 RCTs, including women with a poor ovarian response to
1382 stimulation, investigated direct gonadotropin dose comparisons (Ngwenya, et al., 2024). For live birth
1383 or ongoing pregnancy, the estimates of difference between the dose comparisons were very imprecise,
1384 there is little information about the true treatment effect.

1385 300/450 IU vs. 150 IU

1386 The Cochrane meta-analysis reported no significant difference in live birth/ongoing pregnancy rates (3
1387 RCT, OR 1.20, 95% CI 0.78-1.86, 538 women) between the 150 IU and 300/450 IU dose of gonadotropins
1388 and no cases of moderate or severe OHSS were observed in either group. However, the pooled effect
1389 suggests that slightly more oocytes were retrieved in the higher gonadotropin dose group (3 RCT, ratio
1390 of mean oocytes 1.97, 95% CI 1.70 to 2.29, 947 women) (Ngwenya, et al., 2024).

1391 400/450 IU vs. 300 IU

1392 The Cochrane meta-analysis reported no significant difference in ongoing pregnancy rate (1 RCT, OR
1393 0.77, 95% CI 0.19-3.19, 62 women) or number of oocytes retrieved (2 RCT, ratio of mean oocytes 0.97,
1394 95% CI 0.74 to 1.27, 110 women) between the 300 IU and 400/450 IU dose of gonadotropins and no
1395 cases of moderate or severe OHSS in either group (Ngwenya, et al., 2024).

1396 600 IU vs. 450 UI

1397 The Cochrane meta-analysis reported no significant difference in live birth rate (1 RCT, OR 1.33, 95% CI
1398 0.71-2.52, 356 women), or number of oocytes retrieved (1 RCT, ratio of mean oocytes 1.08, 95% CI 0.96
1399 to 1.22, 356 women) between the 450 IU and 600 IU dose of gonadotropins and one case of moderate
1400 OHSS in the 600 IU dose group (Lefebvre et al., 2015, Ngwenya, et al., 2024).

1401 **Recommendation**

A higher gonadotropin dose is probably not recommended over conventional (equivalent to 150-225 IU) for predicted low responders. [reworded] Conditional ⊕○○○

1402

A gonadotropin dose higher than 300 IU is not recommended for predicted low responders. [2019] Strong ⊕○○○

1403

1404 **Justification**

1405 There is evidence that a higher gonadotropin dose than 150 IU results in a higher number of oocytes in
1406 low responders, and more chances of having an embryo for transfer. However, there was no difference
1407 in live birth/ongoing pregnancy rates. Furthermore, the sample sizes of the studies are small and

⁷ The Cochrane review by Lensen et al. 2017 was replaced by the updated Cochrane review.



1408 therefore not sufficient to provide evidence on the benefits of various dosing levels over the standard
1409 dose for the outcome live birth.

1410 There is unlikely to be significant benefit with doses >300 IU daily, as comparisons with doses >300 IU
1411 did not show significant differences in the above mentioned pre-clinical outcomes.

1412

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Draft for review



1473 5. Pituitary suppression regimes

PICO QUESTION: WHICH PITUITARY SUPPRESSION PROTOCOL IS PREFERABLE?

1474 GNRH AGONIST PROTOCOLS

1475 Evidence

1476 A Cochrane meta-analysis including 40 RCTs compared different GnRH agonist protocols (Siristatidis et
1477 al., 2025).

1478 Long vs short GnRH agonist protocol⁸

1479 The Cochrane meta-analysis found no significant difference in live birth rate/ongoing pregnancy rate
1480 per woman randomised (OR 1.45, 95% CI 0.83-2.52, 5 RCT, 381 women) between the long and the short
1481 GnRH agonist protocol (Siristatidis et al., 2025). None of the included studies for this comparison
1482 reported OHSS rates.

1483 An RCT, not included in the Cochrane meta-analysis, including 131 women also reported no significant
1484 difference in clinical pregnancy rate between the long and the short GnRH agonist protocol (19.6% vs.
1485 8.3%) (Ravhon et al., 2000).

1486 However, another RCT, not included in the Cochrane meta-analysis, including 220 women ≥ 40 years of
1487 age, reported a significantly reduced clinical pregnancy rate with the short GnRH agonist protocol as
1488 compared to the long (10.9% (12/110) vs. 22.7% (25/110)) (Sbracia et al., 2005).

1489 Long vs ultrashort GnRH agonist protocol

1490 The Cochrane meta-analysis found no significant difference in live birth rate when a long protocol was
1491 compared with an ultrashort GnRH agonist protocol (1 RCT, OR 1.78, 95% CI 0.72-4.36, 150 women)
1492 (Kingsland et al., 1992, Siristatidis et al., 2025). There were no data on adverse outcomes reported.

1493 Short vs ultrashort GnRH agonist protocol

1494 The Cochrane meta-analysis reported no significant difference in the clinical pregnancy rate when a
1495 short protocol was compared with an ultrashort protocol (1 RCT, OR 1.33, 95% CI 0.47-3.81, 82 women)
1496 (Berker et al., 2010, Siristatidis et al., 2025). There were no data on adverse outcomes reported.

1497 Long GnRH agonist protocol: luteal vs follicular start

1498 The Cochrane meta-analysis found no significant difference in live birth/ongoing pregnancy rates when
1499 GnRH agonist was commenced in the luteal or follicular phase for the long protocol (1 RCT, OR 1.89,
1500 95% CI 0.87-4.10, 223 women) (Siristatidis et al., 2025, Urbancsek and Witthaus, 1996). There were no
1501 data on adverse outcomes reported.

1502 The RCT by Ravhon et al., including 125 women, also reported no significant difference in pregnancy
1503 rate when GnRH agonist was started on day 2 versus day 21 (19.6% vs. 18.6%) (Ravhon et al., 2000).

⁸ A meta-analysis was cited here in the previous version of the guideline on the long versus short GnRH agonist protocol in women with adenomyosis. The reader is referred to the Good Practice Recommendations paper on Adenomyosis for updated advice on fertility treatment in women with adenomyosis.



1504 Long GnRH agonist protocol: continuation vs stopping GnRH agonist at start of stimulation
1505 The Cochrane meta-analysis found no significant difference in the number of ongoing pregnancies (OR
1506 0.66, 95% CI 0.30-1.49, 2 RCT, 194 women), clinical pregnancy rate (OR 0.76, 95% CI 0.40-1.44, 3 RCT,
1507 264 women) when GnRH agonist was stopped compared with when it was continued (Siristatidis et al.,
1508 2025).

1509 Long agonist protocol: continuation of same-dose vs reduced-dose GnRH agonist until trigger
1510 The Cochrane meta-analysis found no significant difference in live birth/ongoing pregnancy rate (OR
1511 1.59, 95% CI 0.66-3.87, 1 RCT, 96 women) or clinical pregnancy rate when the dose of GnRH agonist
1512 was reduced compared with when the same dose was continued (4 RCT, OR 1.02, 95% CI 0.68-1.52,
1513 407 women) (Siristatidis et al., 2025). There was no significant difference in OHSS rate between
1514 continuing or reducing the GnRH agonist dose (OR 0.47, 95% CI 0.04-5.35, 1 RCT, 96 women).

1515 Recommendation

If GnRH agonists are used, the long GnRH agonist protocol is recommended over the short or ultrashort GnRH agonist protocol. [updated]

Strong

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1516

1517 Justification

1518 The long GnRH agonist protocol has proven to be highly efficient for preventing LH surge. Since its
1519 introduction, there has been a reduction of cycle cancellation, increased number of oocytes retrieved
1520 and higher pregnancy rates. Compared to other GnRH agonist protocols, the long protocol provides
1521 better efficacy and is supported by a larger body of evidence.

1522 The short GnRH agonist protocol appeared as a modification of the classic long protocol with the aim
1523 of improving cycle outcome in low responders and older patients. The current evidence available shows
1524 that this goal is not achieved.

1525 GNRH ANTAGONIST PROTOCOLS

1526 Evidence

1527 A systematic review and meta-analysis⁹ including 36 RCTs in the general IVF population, compared the
1528 GnRH antagonist protocol with the long GnRH agonist protocol. They did not include RCTs reporting on
1529 early follicle phase start-up GnRH antagonist or long-acting follicular GnRH agonist protocols (Liu et al.,
1530 2023). No significant difference was found between the GnRH antagonist and long GnRH agonist
1531 protocol for live birth rate (RR 0.95, 95% CI 0.86-1.06, 10 RCT, 2939 women) or ongoing pregnancy rate
1532 (RR 0.94, 95% CI 0.86-1.03). However, the risk of OHSS was significantly lower with the GnRH antagonist
1533 protocol (RR 0.84, 95% CI 0.75-0.94, 17 RCT, 4892 women), especially the risk of moderate or severe
1534 OHSS (RR 0.56, 95% CI 0.40-0.79, 15 RCT, 4481 women).

⁹ The Cochrane review by Al-Inany et al., 2016 was replaced by a newer meta-analysis. The RCTs by Friedler et al., 2006 and Toftager et al., 2016 are included in the meta-analysis and therefore no longer mentioned separately.



1535 An RCT, not included in the meta-analysis, including 132 women, reported a significantly higher clinical
1536 pregnancy rate with the long GnRH agonist protocol as compared to the GnRH antagonist protocol
1537 (49.2% vs. 26.2%). One case of mild OHSS developed in each group (Verpoest et al., 2017).

1538 Two RCTs including respectively 160 cycles and 96 women, compared the GnRH antagonist protocol
1539 with the short GnRH agonist protocol (Gordts et al., 2012, Maldonado et al., 2013). Gordts *et al.*
1540 reported an ongoing pregnancy rate of 21% and a live birth rate of 19% in GnRH antagonist cycles
1541 compared to 20% and 20% respectively in GnRH agonist cycles, which are both not statistically different
1542 (Gordts et al., 2012). However, Maldonado *et al.* reported a significantly lower clinical pregnancy rate
1543 (31.0% (13/48) vs. 52.1% (25/48)) in the short GnRH agonist protocol as compared to the GnRH
1544 antagonist protocol (Maldonado et al., 2013).

1545 A systematic review and meta-analysis, including 7 RCTs, compared fixed and flexible GnRH antagonist
1546 protocols (Venetis et al., 2023). No significant difference in ongoing pregnancy rate (RR 0.85, 95% CI
1547 0.73-1.00) was observed between the fixed and flexible GnRH antagonist protocol without pre-
1548 treatment.

1549 Recommendation

The GnRH antagonist protocol is recommended over the GnRH agonist protocols given the comparable efficacy and higher safety in the general IVF/ICSI population. [2019]	Strong ⊕⊕⊕○
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The flexible and fixed GnRH antagonist protocol is probably equally recommended. [2025]	Conditional ⊕⊕○○
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1551

1552 Justification

1553 The introduction of GnRH antagonist allowed overcoming the significant undesirable effects of the
1554 GnRH agonist protocols. Although the first studies reported slight but consistent lower pregnancy rates,
1555 which delayed the implementation of the GnRH antagonist protocol, several large meta-analyses
1556 published in the past 10 years support similar live birth rates. There is far less evidence for the short
1557 GnRH agonist protocol, however, results are expected to be similar as for the long GnRH agonist
1558 protocol.

1559 Although there is high heterogeneity in RCTs comparing flexible to fixed GnRH antagonist protocols,
1560 results show that live birth and ongoing pregnancy rates are similar with a flexible GnRH antagonist
1561 protocol (Venetis et al., 2023).

1562 PROGESTIN PROTOCOLS

1563 The use of oral progestins to prevent the LH surge is a novel protocol in which GnRH analogues are not
1564 used. Progestin administration along the whole stimulation will keep the pituitary suppressed and has
1565 shown to prevent untimely LH surges effectively. However, the use of this protocol implies the freezing



1566 of all the embryos and transfer in a subsequent frozen embryo replacement cycle, as the endometrium
1567 would not be receptive in a fresh cycle due to the effect of the progestins.

1568 Evidence

1569 *Progestogens vs. GnRH analogues*

1570 A Cochrane systematic review and meta-analysis including 100 normal responders from 1 RCT,
1571 compared ovarian stimulation with gonadotropins in combination with progestogens with
1572 gonadotropins combined with GnRH antagonist (Ghasemzadeh et al., 2019, Glujovsky et al., 2023).
1573 Significantly more MII oocytes were retrieved after stimulation with progestogens (10.8±5.8 vs. 7±4.2;
1574 MD 3.80, 95% CI 1.82 to 5.78). A more recent RCT, including 200 unselected women undergoing
1575 IVF/ICSI, compared dydrogesterone with the flexible GnRH antagonist protocol for pituitary suppression
1576 (Hossein Rashidi et al., 2020). No significant difference was reported in clinical pregnancy rate per first
1577 embryo transfer (43.95% (40/97) vs. 49.50% (45/95)) between dydrogesterone and GnRH antagonist
1578 for pituitary suppression, however, significantly more MII oocytes were retrieved after dydrogesterone
1579 treatment (7.90±3.62 vs. 6.26±3.64).

1580 In an RCT, 348 women with normal ovarian reserve were randomised to receive ovarian stimulation
1581 with in a progestin protocol with freeze-all (n=174) or GnRH antagonist protocol with fresh transfer first
1582 (n=174) (Ye et al., 2024). No significant difference was reported between the progestin protocol group
1583 and the GnRH antagonist group for cumulative live birth rate per woman (55.7% (97/174) vs. 52.9%
1584 (92/174)) or clinical pregnancy rate per transfer (57% (114/200 vs. 55.9% (109/195)). No cases of OHSS
1585 were reported in either group.

1586 The Cochrane systematic review and meta-analysis including 260 women from 1 RCT, compared ovarian
1587 stimulation with gonadotropins in combination with progestogens with gonadotropins combined with
1588 GnRH agonist (Glujovsky et al., 2023, Xi et al., 2020). No significant difference was found for live
1589 birth/ongoing pregnancy rate (45.3% (59/130) vs. 46.9% (61/130); OR 0.94, 95% 0.58-1.53), OHSS rate
1590 (0% (0/130) vs. 2.3% (3/130); OR 0.14, 95% CI 0.01-2.73), clinical pregnancy rate (50% (65/130) vs.
1591 53.1% (69/130); OR 0.88, 95% CI 0.54-1.44) or number of MII oocytes (10.3±5.8 vs. 10.1±5.2; MD 0.20,
1592 95% CI -1.14 to 1.54).

1593 The Cochrane systematic review and meta-analysis including 340 poor responders from 1 RCT,
1594 compared ovarian stimulation with gonadotropins in combination with progestogens with
1595 gonadotropins combined with GnRH antagonist (Chen et al., 2019, Glujovsky et al., 2023). No significant
1596 difference was found for live birth rate/ongoing pregnancy rate (21.8% (37/170) vs. 18.2% (31/170); OR
1597 1.25; 95% CI 0.73-2.13), clinical pregnancy rate (28.2% (48/170) vs. 22.9% (39/170); OR 1.32; 95% CI
1598 0.81-2.16), or number of MII oocytes (3.2±2.4 vs. 2.8±2.2; MD 0.40; 95% CI -0.09 to 0.89).

1599 In an RCT, 484 predicted suboptimal responders were randomly assigned to receive ovarian stimulation
1600 in a progestin protocol (n=236) compared to a GnRH antagonist protocol (n=248) with freeze-all in both
1601 groups (Cai et al., 2024). Cumulative live birth rate over 12 months was 44.4% (96/216) in the progestin
1602 protocol group compared to 48.9% (114/233) in the GnRH antagonist group (RR 0.91, 95% 0.74-1.11).
1603 Live birth rate after the first transfer was 32.9% (71/216) with the progestin protocol compared to
1604 34.3% (80/240) with the GnRH antagonist protocol (RR 0.96, 95% CI 0.74-1.24).

1605 In a systematic review and meta-analysis 3 RCTs were included with women with PCOS, one comparing
1606 progestogens to the GnRH agonist short protocol and two comparing to the GnRH antagonist protocol



1607 (Yang et al., 2023). No significant difference for live birth rate (OR 1.46, 95% CI 0.79-2.71, 167 cycles),
1608 OHSS rate (OR 0.19, 95% CI 0.01-4.11, 2 RCTs, 240 patients). Also, the number of MII oocytes retrieved
1609 was similar in both groups (MD -0.85; 95% CI -3.40 to 1.71, 3 RCTs, 358 patients).

1610 In an RCT, 784 women with an anticipated high response to ovarian stimulation were randomised to
1611 follow a progestin protocol (n=392) or GnRH antagonist protocol (n=392) for IVF/ICSI with freeze-all in
1612 both groups (Chen et al., 2024). No significant difference was observed in cumulative live birth rate
1613 (54.6% (214/392) vs. 48.5% (190/392); ITT) or live birth rate after the first transfer (37.5% (147/392 vs.
1614 32.7% (128/392); ITT).

1615 Progestogens vs. other progestogens

1616 Cochrane systematic review and meta-analysis¹⁰ 4 mg vs. 10 mg MPA. No significant difference in live
1617 birth/ongoing pregnancy rate (53/150 vs. 39/150; OR 1.56; 95 CI 0.95-2.55), clinical pregnancy rate
1618 (73/150 vs. 87/150; OR 0.69; 95% CI 0.44-1.08) (Dong et al., 2017, Glujovsky et al., 2023). No cases of
1619 moderate or severe OHSS were reported.

1620 One RCT including 516 women compared dydrogesterone with MPA for pituitary suppression and
1621 reported no significant difference in clinical pregnancy rate (57.6 (125/217) vs. 62.3% (132/212); OR
1622 0.82, 95% CI 0.56-1.21) or number of oocytes retrieved (10.8±6.3 vs. 11.1±5.8) (Yu et al., 2018). No
1623 cases of moderate or severe OHSS were reported.

1624 Recommendation

If freeze-all is planned, the use of progestin for pituitary suppression is probably equally recommended to GnRH analogues. [updated]

Conditional

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1625

1626 Justification

1627 Oral progestins are efficient in terms of pituitary suppression, with comparable oocyte yield and
1628 pregnancy outcomes as the GnRH short agonist protocol. This approach is easy, cheap and patient
1629 friendly.

1630 Many of the studies use the term PPOS. The GDG would like to clarify that the terminology PPOS, i.e.
1631 progestin-primed ovarian stimulation is not correct. More correct terminology would be progestin
1632 protocol for pituitary suppression.

1633 The progestin protocol approach is only feasible for OS cycles in which a fresh embryo transfer is not
1634 scheduled, such as fertility preservation, oocyte donors, PGT, or pre-planned freeze-all cycles.

1635 Current evidence shows that euploidy rates and clinical outcomes in PGT are also similar between
1636 progestin and GnRH antagonist protocol (Qin et al., 2025, Wan et al., 2024, Zhou et al., 2025).

1637 A meta-analysis including four retrospective cohort studies found no increased risk of congenital
1638 malformations with the use of progestins for pituitary suppression compared to GnRH agonist protocol
1639 (OR 0.92, 95% CI 0.63-1.34) (Zolfaroli et al., 2020). The results of sensitivity analysis by progestin type

¹⁰ The cohort studies by Chen et al., 2017, Hamdi et al., 2018 and Kuang et al., 2015 were excluded in the presence of several RCTs.



1640 were consistent with the main results. These results are also in line with a more recent, very large
1641 retrospective cohort study, including 15382 PPOS cycles and 1352 GnRH antagonist cycles (Li et al.,
1642 2022). Congenital malformations were observed in 323 of 15,245 (2.1%) in the PPOS group and 27 of
1643 1,248 (2.2%), with a nonsignificant difference.

1644

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1738 6. Types of gonadotropins and other ovarian stimulation drugs

PICO QUESTION: IS THE TYPE OF STIMULATION DRUG ASSOCIATED WITH EFFICACY AND SAFETY?

1739 A. GONADOTROPINS

1740 RECOMBINANT FSH (rFSH)

1741 RECOMBINANT FSH (rFSH) VS HUMAN MENOPAUSAL GONADOTROPIN (hMG)

1742 Evidence

1743 In a systematic review¹¹ and meta-analysis, ovarian stimulation with rFSH was compared to highly
1744 purified (hp)-hMG (Bordewijk et al., 2019). No significant difference was found for cumulative live birth
1745 rate when comparing ovarian stimulation with rFSH and hp-hMG (RR 0.91, 95% CI 0.80-1.04, 3 RCT,
1746 2109 women). Live birth rate (RR 0.88, 95% CI 0.78-0.99, 7 RCT, 3397 women) and clinical pregnancy
1747 rate (RR 0.90, 95% CI 0.81-1.00, 7 RCT, 3397 women) were lower with rFSH for ovarian stimulation
1748 compared to hp-hMG.

1749 An RCT, not included in the meta-analysis, included 160 women and also compared hMG to rFSH in the
1750 GnRH agonist protocol. No significant differences were reported for live birth rate (27.5% (11/40) vs.
1751 40% (16/40)) between hMG and rFSH for OS (Parsanezhad et al., 2017).

1752 An RCT compared the efficacy and safety of highly purified hMG (150 IU) and rFSH (150 IU) for ovarian
1753 stimulation with the GnRH antagonist protocol in a population of patients predicted to be high
1754 responders (Witz et al., 2020). Cumulative live birth rates per cycle start were 50.6% and 51.5% in hMG
1755 treated and rFSH-treated patients (difference: -0.8%, 95% CI -8.7% to 7.1%). Similarly, comparing hMG
1756 and rFSH, there was no significant difference in live birth rate after fresh (52.2% vs. 48.7%; difference
1757 3.6, 95% CI -6.4 to 13.4) or frozen (63.4% vs. 50.8%; difference 12.7, 95% CI -0.9 to 26.2) embryo
1758 transfer. The incidence of OHSS was significantly lower with hMG compared to rFSH (9.7% (30/310) vs.
1759 21.4% (66/309); difference -11.7%, 95% CI -17.3% to -6.1%).

1760 A small RCT including 80 PCOS patients reported no significant difference in live birth rate (23.1% vs.
1761 35.7%) or mild OHSS rate (0.0% (0/38) vs. 11.9% (5/42)) between hMG and rFSH for OS (Figen Turkcapar
1762 et al., 2013).

1763 Recommendation

The use of recombinant FSH (rFSH) and human menopausal gonadotropin (hMG) for ovarian stimulation is equally recommended. [2019]

Strong

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1764

¹¹ The Cochrane systematic review (van Wely et al., 2011) that was mentioned here in the 2019 version of the guideline was replaced by a more recent meta-analysis. The RCTs by Devroey et al., 2012 and Ye et al., 2012 are included in the meta-analysis and therefore no longer mentioned separately.



1765 **Justification**

1766 The results from the meta-analysis suggest no significant difference in cumulative live birth rate and a
1767 slightly higher efficacy (LBR/PR) with hMG compared to rFSH in GnRH agonist cycles. Effects on OHSS
1768 rates were not reported in the meta-analysis.

1769 For GnRH antagonist cycles, the evidence is less extensive, however the RCTs by Bosch et al. and
1770 Devroey et al. showed highly purified hMG to be at least as effective as rFSH in antagonist cycles (Bosch
1771 et al., 2008, Devroey et al., 2012). Similar results were reported by Witz et al. in high responders (Witz
1772 et al., 2020).

1773 Studies for this question in PCOS and women of advanced age were limited, so that a potential
1774 difference between compounds in these subgroups cannot be ruled out based on the current evidence.

1775 **RECOMBINANT FSH (rFSH) VS PURIFIED URINARY FSH (p-FSH)**

1776 **Evidence**

1777 In a Cochrane systematic review and meta-analysis, use of rFSH was not associated with a higher
1778 probability of live birth as compared to p-FSH when downregulation was achieved with GnRH agonists
1779 (5 RCT, OR 1.26, 0.96-1.64, 1430 women). The meta-analysis reported no significant difference in OHSS
1780 rate between rFSH and p-FSH (6 RCT, OR 1.79, 95% CI 0.89 to 3.62, 1490 women) (van Wely et al.,
1781 2011).

1782 **Recommendation**

The use of recombinant FSH (rFSH) and purified FSH (p-FSH)
for ovarian stimulation in GnRH agonist protocol is equally
recommended. [2019]

Strong



1783

1784 **Justification**

1785 In patients undergoing ovarian stimulation for IVF/ICSI, the use of p-FSH is not preferable to rFSH when
1786 downregulation is achieved with GnRH agonists, according to the Cochrane meta-analysis. Studies
1787 comparing the use of the two FSH preparations (p-FSH and rFSH) in GnRH antagonist cycles are not
1788 present to allow evaluation of this statement in such a setting.

1789 **RECOMBINANT FSH (rFSH) VS HIGHLY PURIFIED URINARY FSH (hp-FSH)**

1790 **Evidence**

1791 In a systematic review and meta-analysis¹², ovarian stimulation with rFSH was compared to hp-FSH
1792 (Bordewijk et al., 2019). No significant difference was found between rFSH and hp-FSH for ovarian
1793 stimulation for live birth rate (RR 1.03, 95% CI 0.90-1.18, 12 RCTs, 2458 women) or clinical pregnancy
1794 rate (RR 1.03, 95% CI 0.94-1.13, 21 RCTs, 4165 women).

¹² The Cochrane systematic review (van Wely et al., 2011) that was mentioned here in the 2019 version of the guideline was replaced by a more recent meta-analysis.



1795 These observations are in line with the findings of several other RCTs not included in the systematic
1796 review in GnRH agonist cycles (Gholami et al., 2010, Murber et al., 2011, Parsanezhad et al., 2017,
1797 Selman et al., 2010, Selman et al., 2013). Three RCTs including respectively 70, 127 and 160 women
1798 reported no significant difference in live birth rate between rFSH and hp-FSH (respectively 31.3% vs.
1799 31.4%; 16.1% vs. 18.4% and 40% vs. 22.5%) (Murber et al., 2011, Parsanezhad et al., 2017, Selman et
1800 al., 2013). Two RCTs reported no difference in clinical pregnancy rate between rFSH and hp-FSH
1801 (respectively 39.6% vs. 38.7% and 33.3% (21/65) vs. 39% (23/60)) (Gholami et al., 2010, Selman et al.,
1802 2010).

1803 Two RCTs including respectively 84 and 160 women investigated the comparison of rFSH compared to
1804 hp-FSH in PCOS patients. There was no difference in clinical pregnancy rate (50% (21/42) vs. 50.2%
1805 (22/42) and 41.2% (33/80) vs. 45% (36/80)) or number of oocytes retrieved (13.83±7.07 vs. 17.1±8.66
1806 and 13.03±5.56 vs. 14.17±4.89) between both groups (Aboulghar et al., 2010, Sohrabvand et al., 2012).
1807 Sohrabvand et al. also reported no difference in live birth rate (21.3% (17/80) vs. 23.8% (19/80)), slight
1808 OHSS (5% (4/80) vs. 6.3% (5/80)) or moderate to severe OHSS (2.5% (2/80) vs. 2.5% (2/80)) between
1809 groups (Sohrabvand et al., 2012).

1810 Recommendation

The use of recombinant FSH (rFSH) and highly purified FSH (hp-FSH) for ovarian stimulation in GnRH agonist protocol is equally recommended. [2019]

Strong

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1811

1812 Justification

1813 In patients undergoing ovarian stimulation, the use of hp-FSH is not preferable to rFSH, when
1814 downregulation is achieved by GnRH agonists according to a Cochrane meta-analysis and confirmed in
1815 subsequently published studies. Studies comparing the use of the two FSH preparations (hp-FSH and
1816 rFSH) in GnRH antagonist cycles are not present to allow evaluation of this statement in such a setting.

1817 Studies for this question in PCOS patients were limited, so that a potential difference between
1818 compounds in this subgroup cannot be ruled out based on the current evidence.

1819 RECOMBINANT (rFSH) VS RECOMBINANT FSH + RECOMBINANT LH (rFSH+rLH)

1820 Evidence

1821 A Cochrane meta-analysis including 499 women found insufficient evidence to determine if there was
1822 a difference in patients treated with rFSH+rLH compared to those treated with rFSH only (4 RCT, OR
1823 1.32, 95% CI 0.85-2.06) (Mochtar et al., 2017). In a subgroup analysis in patients treated with GnRH
1824 agonists, although no difference has been observed in live birth rates between the two treatment
1825 groups compared (3 RCT, OR 1.73, 95% CI 0.95-3.16, 259 women), a higher probability of ongoing
1826 pregnancy has been observed with rLH addition (12 RCT, OR 1.27, 95% CI 1.02-1.57, 1980 women). The
1827 meta-analysis reported no difference in OHSS rate with rLH supplementation to rFSH compared to rFSH
1828 alone (6 RCT, OR 0.38, 95%CI 0.14-1.01, 2178 women). In a subgroup analysis in patients treated with
1829 GnRH agonists, a lower probability of OHSS has been observed with rLH addition (Mochtar et al., 2017).



1830 An RCT, more recent than the meta-analysis, including 238 women also reported no difference in live
1831 birth rate with rLH supplementation to rFSH (RR 0.78, 95% CI 0.4-1.53) (Lahoud et al., 2017).

1832 In a sub-analysis of the meta-analysis, a small RCT in poor responders showed a beneficial effect of rLH
1833 pre-treatment to rFSH on live birth rate (OR 9.33, 95% CI 1.03-84.20, 43 women) (Ferraretti et al., 2014,
1834 Mochtar et al., 2017). However, a large RCT (939 women), more recent than the meta-analysis,
1835 reported no effect of rLH addition to rFSH in Bologna poor responders on live birth rate (10.6% (49/462)
1836 vs. 11.7% (56/477)) (Humaidan et al., 2017). In this trial, only one event of mild early OHSS occurred in
1837 the rFSH+rLH group.

1838 A systematic review and meta-analysis focussing on women of advanced age (≥ 35 years) on the effect
1839 of rLH supplementation to rFSH in fresh IVF cycles included 12 RCTs and 1821 participants (Conforti et
1840 al., 2021). Live birth rates were evaluated in only two RCTs, and no differences were detected between
1841 ovarian stimulation with rLH supplementation and rFSH alone (OR 1.53, 95% CI 0.50-4.65, 2 RCT, 371
1842 women). Similarly, no significant differences were seen for clinical pregnancy rate (OR 1.11, 95% CI
1843 0.89-1.38, 11 RCT, 1670 women) and number of oocytes retrieved (MD -0.47, 95% CI -1.07 to +0.12, 7
1844 RCT, 997 women).

1845 Recommendation

The combination of rFSH with rLH and rFSH alone are probably equally recommended for the general IVF population. [updated]	Conditional ⊕⊕○○
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1846

The combination of rFSH with rLH and rFSH alone are probably equally recommended for low responders. [updated]	Conditional ⊕⊕○○
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1847

The combination of rFSH with rLH and rFSH alone are probably equally recommended for women of advanced age (≥ 35 year). [updated]	Conditional ⊕⊕○○
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1848

1849 Justification

1850 According to the best available evidence, the combination of rFSH with rLH results in similar live birth
1851 rates compared to rFSH alone.

1852 Current evidence from a large RCT in low responders indicated no beneficial effect of the combination
1853 of rFSH with rLH and rFSH alone on live birth rate.

1854 Similarly, a systematic review and meta-analysis focussing on women of advanced age (≥ 35 years) found
1855 no evidence of a benefit of adding rLH to ovarian stimulation with rFSH (Conforti et al., 2021).

1856 The GDG would also like to point to the importance of 'simplicity of ovarian stimulation'. When
1857 comparing compounds, dosages or add-on treatments for ovarian stimulation in this guideline
1858 document, preference was always given to the more basic option, unless a clear benefit was shown.



1859 **RECOMBINANT (rFSH) VS RECOMBINANT FSH + HUMAN MENOPAUSAL GONADOTROPIN (hMG)**

1860 **Evidence**

1861 rFSH vs. rFSH+hMG

1862 An RCT compared the clinical efficacy of highly purified hMG (75 IU) combined with rFSH (75-150 IU;
1863 n=305) to rFSH alone (150-225 IU; n=305) on ovarian stimulation for IVF in a long GnRHa protocol (Shu
1864 et al., 2019). No significant difference was reported between ovarian stimulation with or without hMG
1865 supplementation for moderate/severe OHSS (3.3% (10/305) vs. 3.6% (11/305)), clinical pregnancy rate
1866 per initiated cycle (29.2% (89/305) vs. 23.9% (73/305)) or number of MII oocytes retrieved (10.6±5.7
1867 vs. 11.4±5.2).

1868 An RCT evaluated whether the addition of hMG (75 IU; n= 78) to rFSH (225-300 IU) during the early
1869 follicular phase of ovarian stimulation improves clinical outcomes compared to no supplementation
1870 (n=94) in group 4 Bologna poor responders with the long GnRH agonist or GnRH antagonist (97%)
1871 protocol (35-44 year) (Qiu et al., 2023). No significant difference was noted with hMG supplementation
1872 compared to no supplementation for ongoing pregnancy rate per completed cycle (26.1% (23/88) vs.
1873 27.1% (19/70)) or clinical pregnancy rate per completed cycle (29.5% (26/88) vs. 28.6% (20/70)).

1874 Long-acting rFSH vs. long-acting rFSH + mid-follicular hMG

1875 In an RCT, women underwent ovarian stimulation with long-acting rFSH, in combination with either
1876 hCG (150 IU) or hMG (225 IU) starting from day 7 of stimulation until final oocyte maturation in the
1877 GnRH antagonist protocol (Decler et al., 2020). There were no significant differences between hCG
1878 and hMG supplementation for live birth rate (fresh+frozen; 11/61 vs. 9/67), clinical pregnancy rate
1879 (fresh+frozen; 15/61 vs. 12/67) or number of MII oocytes (6.6±4.4 vs. 6.1±4.8).

1880 An RCT compared the results of two ovarian stimulation protocols for IVF in patients at risk of low
1881 ovarian response: long-acting rFSH followed by hMG (300 IU; n=112) versus daily administration of hMG
1882 (300 IU; n=109) in a GnRH antagonist protocol (Taronger et al., 2018). There was no difference reported
1883 between the hMG/rFSH combination group and hMG only group for cumulative ongoing pregnancy
1884 rate and live birth rate (15.2% vs. 22%), ongoing pregnancy and live birth rate per started cycle (15.2%
1885 (17/112) vs. 20.2% (22/109)) or cumulative clinical pregnancy rate (19.6% (22/112) vs. 26.6% (29/109)).

1886 **Recommendation**

The combined use of recombinant FSH with human menopausal gonadotropin, either from the start or mid-phase of ovarian stimulation, is probably not recommended over the use of either recombinant FSH or hMG alone in normal and low responders. [2025]

Conditional ⊕⊕○○

1887

1888 **Justification**

1889 From only a handful studies it appears that, adding hMG either in the beginning of the stimulation with
1890 rFSH or after a rFSH stimulation period of 5-8 days, does not create any benefits in patients using either
1891 the GnRH agonist or antagonist pituitary suppression protocol.



1892 LONG-ACTING VS DAILY RECOMBINANT FSH

1893 Evidence

1894 In a systematic review¹³ and meta-analysis, RCTs were included of infertile women undergoing a single
1895 IVF/ICSI cycle with either long-acting or a conventional ovarian stimulation protocol based on daily
1896 injections (Cozzolino et al., 2019). No significant differences were seen between long-acting and daily
1897 rFSH for live birth rate/ongoing pregnancy rate (RR 0.92, 95% CI 0.80–1.05, 8 RCT, 4340 cycles) or
1898 incidence of overall OHSS (RR 1.15, 95% CI 0.83-1.57, 5 RCT, 3749 cycles) or moderate/severe OHSS (RR
1899 1.17, 95% CI 0.54-2.56, 4 RCT, 3349 cycles). However, significantly more oocytes were retrieved after
1900 ovarian stimulation with the long-acting formulation (MD 1.13, 95% CI +0.33 to +1.92, 5 RCT, 3848
1901 cycles).

1902 In an RCT, 283 women were randomly assigned to either rFSH-CTP (n=142) or rFSH groups (n=141) for
1903 ovarian stimulation in a GnRH antagonist protocol for IVF/ICSI (Wu et al., 2025). There was no significant
1904 difference in live birth rate (23.2% (33/142) vs. 29% (41/141)) or ongoing pregnancy rate (31.7% vs.
1905 36.9%) when comparing rFSH-CTP to rFSH. No cases of severe OHSS were reported in the rFSH-CTP
1906 group compared to 2 in the rFSH group.

1907 In an RCT, 117 women with poor ovarian response were randomly assigned to long-acting (n=59) or
1908 daily rFSH (n=58) for ovarian stimulation in a GnRH antagonist protocol for IVF/ICSI (Saharkhiz et al.,
1909 2024). The number of MII oocytes retrieved was significantly higher with long-acting rFSH compared to
1910 daily rFSH (5.0±2.1 vs. 4.2±1.7). However, there was no statistically significant difference in clinical
1911 pregnancy rate between long-acting and daily rFSH (28.8% vs. 22.0%).

1912 Recommendation

The use of long-acting and daily recombinant FSH (rFSH) is equally recommended in GnRH antagonist cycles for normal responders. [2019]

Strong

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1913

1914 Justification

1915 No differences have been observed in several large RCTs and in a small RCT in low responders regarding
1916 the probability of pregnancy, or the number of COCs retrieved and the incidence of OHSS.

1917 There are no controlled studies in high responders.

1918 The GnRH antagonist protocol is recommended for the use of long-acting rFSH.

1919 FOLLITROPIN DELTA

1920 Evidence

1921 *Follitropin delta requires the use of a dosing algorithm. There are no RCTs comparing individualised*
1922 *follitropin alpha/beta to individualised follitropin delta.*

¹³ The meta-analysis cited here in the 2019 version of the guideline is replaced by a more recent meta-analysis. The RCT by Kolibianakis et al., 2015 cited here in the 2019 version of the guideline is included in the new meta-analysis and therefore no longer mentioned separately.



1923 Recommendation

Follitropin delta and follitropin alfa/beta are probably equally recommended for ovarian stimulation. [2025]

Conditional ⊕○○○

1924

1925 Justification

1926 Several systematic reviews were published including 3 RCTs and 2682 women, comparing individualised
1927 follitropin delta compared to follitropin alfa/beta (Komiya et al., 2024, Nelson et al., 2024, Palomba et
1928 al., 2024). The live birth rates and ongoing pregnancy rates were found to be similar between women
1929 treated with follitropin delta compared to those treated with follitropin alfa/beta. However, the RCTs
1930 included in the systematic reviews include two interventions: a) different follitropin medications, and
1931 b) individualised versus fixed dosing. Therefore, it is uncertain that the effect on OHSS rate is due to the
1932 gonadotropin or the dosing regimen. Therefore, both gonadotropins are probably equally
1933 recommended.

1934 HIGHLY PURIFIED FSH (hp-FSH) VS HUMAN MENOPAUSAL GONADOTROPIN (hMG)

1935 Evidence

1936 Three RCTs including resp. 20, 80 and 218 women, compared hp-FSH with hMG for ovarian stimulation
1937 in the long GnRH agonist protocol and reported similar clinical pregnancy rate (10% (1/10) vs. 10%
1938 (1/10); 37.5% (15/40) vs. 45% (18/40) and 34% (35/104) vs. 36% (41/114)) and number of oocytes
1939 retrieved (8 (4-11) vs. 13 (4-23); 13.4±0.6 vs. 13.7±0.7 and 8.2±4.7 vs. 9.5±4.83) between both groups
1940 (Duijkers et al., 1993, Parsanezhad et al., 2017, Westergaard et al., 1996).

1941 Recommendation

The use of highly purified FSH (hp-FSH) and human menopausal gonadotropin (hMG) for ovarian stimulation in GnRH agonist protocols is equally recommended. [2019]

Conditional ⊕⊕○○

1942

1943 Justification

1944 In patients undergoing OS for IVF/ICSI, the use of hp-FSH does not appear to be preferable over hMG,
1945 if downregulation is achieved by GnRH agonists, according to three RCTs.

1946 HUMAN MENOPAUSAL GONADOTROPIN (hMG) VS RECOMBINANT FSH + RECOMBINANT LH (rFSH+rLH)

1947 Evidence

1948 In a small RCT including 122 patients undergoing ovarian stimulation with GnRH agonists, use of
1949 rFSH+rLH was not associated with increased pregnancy rate compared to hMG (28.3% (15/53) vs. 29.3
1950 (17/58)). However, significantly more cycles were cancelled to prevent OHSS in the rFSH+rLH group
1951 compared to the hMG group (11.1% (7/53) vs. 1.7% (1/58)) (Pacchiarotti et al., 2010).



1952 Recommendation

The use of recombinant LH (rLH)+recombinant FSH (rFSH+LH) for ovarian stimulation is probably not recommended over human menopausal gonadotropin (hMG) in GnRH agonist protocols with regards to safety. [2019]

Conditional ⊕○○○

1953

1954 Justification

1955 HMG and rFSH+LH appear to result in an equal probability of pregnancy in GnRH agonist protocols.
1956 However, the risk of OHSS appears to be higher with the use of rFSH+rLH. The recommendation is not
1957 applicable to GnRH antagonist cycles.

1958 GONADOTROPIN COMBINATION WITH HCG

1959 Evidence

1960 In a large RCT, addition of hCG to rFSH was investigated in women undergoing their first IVF/ICSI cycle
1961 in the long GnRH agonist protocol (Fernández Sánchez et al., 2022). hCG was administered in a fixed
1962 daily dose of 1 (n=104), 2 (n=101), 4 (n=99), 8 (n=107), or 12 µg (n=104) daily and compared to a control
1963 group receiving placebo (n=104) in 5 different injection volumes to match the injection volume of the
1964 different hCG dosages. The incidence of OHSS was lower in the hCG groups compared with the placebo
1965 group (2-6 cases per group vs. 12 in the control group) and the risk of OHSS was statistically significantly
1966 lower in the 12 µg dose group compared with the placebo group. The ongoing pregnancy rate was
1967 significantly lower in the 1 and 2 µg hCG groups compared to placebo (28.4% vs. 29.1% vs. 42.9%). No
1968 significant difference was seen with the higher dosages of hCG (4, 8, 12 µg) compared to placebo (39.2%
1969 vs. 37.4% vs. 30.4% vs. 42.9%). Significantly less MII oocytes were retrieved in all hCG treatment groups
1970 compared to placebo (8.2 vs. 8.3 vs. 8.0 vs. 8.4 vs. 7.3 vs. 9.7).

1971 In an RCT, supplementation with low-dose hCG (100 IU; n=40) to rFSH (200 IU) throughout stimulation
1972 was investigated and compared to placebo (n=41) in infertile women (35-40 years) undergoing IVF with
1973 a short GnRH agonist protocol (Siristatidis et al., 2022). Three cases of OHSS were noted in the study
1974 group (7.5%), compared to one in the control group (2.4%). No significant differences were seen when
1975 comparing the study and control groups for clinical pregnancy rate (25% (10/40) vs. 24.4% (10/41)) or
1976 number of MII oocytes retrieved (3 (IQR 5) vs. 3 (IQR 2)).

1977 In an RCT, hCG supplementation to rFSH (150 IU) from the start of stimulation at different dosages (50
1978 IU, n=15; 100 IU n=16; 150 IU, n=13) was compared to no supplementation (n=16) in the long GnRH
1979 agonist protocol (Thuesen et al., 2012). There were no cases of OHSS in the two highest dose groups of
1980 hCG, one case of moderate OHSS in the lowest hCG dose group and one case of mild OHSS in the control
1981 group. No significant differences were found when comparing the different hCG dosages (50, 100, 150
1982 IU) to no supplementation for cumulative live birth rate per started cycle (33% (5/15) vs. 44% (7/16) vs.
1983 39% (5/13) vs. 31% (5/16)) or live birth rate per started cycle (27% (4/15) vs. 25% (4/16) vs. 31% (4/13)
1984 vs. 25% (4/16)).



1985 An RCT investigated whether low-dose hCG added to rFSH (n=58) in regimens of ovarian stimulation
1986 could improve reproductive outcomes compared to the addition of rLH (n=56) in a GnRH agonist
1987 protocol in women aged 36-42 years, entering IVF-ET, especially in those women who had previous IVF
1988 failures (Drakakis et al., 2009). Clinical pregnancy rate per protocol was significantly higher with hCG
1989 supplementation compared to LH (27.6% (16/58) vs. 10.7% (6/56).

1990 In an RCT, the efficacy of low-dose hCG was investigated using a GnRH antagonist protocol (Koichi et
1991 al., 2006). All women were treated with purified urinary FSH (225-300 IU daily) until a follicular diameter
1992 of 14 mm was reached. Subsequently, the dose of purified urinary FSH was decreased (75 UI daily) and
1993 low-dose hCG (200 IU daily) and GnRH antagonist were initiated in the study group (n=63). In the control
1994 group (n=63), the purified urinary FSH dose was increased (300 IU daily) and GnRH antagonist was
1995 initiated. One case of severe OHSS was reported in both groups. No significant difference was seen for
1996 clinical pregnancy rate (39% (23/59) vs. 36.8% (21/57)).

1997 In an RCT, the efficacy of low-dose hCG was investigated using a GnRH antagonist protocol (Serafini et
1998 al., 2006). All women were treated with rFSH until a follicular diameter of 14 mm was reached.
1999 Subsequently, the dose of purified urinary FSH was decreased (75 UI daily) and low-dose hCG (200 IU
2000 daily) and GnRH antagonist were initiated in the study group (n=102). In the control group, the dosage
2001 of rFSH was continued and GnRH antagonist initiated (n=86). Three cases of OHSS were reported in the
2002 study group and four in the control group. No significant differences were reported between the study
2003 and control group for clinical pregnancy rate (54.9 (56/102) vs. 40.7% (35/86)) or number of MII oocytes
2004 (10.3±0.5 vs. 11.6±0.8).

2005 *Low responders*

2006 An RCT investigated the effect of late follicular (day 6) supplementation with low-dose hCG (100 IU,
2007 n=24 or 200 IU, n=23) on reproductive outcomes and compared them to rFSH alone (300 IU, n=26) in
2008 poor responder women undergoing ovarian stimulation for ICSI with a GnRH antagonist protocol
2009 (Madani et al., 2012). No significant differences were found between the 100 IU and 200 IU hCG groups
2010 and control group for live birth rate (14.3% (3/21) vs. 21.1% (4/19) vs. 13% (3/23)), clinical pregnancy
2011 rate (19.0% (4/21) vs. 26.3% (5/19) vs. 13% (3/23)) or number of MII oocytes retrieved (5.2±2.1 vs.
2012 5.2±4.4 vs. 3.4±1.7).

2013 In an RCT, the clinical effects of low-dose rhCG (75 IU) supplementation to rFSH (600 IU) in the
2014 midfollicular phase (n=48) were compared to stimulation with rFSH only (600 IU, n=51) in the GnRH
2015 agonist protocol for poor responders (Berkkanoglu et al., 2007). No significant differences were found
2016 in clinical pregnancy rate per transfer (21.8% vs. 27.1%) or number of MII oocytes retrieved (3.8±0.4 vs.
2017 5.6±0.7) between the rFSH and rhCG combination group and the rFSH only group.

2018 In an RCT, women underwent ovarian stimulation with long-acting rFSH, in combination with hCG (150
2019 IU) starting from day 7 of stimulation until final oocyte maturation in the GnRH antagonist protocol
2020 (Decler et al., 2020). There were no significant differences between hCG supplementation for live birth
2021 rate (fresh+frozen; 11/61 vs. 9/67), clinical pregnancy rate (fresh+frozen; 15/61 vs. 12/67) or number
2022 of MII oocytes (6.6±4.4 vs. 6.1±4.8).

2023 *High responders*

2024 In an RCT, the clinical effects of low-dose hCG supplementation from the start of ovarian stimulation
2025 with rFSH were investigated and compared to no hCG supplementation in PCOS patients in their first



2026 IVF/ICSI cycle with freeze-all (Zhu and Fu, 2019). All patients were treated with progesterone (100 mg
2027 daily) and hMG (150 IU daily), the study group also received low-dose hCG (200 IU every 3 days). There
2028 was no significant difference found between the study and control group for live birth rate per cycle
2029 (48.26% (14/29) vs. 35.48% (11/31)), clinical pregnancy rate per transfer (65.52% (19/29) vs. 41.94%
2030 (13/31)) or number of MII oocytes retrieved (13.55±6.56 vs. 13.4±6.34).

2031 Recommendation

Adding low dosages of hCG to the FSH stimulation is probably not recommended. [2025]	Conditional ⊕○○○
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2032

2033 Justification

2034 No significant benefit was observed for hCG supplementation during ovarian stimulation in the general
2035 population, low responders or in the one RCT including women with PCOS. Furthermore, there was
2036 large heterogeneity between studies for hCG dosing and timing of initiation.

2037 B. COMBINATIONS OF GONADOTROPINS WITH OTHER STIMULATION 2038 DRUGS

2039 LETROZOLE

2040 The combining of the aromatase inhibitor letrozole with gonadotropin during OS has been suggested
2041 as a method to reduce the total gonadotropin requirement in IVF. In recent years, the use of letrozole
2042 along with gonadotropins has grown, particularly in women predicted to respond poorly to OS
2043 (Goswami et al., 2004).

2044 Evidence

2045 Gonadotropin and letrozole combination

2046 *High responder*

2047 In a small RCT, the effect of letrozole (5 mg) in reducing the risk of OHSS was investigated in women
2048 with PCOS (n=27) and compared to placebo (n=28) (Ghasemi Tehrani et al., 2022). All women
2049 underwent ovarian stimulation with rFSH (150 IU daily) combined with hMG (75 daily) from day 4 of
2050 stimulation in the GnRH antagonist protocol. Patients in the study group received letrozole (5 mg) daily
2051 for 5 consecutive days, patients in the control group received placebo in an identical manner.
2052 Significantly less cases of moderate OHSS were seen in the letrozole group compared to placebo (1/25
2053 vs. 9/25). No significant difference was seen in clinical pregnancy rate with or without letrozole (60%
2054 (15/25) vs. 52% (13/25)).

2055 In an RCT, women with PCOS undergoing ovarian stimulation for ICSI were randomised to either receive
2056 combined letrozole (5 mg) and hMG (75 IU) (n=50) or hMG (75-225 IU) and placebo (n=50) in a GnRH
2057 antagonist protocol (Lotfy et al., 2022). No significant difference was found between letrozole and
2058 placebo supplementation for OHSS (2% (1/50) vs. 10% (5/50)), live birth rate (20% (10/50) vs. 28%
2059 (14/50)) or clinical pregnancy rate (46.0% (23/50) vs. 52.0% (26/50)).



2060 In an RCT, the clinical outcomes of PCOS patients at very high risk of OHSS undergoing ovarian
2061 stimulation with (n=24) or without (n=24) letrozole supplementation (5 mg) to gonadotropins (rFSH
2062 150 IU for 6 days followed by hMG 150 IU from day 4) were compared in a GnRH antagonist protocol
2063 (Tshzmachyan and Hambartsoumian, 2020). Significantly less cases of OHSS were reported in the study
2064 group (2 mild cases) compared to controls (9 mild cases and 1 moderate) (OR 7.86, 95% CI 1.49-41.3).
2065 However, live birth rate (33.3% (8/24) vs. 37.5% (9/24) and pregnancy rate per retrieval (58.3% (14/24)
2066 vs. 54.2% (13/24)) were comparable with and without letrozole for ovarian stimulation.

2067 In an RCT, it was investigated whether letrozole (2.5 mg daily) supplementation (n=65) to rFSH (100-
2068 225 IU) stimulation in a GnRH agonist protocol can positively influence the endometrial receptivity
2069 compared to conventional stimulation (n=65) in women with an expected high response to ovarian
2070 stimulation (Yang et al., 2019). No significant differences were reported with or without letrozole
2071 supplementation for incidence of OHSS (0 vs. 1.5% (1/65)) or live birth rate (42.9% (21/49) vs. 62.5%
2072 (30/48)).

2073 *Normal responder*

2074 In an RCT, the impact of letrozole co-treatment (rFSH 150 IU + Ltz 5 mg per day; n=67) on reproductive
2075 outcomes was investigated in expected normal responders and compared to placebo co-treatment
2076 (rFSH 150 IU + placebo; n=62) in the GnRH antagonist protocol (Bülow et al., 2022). No significant
2077 differences were found between letrozole co-treatment and placebo for live birth rate per woman
2078 randomised (24% (19/67) vs. 30% (24/62)), ongoing pregnancy rate per women randomised (26%
2079 (21/67) vs. 33% (26/62)) or number of MII oocytes retrieved per protocol (5.8±3.9 vs. 6.6±3.4). Similarly,
2080 there was no significant difference in cumulative clinical pregnancy rate after 4.8 years (38% (53/140)
2081 vs. 34% (50/147)) (Bülow et al., 2022, Bülow et al., 2023).

2082 An RCT compared the IVF outcomes of normal responders who have received gonadotropin both with
2083 (n=50) and without (n=50) the addition of letrozole (5 mg/day) from the start of stimulation until final
2084 oocyte maturation in the GnRH antagonist protocol (Eftekhar and Saeed, 2020). There was no
2085 significant difference with and without letrozole supplementation for incidence of OHSS (4% (2/50) vs.
2086 4% (2/50)). There was also no difference in clinical pregnancy rate (20.0% (10/50) vs. 22.0% (11/50)) or
2087 number of MII oocytes retrieved (8.46±4.73 vs. 6.96±4.09) with or without letrozole supplementation.

2088 A small RCT with only 20 patients randomized, investigated the addition of letrozole to FSH in an GnRH
2089 antagonist protocol for OS (Verpoest et al., 2006). No significant differences were reported in ongoing
2090 pregnancy rate (50% (5/10) vs. 20% (2/10)) or number of oocytes retrieved (13.8±9.2 vs. 9.6±7.7) in the
2091 letrozole + FSH group compared to the FSH only group (Verpoest et al., 2006).

2092 A small RCT including 94 women also investigated the addition of letrozole to FSH in an GnRH antagonist
2093 protocol for OS (Mukherjee et al., 2012). No differences were reported in clinical pregnancy rate (36%
2094 (15/42) vs. 33% (17/52)) or number of mature oocytes (4.6±2.5 vs. 4.9±2.3). There were no cases of
2095 OHSS in the letrozole group compared to 7 in the control group (Mukherjee et al., 2012).



2096 *Low responder*

2097 A systematic review and meta-analysis¹⁴ compared ovarian stimulation, with a combination of letrozole
 2098 and gonadotropins to gonadotropins alone in the GnRH antagonist protocol (Qin, 2021). The clinical
 2099 pregnancy rate (per cycle) was not statistically significant higher with administration of letrozole than
 2100 that in the control groups (RR 1.57, 95% CI 1.00–2.44, 6 RCT, 564 women). Furthermore, in low- (2.5
 2101 mg/day, 5 days) or high-dose (5 mg/day, 5 days) subgroups, no significant differences were indicated
 2102 in the clinical pregnancy rate with administration of letrozole compared to that in the control groups
 2103 (RR 1.65, 95% CI 0.85–3.18, 3 RCT, 270 women; RR 1.5, 95% CI 0.82–2.73, 3 RCT, 294 women).

2104 One RCT was found comparing the addition of letrozole with the addition of CC to gonadotropins in an
 2105 GnRH antagonist protocol in 184 poor responder women and reported no significant difference in
 2106 clinical pregnancy rate between groups (11.3% (9/87) vs. 8% (7/80)) (Eftekhari et al., 2014).

2107 Gonadotropin substitution by letrozole

2108 Three RCTs, including resp. 70, 20 and 50 women, investigated the effect of FSH substitution with
 2109 letrozole for OS (Ebrahimi et al., 2017, Verpoest et al., 2006, Yasa et al., 2013). Ebrahimi et al. and
 2110 Verpoest et al. reported no difference in clinical pregnancy rate with letrozole substitution compared
 2111 to no letrozole (resp. 14.3% (5/35) vs. 11.3% (4/35) and 50% (5/10) vs. 20% (2/10)) (Ebrahimi et al.,
 2112 2017, Verpoest et al., 2006). Yasa et al. reported no difference in ongoing pregnancy rate with letrozole
 2113 compared to no letrozole (20% (5/25) vs. 20% (5/25)) (Yasa et al., 2013).

2114 **Recommendation**

A stimulation scheme starting with gonadotropins followed by letrozole is probably not recommended over gonadotropins alone in low responders. [updated]	Conditional ⊕○○○
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2115

The addition of letrozole to gonadotropins in stimulation protocols for predicted high responders is probably not recommended. [updated]	Conditional ⊕○○○
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2116

The addition of letrozole to gonadotropins in stimulation protocols is probably not recommended for predicted normal responders. [2019]	Conditional ⊕○○○
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2117

The addition of letrozole to gonadotropins in stimulation protocols is probably not recommended for predicted low responders. [2019]	Conditional ⊕⊕○○
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¹⁴ The meta-analysis by Bechtejew et al., 2017 has been replaced by a more recent meta-analysis. The RCT by Ebrahimi et al., 2017 described here in the 2019 version of the guideline is included in the meta-analysis and therefore no longer described separately.



2118 Justification

2119 Due to the small number and size of RCTs available, no solid recommendation can be made for letrozole
2120 substitution of gonadotropins.

2121 Addition of letrozole to FSH in an GnRH antagonist protocol does not improve efficacy of OS in high,
2122 normal or low responders. The use of letrozole may reduce the risk of OHSS, however this was only
2123 shown in two small RCTs in high responders.

2124 In addition, safety concerns have been raised regarding possible teratogenicity associated with
2125 letrozole. The use of letrozole is off-label for OS.

2126 CLOMIPHENE CITRATE

2127 Evidence

2128 Gonadotropin and clomiphene citrate combination

2129 *High responder*

2130 In an RCT, women with PCOS undergoing ovarian stimulation for ICSI were randomised to either receive
2131 combined clomiphene citrate (5 mg) and hMG (75 IU) (n=50) or hMG (75-225 IU) and placebo (n=50)
2132 in a GnRH antagonist protocol (Lotfy et al., 2022). No significant difference was noted for OHSS rate (0
2133 vs. 10% (5/50)), live birth rate (24% (12/50) vs. 28% (14/50)) or clinical pregnancy rate (48% (24/50) vs.
2134 52.0% (26/50)) between clomiphene and placebo supplementation.

2135 In the prospective study by Saleh et al. (including 128 PCOS patients) the study group received a
2136 stimulation protocol consisting of CC, combined with a GnRH antagonist and rFSH, compared to GnRH
2137 antagonist with rFSH in the control group (Saleh et al., 2014). There was no significant difference in the
2138 clinical pregnancy rate (43.8% vs. 45.3%), number of oocytes retrieved (7.7 ± 1.3 vs. 8.1 ± 1.4) or number
2139 of mature oocytes (5.7 ± 1.1 vs. 6.1 ± 1.3) between the study group and the control group (Saleh et al.,
2140 2014).

2141 In the retrospective study by Jiang et al. (174 PCOS patients) the study group received a stimulation
2142 protocol consisting of CC combined with progestin protocol (MPA) and hMG, compared to MPA with
2143 hMG in the control group (Jiang and Kuang, 2017). There were significantly more oocytes retrieved (13
2144 (0–42) vs. 5 (0–30)) and mature oocytes (11 (0–35) vs. 4 (0–26)) in the control group as compared to
2145 the study group. There were no cases of moderate or severe OHSS in either group (Jiang and Kuang,
2146 2017).

2147 *Normal responder*

2148 A systematic review and meta-analysis¹⁵ investigated efficacy of ovarian stimulation with a combination
2149 of CC and reduced dose gonadotropins compared to conventional stimulation without oral medication
2150 (Datta et al., 2021). No significant difference was found between stimulation with CC and conventional
2151 gonadotropin stimulation for live birth rate (RR 0.88, 95 % CI 0.69-1.12, 3 RCTs, 573 women). However,

¹⁵ A more recent meta-analysis was found with the literature update of 2024, therefore the meta-analysis by Bechtejew et al., 2017 was removed.



2152 the risk of OHSS was significantly lower with the use of CC supplementation compared to the
2153 conventional (RR 0.12, 95% CI 0.03-0.51, 3 RCTs, 623 women).

2154 In an RCT, the effect of ovarian stimulation with (n=144) or without (n=132) clomiphene citrate (50 mg)
2155 supplementation to hMG stimulation (150 IU) was investigated in normal ovulatory women undergoing
2156 IVF/ICSI with the progestin-primed stimulation protocol (Liu et al., 2018). No significant differences
2157 were seen when comparing ovarian stimulation with or without clomiphene citrate for cumulative
2158 ongoing pregnancy rate per patient (60.6% (97/160) vs. 53.1% (85/160)), cumulative clinical pregnancy
2159 rate per patient (68.8% (110/160) vs. 66.9% (107/160)) or number of MII oocytes retrieved (8.71±5.28
2160 vs. 8.9±6.59).

2161 *Low responder*

2162 A systematic review and meta-analysis¹⁶ compared ovarian stimulation with a combination of
2163 clomiphene citrate and gonadotropins to gonadotropins alone, both in the GnRH agonist and
2164 antagonist protocol (Montoya-Botero et al., 2021). There was no significant difference in the clinical
2165 pregnancy rates (CC+GnRH antagonist vs conventional stimulation in GnRH agonist: RR 1.00, 95% CI
2166 0.96-1.04, 4 RCT, 1228 women; CC+GnRH antagonist vs conventional stimulation in GnRH antagonist:
2167 RR 1.00, 95% CI 0.93-1.08, 1 RCT, 77 women) or number of oocytes retrieved with clomiphene
2168 supplementation in the GnRH antagonist protocol versus conventional stimulation in the GnRH agonist
2169 protocol (MD -0.45, -1.49 to 0.59, 5 RCT, 1239 cycles) or conventional stimulation in the GnRH
2170 antagonist protocol (MD -0.59, -1.42 to 0.24, 1 RCT, 77 cycles).

2171 An RCT not included in the meta-analysis, also investigating the combination of CC and gonadotropins
2172 in an antagonist protocol in 250 poor responders. A significantly lower clinical pregnancy rate (5.9% vs.
2173 14.1%) was reported with CC addition compared to no CC, which was not associated with a difference
2174 in the number of oocytes retrieved (3.8 ± 2.9 vs. 3.41±1.9) (Schimberni et al., 2016).

2175 Gonadotropin substitution by clomiphene citrate

2176 *Studies comparing CC with the standard of care (FSH ovarian stimulation) are very scarce. We did not*
2177 *retrieve any RCTs comparing clomiphene citrate (CC) alone in high responders.*

2178 *Normal responder*

2179 One cohort study was identified, including 25 'good prognosis patients', comparing a protocol with
2180 clomiphene citrate addition to GnRH antagonist protocol. Significantly less oocytes were retrieved with
2181 the CC addition protocol (6.4±0.7 vs. 10.7±0.9). However, there was no difference in clinical pregnancy
2182 rate between CC addition and GnRH antagonist protocol (27.3% (6/22) vs. 49.0% (24/49) (Zander-Fox
2183 et al., 2018).

2184 *Poor responder*

2185 Only one RCT, including 249 poor responder women, has compared CC with a short GnRH agonist FSH
2186 protocol and showed similar live birth rate (5/145 vs. 7/146; RR 0.72, 95% CI 0.23-2.21) (Ragni et al.,
2187 2012).

¹⁶ The meta-analysis by Bechtejew et al., 2017 has been replaced by a more recent meta-analysis.



2188 **Conclusion**

2189 There is insufficient evidence available to recommend the substitution of FSH by Clomiphene Citrate in
2190 ovarian stimulation.

2191 **Recommendation**

The addition of Clomiphene Citrate to gonadotropins in stimulation protocols is probably not recommended for predicted high responders. [2019] Conditional ⊕⊕○○

2192

The addition of Clomiphene Citrate to gonadotropins in stimulation protocols is probably not recommended for predicted normal responders. [2025] Conditional ⊕⊕⊕○

2193

Clomiphene citrate alone or in combination with gonadotrophins, and gonadotropin stimulation alone are probably equally recommended for predicted low responders. [updated] Conditional ⊕⊕○○

2194

2195 **Justification**

2196 In women with normal ovarian response, current evidence shows no benefit in terms of efficacy with
2197 CC supplementation to gonadotropins. The systematic review reported a significantly lower OHSS rate,
2198 however, this is due to the lower dose of gonadotropins that was used in the CC and gonadotropins
2199 combination arm.

2200 In women with low ovarian response, no differences were reported in terms of safety and efficacy
2201 between CC alone, CC in combination with gonadotropins or gonadotropin stimulation alone.

2202 In women with high ovarian response, limited evidence shows no benefit of CC supplementation to
2203 gonadotropins in terms of efficacy.

2204

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- 2410



2411 7. Adjustment of gonadotropin dose

PICO QUESTION: IS ADJUSTMENT OF THE GONADOTROPIN DOSAGE DURING THE STIMULATION PHASE MEANINGFUL IN TERMS OF EFFICACY AND SAFETY?

2412 A systematic review studied the incidence of dose adjustments in clinical trials (Fatemi et al., 2021).
2413 Eighteen RCTs out of 1073 RCTs investigated were identified that reported dose adjustments: in 10
2414 RCTs (3952 cycles), dose increases were reported, in 11 RCTs (5123 cycles), dose reductions were
2415 reported and five RCTs reported unspecified dose changes (1359 cycles). However, the systematic
2416 review was unable to provide evidence of the impact of gonadotropin dose adjustments on clinical
2417 outcomes. These results are in agreement with a real-world study reporting on 33,962 ovarian
2418 stimulation cycles (23,582 patients), of which 40.7% had at least one dose adjustment. Among cycles
2419 with dose changes, 57.4% had at least one dose increase, 62.5% had at least one dose decrease, and
2420 19.9% of cycles included both increases and decreases (Mahony et al., 2021).

2421 Evidence

2422 An RCT investigated the effect of a modified flexible GnRH antagonist protocol by reducing rFSH dose
2423 by 30-50% as soon as the leading follicles reached 14 mm. Additionally, the GnRH antagonist
2424 administration was suppressed on final oocyte maturation day in the study group. The control group
2425 underwent a conventional flexible GnRH antagonist protocol (Xu et al., 2024). Comparing the modified
2426 to the conventional flexible GnRH antagonist protocol, a significantly higher live birth rate (38.1%
2427 (104/273) vs. 27.5% (75/273); RR 1.39 (1.09-1.77)) was seen. No significant differences were noted in
2428 risk of OHSS (1.1% (3/273) vs. 1.8% (5/273)) or number of MII oocytes (10.95±4.43 vs. 10.75±4.53))
2429 between the modified and conventional GnRH antagonist protocol.

2430 Another RCT investigated the effect of reducing the rFSH dose as soon as ≥ 3 follicles ≥ 14 mm were
2431 present until the criteria for final oocyte maturation were met (Lawrenz et al., 2021) and compared to
2432 conventional rFSH dosing. No significant difference was found in number of MII oocytes between the
2433 dose reduction group and the conventional dosing group (Lawrenz et al., 2021).

2434 An RCT including 151 women compared increasing hMG dose (with 75 IU) on the day of GnRH
2435 antagonist initiation with not increasing hMG dose and reported no difference in clinical pregnancy rate
2436 (36.2% vs. 32.1%, OR 1.3, 95% CI 0.63-2.6) or number of oocytes retrieved (9.2±2.1 vs. 10.1±3.8)
2437 between both groups (Aboulghar et al., 2004).

2438 A more recent retrospective study reported that changing the dose of gonadotropins during stimulation
2439 (increasing or decreasing) had no effect on clinical or ongoing pregnancy rates. Clinical pregnancy rate
2440 was 28.2% (11/39) with dose increase vs. 32.1% (27/84) with dose decrease vs. 25.8% (110/427) with
2441 no dose adjustments. Similarly, ongoing pregnancy rate was resp. 23.1% (9/39) vs. 25.0% (21/84) vs.
2442 22.5% (96/427) (Martin et al., 2006).

2443 Two RCTs investigated the effect of gonadotropin dose modulation in poor responder patients. Van
2444 Hooff et al. investigated the effect of doubling hMG dose on day 6 of OS in 47 low responders and
2445 reported no difference in pregnancy rate (2/25 vs. 1/22) or number of oocytes retrieved (4.7±1.0 vs.
2446 4.6±0.8). No cases of severe OHSS were reported (van Hooff et al., 1993). A more recent RCT including
2447 73 poor responders investigated the effect of reducing gonadotropin dose (step-down FSH protocol:
2448 450 IU starting dose, reduced to 300 IU/d when serum E2 values reached 200 pg/mL and again reduced



2449 to 150 IU/d when 2 follicles of 12 mm in diameter were detected on ultrasound) during OS and reported
2450 no difference in number of pregnancies (3/34 vs. 4/39) or number of oocytes retrieved (6.4±0.6 vs.
2451 6.3±0.6) (Cedrin-Durnerin et al., 2000).

2452 Aboulghar et al. investigated the effect of reducing hMG dose before coasting in 49 women at risk for
2453 developing OHSS. They found that reducing the hMG dose before coasting compared to not reducing
2454 hMG dose significantly reduced the duration of coasting (1.8±0.65 vs. 2.92±0.92 days) without
2455 influencing pregnancy rate (33.3% (8/25) vs. 35% 7/24) (Aboulghar et al., 2000).

2456 Recommendation

Adjustment (increase or decrease) of the gonadotrophin dose in the mid-stimulation phase during ovarian stimulation is probably not recommended. [2019]	Conditional ⊕○○○
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2457

Given the lack of evidence on the value of dose adjustments during ovarian stimulation, it is important that the gonadotropin starting dose is appropriate based on patient characteristics and desired outcome. [2025]	GPP
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2458

2459 Justification

2460 It is considered good practice to use ovarian reserve testing, patient preferences etc to determine the
2461 appropriate gonadotropin starting dose. The current evidence does not support changing gonadotropin
2462 dose during OS in the mid-stimulation phase. Modification (higher or lower) of gonadotrophin dose
2463 during ovarian stimulation for IVF/ICSI does not influence pregnancy rate. There is no evidence
2464 regarding dose modifications before the mid-stimulation phase during OS.

2465 The RCT by Xu et al. and Lawrentz et al are not specifically addressing the question, however, it is the
2466 best evidence found (Lawrenz et al., 2021, Xu et al., 2024).

2467

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- 2498

Draft for review



2499 8. Adjunct therapies

PICO QUESTION: IS THE ADDITION OF ADJUNCTS IN OVARIAN STIMULATION MEANINGFUL IN TERMS OF EFFICACY AND SAFETY?

2500 METFORMIN

2501 Evidence

2502 *Systematic reviews, meta-analyses of RCTs and RCTs comparing adjuvant metformin compared to*
2503 *control or placebo were considered for inclusion to address the efficacy and safety of metformin use*
2504 *during ovarian stimulation in IVF/ICSI treatment. All studies addressing the role adjuvant metformin*
2505 *were in women with PCOS.*

2506 A Cochrane systematic review and meta-analysis¹⁷ found no conclusive evidence that metformin before
2507 or during ovarian stimulation improves live birth rate compared to placebo/no treatment in women
2508 with PCOS (Tso et al., 2020). Substantial heterogeneity was found between studies, therefore the
2509 results were analysed based on the type of ovarian stimulation protocol. Six RCTs compared metformin
2510 to placebo/no treatment in a long GnRH agonist protocol, pooling of these RCTs showed no statistically
2511 significant evidence of improvements in live birth rate with metformin (OR 1.30, 95% CI 0.94-1.79, 651
2512 women). One RCT compared metformin to placebo/no treatment in a GnRH antagonist protocol and
2513 showed that metformin may reduce the live birth rate compared to placebo/no treatment (OR 0.48;
2514 95% CI 0.29-0.79, 153). A lower incidence of OHSS (severity of OHSS not specified) was found in the
2515 metformin group as compared to placebo/no treatment (11 RCT, RR 0.46; 95% CI 0.29-0.72, 1091
2516 women). The majority of the studies in the meta-analysis involved the use of GnRH agonist and only
2517 two studies used the GnRH antagonist protocol. Subgroup analysis based on the type of GnRH analogue
2518 showed only a significant difference in OHSS between the metformin group compared to control group
2519 when used with a long GnRH agonist protocol (9 RCT, OR 0.40, 95% CI 0.26-0.60), not with a GnRH
2520 antagonist protocol (2 RCT, OR 0.97, 95% CI 0.32-2.98, 193 women). The Cochrane meta-analysis also
2521 showed no significant difference in number of oocytes retrieved in the metformin compared to control
2522 group (11 RCT, MD 0.03; 95% CI -1.42 to 1.48) (Tso et al., 2020).

2523 An RCT, more recent than the Cochrane review, included 320 PCOS women randomised to receive
2524 either metformin (n=160) or placebo (n=160) during ovarian stimulation for IVF in a GnRH antagonist
2525 protocol (Hussein et al., 2021). This RCT reported that women receiving metformin had a significantly
2526 higher live birth rate (38.1% (61/160) vs. 27.5% (44/160) compared to placebo. One case of severe
2527 OHSS was reported in each group.

2528 Another RCT (102 PCOS women), not included in the Cochrane review, of metformin compared to
2529 placebo in an GnRH agonist protocol, reported no significant difference in live birth rate (25.5% (13/51)
2530 vs. 17.6% (9/51)) with adjuvant metformin compared to placebo treatment. However, significantly less
2531 oocytes were retrieved in the metformin group compared to placebo (9.06±4.23 16.86±8.3)
2532 (Abdalmageed et al., 2019).

¹⁷ The Meta-analysis by Tso et al. 2014 was replaced by the updated version. Jacob et al., 2016 is included in the updated meta-analysis and therefore no longer mentioned separately.



2533 Recommendations

Routine use of adjuvant metformin before and/or during ovarian stimulation is probably not recommended when using the GnRH antagonist protocol for women with PCOS. [updated]

Conditional



2534

2535 Justification

2536 The GDG recommends the use of GnRH antagonist for high responders and in women with PCOS. As
2537 current evidence does not show a beneficial effect of metformin in reducing OHSS when used with
2538 GnRH antagonist protocols and given the inconsistent evidence for live birth outcome, metformin is
2539 probably not recommended in women with PCOS.

2540 **GROWTH HORMONE (GH)**

2541 Evidence

2542 *Systematic reviews, meta-analyses of RCTs and RCTs comparing adjuvant growth hormone (GH)*
2543 *compared to control or placebo were considered for inclusion to address the efficacy and safety of GH*
2544 *use during ovarian stimulation in IVF/ICSI treatment.*

2545 Dose and administration of GH that was administered varied among studies from 4-12 IU
2546 subcutaneously daily to 4-24 IU on alternate days. The timing of GH administration varied between
2547 trials from daily administration pre-stimulation to alternate doses after the start of stimulation.

2548 GH for normal responders

2549 A Cochrane meta-analysis including 80 women considered as normal responder undergoing IVF
2550 treatment reported no significant difference in live birth rate (2 RCT, OR 1.32, 95% CI 0.40–4.43) with
2551 routine use of GH in women undergoing IVF treatment compared to placebo (Duffy et al., 2010). The
2552 updated Cochrane systematic review and meta-analysis included the same two RCTs in women
2553 considered as normal responders (Sood et al., 2021).

2554 An RCT included 288 normal responder women randomised to receive either GH adjunct therapy
2555 (n=144) compared to no adjunct treatment (n= 144) in a GnRH antagonist protocol (Mourad et al.,
2556 2025). There was no significant difference observed between adjunct GH treatment and no adjunct
2557 treatment for live birth rate after fresh transfer (32% (25/78) vs. 33% (30/90)) or clinical pregnancy rate
2558 after fresh transfer (44% (34/78) vs. 50% (45/90)). In addition, no significant difference was observed
2559 in the number of MII oocytes retrieved (8.5±6.2 vs. 8.6±6.3, ITT).

2560 GH for low responders

2561 A systematic review and meta-analysis¹⁸ investigated the effect of growth hormone supplementation
2562 on reproductive outcomes in women experiencing a poor ovarian response to stimulation (Liu et al.,
2563 2025). Comparing women receiving GH treatment to women receiving placebo/no treatment, a
2564 significantly higher live birth rate (OR 1.80, 95% CI 1.22-2.64, 9 RCTs, 945 women) and clinical pregnancy

¹⁸ The meta-analysis by Li et al., 2017 is replaced by the more recent Cochrane meta-analysis. The RCT by Choe et al., 2018 is included in this meta-analysis and therefore no longer mentioned separately.



2565 rate (OR 1.92, 95% CI 1.51-2.43, 19 RCTs, 1763 women) was observed. Furthermore, significantly more
2566 MII oocytes were retrieved in women receiving GH treatment (MD 1.63, 95% CI 1.13-2.13, 11 RCTs,
2567 1358 women).

2568 An RCT investigated the effect of GH co-treatment during ovarian stimulation on IVF outcomes in 158
2569 women who had at least one previous IVF cycle failure with no top-quality embryos (Li et al., 2020). Live
2570 birth rate was significantly higher in women receiving GH co-treatment compared to controls (41.1%
2571 (44/107) vs. 17.7% (9/51)). However, twelve women in the study group experienced OHSS, compared
2572 to only one in the control group.

2573 GH for PCOS

2574 An RCT investigated the effect of GH supplementation on reproductive outcomes in women with PCOS
2575 (Gong et al., 2020). No significant difference was found in clinical pregnancy rate between women with
2576 GH treatment versus controls (54% (27/50) vs. 42% (21/50)) or number of MII oocytes (12.30±6.80 vs.
2577 10.02±6.48).

2578 Recommendations

2579	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for normal responders. [2025]	Strong	⊕○○○
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2580	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for low responders. [updated]	Strong	⊕○○○
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2581	Use of adjuvant growth hormone before and/or during ovarian stimulation is not recommended for women with PCOS. [2025]	Strong	⊕⊕○○
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2582 Justification

2583 In general, there was a lack of data showing a beneficial effect. It is of great importance to point out
2584 that GH has the potential for serious harm and no long-term safety data are available. Furthermore, GH
2585 dosing schemes were very heterogenous. One new RCT in normal responders showed no benefit and
2586 the conclusion of the updated Cochrane review (Sood et al., 2021) was similar to the previous one
2587 (Duffy et al., 2010). Collective evidence from 2 small RCTs (included in meta-analysis by Duffy et al.)
2588 reported no effect on live birth rate (Duffy et al., 2010). One RCT involving women with PCOS showed
2589 no improvement in live birth rates following GH supplementation (Gong et al., 2020). There is collective
2590 evidence from the updated Cochrane review (Sood et al., 2021) and confirmed in the most recent meta-
2591 analysis (Liu et al., 2025) that adjuvant GH before and/ or during ovarian stimulation improves live birth
2592 rates in low responders following IVF treatment. Similar results were also reported by older meta-
2593 analysis (Duffy et al., 2010, Kolibianakis et al., 2009, Kyrrou et al., 2009, Li et al., 2017). Despite the



2594 possible beneficial effects in low responders on live birth rate, the evidence is of too limited quality to
2595 recommend GH during OS. The studies in the systematic review were generally underpowered and the
2596 definition of low response very heterogenous among studies. The GDG encourages further research on
2597 the use and dosing of GH in low responders, focussing on long-term safety data, both in the woman
2598 and offspring.

2599 TESTOSTERONE

2600 Evidence

2601 *Systematic reviews, meta-analyses of RCTs and RCTs comparing adjuvant testosterone pre-treatment*
2602 *compared to control or placebo were considered for inclusion to address the efficacy and safety of pre-*
2603 *treatment testosterone during ovarian stimulation in IVF/ICSI treatment. All studies addressing the role*
2604 *adjuvant testosterone were in predicted low responders.*

2605 Testosterone was administered transdermally, mostly as gel. Duration and dose of testosterone pre-
2606 treatment was either 12.5 mg/day of testosterone gel during pituitary downregulation, or testosterone
2607 gel 1%, with varying dose between studies between 10 mg/day to 12.5 mg/day preceding gonadotropin
2608 stimulation. Duration varied between studies ranging from 10-56 days.

2609 A Cochrane systematic review and meta-analysis¹⁹ studied the effect of testosterone pre-treatment
2610 versus placebo/no treatment (Naik et al., 2024). A significantly higher live birth rate/ongoing pregnancy
2611 rate was found in women pretreated with transdermal testosterone compared with those who were
2612 not (OR 2.53, 95% CI 1.61-3.99, 8 RCT, 716 women).

2613 In an RCT, 120 poor responder women were randomly assigned to receive methyltestosterone (n=60)
2614 or placebo pre-treatment (n=60) before ovarian stimulation for IVF/ICSI (Aliakbar et al., 2024). There
2615 was no significant difference in ongoing pregnancy rate (13.3% (8/60) vs. 3.3% (2/60)) or clinical
2616 pregnancy rate (15% (9/60) vs. 6.67% (4/60)) with testosterone pre-treatment compared to placebo.

2617 In a pilot RCT, not included in the meta-analysis, testosterone treatment administrated during ovarian
2618 stimulation in women experiencing poor ovarian response (Saharkhiz et al., 2018) showed that the
2619 pregnancy rate was significantly higher in the treatment group compared to controls (16% (4/25) vs.
2620 0% (0/23). Number of oocytes was also significantly higher in the study group vs controls (2.48±1.64 vs.
2621 1.17±1.27).

2622 Recommendations

Use of testosterone before ovarian stimulation is not recommended for low responders. [updated]

Strong



2623

¹⁹ The Cochrane systematic review and meta-analysis by Nagels et al., 2015 was replaced by a more recent systematic review. The RCTs by Kim et al., 2014 and Bosdou et al., 2016 are included in this meta-analysis and therefore no longer mentioned separately.



2624 **Justification**

2625 There is currently inconsistent evidence that testosterone pre-treatment before ovarian stimulation
2626 improves ovarian response in terms of number of oocytes retrieved and clinical outcomes of live birth
2627 rates in low responders undergoing IVF treatment. Also, due to insufficient data on dosage,
2628 administration duration and safety we cannot recommend testosterone use until a large RCT has been
2629 conducted.

2630 **DEHYDROEPIANDROSTERONE (DHEA)**

2631 **Evidence**

2632 *Systematic reviews, meta-analyses of RCTs and RCTs comparing adjuvant Dehydroepiandrosterone*
2633 *(DHEA) compared to control or placebo were considered for inclusion to address the efficacy and safety*
2634 *of DHEA use during ovarian stimulation in IVF/ICSI treatment.*

2635 The dose of DHEA used was 75 mg/day and varied in duration, starting either 6, 8 or 12 weeks before
2636 the start of ovarian stimulation and continued during ovarian stimulation. Most studies started DHEA
2637 12 weeks prior to ovarian stimulation.

2638 A systematic review and meta-analysis²⁰ investigated the effects of DHEA priming in women undergoing
2639 ovarian stimulation for IVF/ICSI (Huang et al., 2025). No significant difference was found between DHEA
2640 treatment or placebo/no treatment in live birth rate (OR 1.33, 95% CI 0.98-1.82, 10 RCTs, 1217 women).
2641 DHEA pre-treatment did also not increase the number of MII oocytes retrieved (MD 0.56, CI -0.06 to
2642 1.18, 8 RCTs, 842 women).

2643 **Recommendations**

Use of DHEA before and/or during ovarian stimulation is not recommended for low responders. [updated]

Strong ⊕⊕○○

2644

Use of DHEA before and/or during ovarian stimulation is not recommended for normal responders. [2025]

Strong ⊕⊕○○

2645

2646 **Justification**

2647 The systematic review including 16 RCTs showed that adjuvant DHEA use before and during ovarian
2648 stimulation does not improve live birth/ongoing pregnancy rate (Huang et al., 2025). Two RCTs involving
2649 normal responders showed that DHEA use before and during ovarian stimulation did not improve
2650 clinical pregnancy rates and number of oocytes retrieved (Mostajeran et al., 2018, Yeung et al., 2016).
2651 The studies varied in duration of DHEA treatment, possibly contributing towards the inconsistency in

²⁰ The Cochrane systematic review and meta-analysis was replaced by a more recent systematic review. The RCTs by Kotb et al., 2016, Narkwicheckan et al., 2017, Mostajeran et al., 2018 and Yeung et al., 2016 are included in this meta-analysis and therefore no longer mentioned separately.



2652 observed results. Also, due to insufficient data on administration duration and safety we cannot
 2653 recommend DHEA use until a large RCT has been conducted.

2654 **ASPIRIN**

2655 **Evidence**

2656 *To address the efficacy and safety of adjuvant aspirin use with ovarian stimulation in IVF/ICSI treatment,*
 2657 *studies were selected if aspirin was used before and/or during ovarian stimulation. Studies commencing*
 2658 *aspirin after ovarian stimulation were excluded. Systematic reviews, meta-analyses and eligible RCTs*
 2659 *(not included in the selected systematic reviews or meta-analyses) comparing adjuvant aspirin alone*
 2660 *(without other co-interventions) compared to control or placebo were included.*

2661 Doses of aspirin used in the studies varied between 75 mg daily, 80 mg daily or 100 mg daily and aspirin
 2662 was continued until hCG administration for final oocyte maturation, 12 weeks of pregnancy or until
 2663 delivery.

2664 A Cochrane meta-analysis combining 3 RCTs with 1053 women reported no significant difference in the
 2665 live birth rate (3 RCT, RR 0.91, 95% CI 0.72-1.15) or ongoing pregnancy rate (2 RCT, RR 0.94, 95% CI
 2666 0.69-1.27) between the aspirin and control group (Siristatidis et al., 2016). Due to technical limitations
 2667 of the meta-analysis to specifically address the role of adjuvant aspirin use before and/or during ovarian
 2668 stimulation, all other outcomes were assessed from individual studies.

2669 Results from 4 RCTs in the general IVF/ICSI population showed that adjuvant aspirin has no beneficial
 2670 effect on the number of oocytes retrieved (Table 7) (Dirckx et al., 2009, Lambers et al., 2009, Moini et
 2671 al., 2007, Pakkila et al., 2005). One RCT, Rubinstein et al. reported a significantly higher number of
 2672 oocytes with aspirin compared to placebo treatment (16.2±6.7 vs. 8.6±4.6) (Rubinstein et al., 1999).

2673 There was one RCT including poor responders which demonstrated no significant difference in number
 2674 of oocytes retrieved and clinical pregnancy rate between the aspirin compared to control group (Lok et
 2675 al., 2004).

2676
 2677 An RCT investigated the effect of pre-treatment with low-dose aspirin on the risk of OHSS in the long
 2678 GNRH agonist protocol in 232 women with PCOS (Namavar Jahromi et al., 2019). No significant
 2679 difference was found between aspirin and placebo pre-treatment for moderate to severe OHSS (34.9%
 2680 (38/109) vs. 34.9% (38/109)) and clinical pregnancy rate (28.4% (31/109) vs. 22.9% (24/105)).

2681 *Table 8: Number of oocytes retrieved.*

Study	Cohort (n)	Aspirin	Placebo
Dirckx 2009	193	12.6 ± 7.6	12.9 ± 7.9
Lambers 2009	169	13.7	13.5
Moini 2007	145	6.9 ± 5.6	8.6 ± 6.8
Pakkila 2005	374	12.0 ± 7.0	12.7 ± 7.2
Lok 2004	60	3.0 (2.0–7.25)	4.0 (3.0–7.25)
Rubinstein 1999	298	16.2 ± 6.7	8.6 ± 4.6

2682



2683 Recommendation

Use of aspirin before and/or during ovarian stimulation is not recommended in the general IVF/ICSI population nor for low responders. [2019]

Strong ⊕⊕⊕○

2684

2685 Justification

2686 The existing evidence suggests that adjuvant aspirin before and/ or during ovarian stimulation does not
2687 improve ovarian response in terms of number of oocytes retrieved and clinical outcomes of clinical or
2688 ongoing pregnancy, or live birth rates following IVF treatment.

2689 Evidence could not be formulated on the outcome of OHSS due to poor study quality and reporting
2690 method (Varnagy et al., 2010).

2691 **INDOMETACIN**

2692 Evidence

2693 Current evidence is limited to one case report (Nargund and Wei, 1996).

2694 Conclusion

2695 There are no controlled studies nor RCT addressing the efficacy and safety of adjuvant indomethacin
2696 use during ovarian stimulation in IVF treatment. Thus, there is no evidence to recommend the use of
2697 indomethacin during OS.

2698 **SILDENAFIL**

2699 Sildenafil is used in ovarian stimulation to increase ovarian vascularization and hence increase live birth.

2700 Evidence

2701 Studies on sildenafil administered (for improving endometrial thickness) after oocyte pick-up were not
2702 included.

2703 A small RCT evaluated the effect of vaginal sildenafil during ovarian stimulation on IVF success rate in
2704 72 women (Tehraninejad et al., 2018). No significant difference was found between the study group
2705 and the control group for clinical pregnancy rate (33.3% (12/36) vs. 27.8% (10/36)).

2706 A small pseudo-randomised RCT including 60 patients classified as poor responders reported no
2707 significant difference in the clinical pregnancy rate (16.7% (5/30) vs. 13.3% (4/30)) or number of oocytes
2708 retrieved between the sildenafil and control group (3.95±1.40 vs. 3.65± 1.14) (Ataalla et al., 2017).

2709 Recommendations

Use of sildenafil before and/or during ovarian stimulation is not recommended for low responders. [2019]

Strong ⊕○○○

2710



2711 **Justification**

2712 Current evidence from one low-quality, pseudo-randomized study involving women considered as low
2713 responders undergoing IVF showed no improvement in ovarian response with adjuvant sildenafil use
2714 during ovarian stimulation. Furthermore, a Dutch trial using sildenafil to try to correct foetal growth
2715 restriction (STRIDER study) has been halted after 11 babies subsequently died (Ganzevoort et al., 2014,
2716 Hawkes, 2018).

2717 **ANTI-OXIDANTS (MYO-INOSITOL)**

2718 **Evidence**

2719 A Cochrane systematic review and meta-analysis investigated the effect of inositol on IVF outcomes in
2720 women with PCOS (Showell et al., 2018). The start and duration of pre-treatment varied between eight
2721 to twelve weeks before IVF/ICSI treatment. The treatment period for one study lasted from the first
2722 day of the cycle to 14 days after embryo transfer, and another study started treatment on the first day
2723 of GnRH agonist administration. No significant difference in live birth rates have been found with myo-
2724 inositol compared to standard treatment (folic acid) (2 RCT, OR 2.42; 95% CI 0.75-7.83; 84 women).

2725 An RCT investigated the effect of myo-inositol pre-treatment on pregnancy outcomes in 60 women
2726 referred for IVF (Seyedshohadaei et al., 2022). Live birth rate was significantly higher in women
2727 receiving myo-inositol pre-treatment compared to standard treatment (folic acid) (26.7% (8/30) vs. 10%
2728 (3/30)). Similarly, clinical pregnancy rate (56.7% (17/30) vs. 23.3% (7/30)) and the number of MII
2729 oocytes (7.53±3.71 vs. 5.43±2.50) were higher in the study group compared to controls.

2730 An RCT investigated the effect of myo-inositol treatment before and during ovarian stimulation on IVF
2731 outcomes in non-PCOS women (Lisi et al., 2012). There was no significant difference in clinical
2732 pregnancy rate between myo-inositol treatment and standard treatment (folic acid) (28% (14/50) vs.
2733 24% (12/50)). However, significantly less MII oocytes were retrieved after myo-inositol treatment
2734 compared to standard treatment (4.8±2.2 vs. 6.3±2.9).

2735 Two RCTs investigated the effect of myo-inositol treatment before and during ovarian stimulation on
2736 reproductive outcomes respectively 60 and 112 women experiencing poor ovarian response to
2737 stimulation (Mohammadi et al., 2021, Nazari et al., 2020). No significant differences were found
2738 between women receiving myo-inositol and women receiving standard treatment (folic acid) for
2739 ongoing pregnancy rate (7.1% vs. 3.6%) (Nazari et al., 2020), clinical pregnancy rate (6.6% (2/30) vs. 0%
2740 (0/30)) or number of MII oocytes (2.36±1.64 vs. 1.87±1.07) (Mohammadi et al., 2021).

2741 **Recommendations**

Use of myo-inositol before and/or during ovarian stimulation is probably not recommended for women with PCOS undergoing IVF. [2025]	Conditional ⊕○○○
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2742

Use of myo-inositol before and/or during ovarian stimulation is not recommended in low responders. [2025]	Strong ⊕⊕○○
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2743

Use of myo-inositol before and/or during ovarian stimulation is not recommended in non-PCOS women undergoing IVF. [2025]

Strong ⊕⊕○○

2744

2745 Justification

2746 Studies varied in the duration of pre-treatment and timing of myo-inositol treatment. The Cochrane
2747 review concluded that it is uncertain whether myo-inositol improves live birth rates in women
2748 undergoing IVF (Showell et al., 2018). An RCT involving non-PCOS women undergoing IVF showed no
2749 improvement in clinical pregnancy rates but a lower number of MII oocytes in the myo-inositol group
2750 (Lisi et al., 2012). Two RCTs involving low responders undergoing IVF showed no improvement in the
2751 pregnancy rates and number of MII oocytes in the myo-inositol group (Mohammadi et al., 2021, Nazari
2752 et al., 2020). For non-PCOS women and low responders, there is no biological rationale for using myo-
2753 inositol to the treatment scheme.

2754

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- 2869



2870 9. Non-conventional start of ovarian stimulation

PICO QUESTION: WHAT IS THE SAFETY AND EFFICACY OF NON-CONVENTIONAL START STIMULATION COMPARED TO STANDARD EARLY FOLLICULAR PHASE STIMULATION?

2871 NON-CONVENTIONAL START

2872 Evidence

2873 A retrospective study in 150 normal responders reported comparable ongoing pregnancy rates (39.4%
2874 (13/33) vs. 33.3% (12/36) vs. 39.0% (16/41)) and number of oocytes retrieved (6.6 ± 3.8 vs. 5.9 ± 4.3 vs.
2875 5.9 ± 4.2) when stimulation was started in the late follicular or luteal phase as compared to conventional
2876 start (day 2-5) (Qin et al., 2016). Similarly, a more recent, large retrospective study in 1302 normal
2877 responders (non-oncologic fertility preservation) reported no difference in number of oocytes retrieved
2878 (12.7 ± 2.7 vs. 13.0 ± 3.1 vs. 13.2 ± 2.9 vs. 13.1 ± 2.3) between early follicular (day 4-7), late follicular (> day
2879 7), and luteal start stimulation as compared to conventional start (day 2/3) (Pereira et al., 2017).

2880 Recommendation

Random-start ovarian stimulation could be used when a fresh transfer is not intended and there is no possibility of natural conception. [Updated]

GPP

2881

2882 Justification

2883 Current evidence in normal responders reported no difference in efficacy in terms of number of oocytes
2884 retrieved with non-conventional start stimulation as compared to conventional (early follicular) start
2885 stimulation. This validates the feasibility of random-start protocols; however, freeze-all oocytes or
2886 embryos is mandatory. A medico-economic study is needed as non-conventional stimulation might
2887 require a higher consumption of FSH and the long-term child health has to be carefully monitored as
2888 the hormonal environment of the oocytes is modified. The risk of an undetected, natural conception
2889 pregnancy in non-conventional start stimulation is low (Lawrenz et al., 2024), however, they could lead
2890 to severe OHSS and hospitalisation (Semrl et al., 2024).

2891 LUTEAL PHASE STIMULATION

2892 Luteal phase stimulation can be regarded as an extension to urgent oncologic fertility preservation. A
2893 distinction must be made between gonadotropin pre-treatment in the luteal phase before follicular
2894 stimulation with fresh transfer, and ovarian stimulation in the luteal phase (day 15-19) with mandatory
2895 frozen oocytes/embryos.

2896 Evidence

2897 Late luteal gonadotropin start with intention of fresh transfer

2898 Three very small RCTs in poor ovarian reserve patients reported conflicting results on the number of
2899 oocytes retrieved (Kansal Kalra et al., 2008, Kucuk et al., 2008, Rombauts et al., 1998). A very small RCT



2900 (18 women) reported no difference in number of oocytes retrieved (5.0 (3-8) vs. 5.5 (1-14)) between
2901 gonadotropin pre-treatment and normal-start stimulation in GnRH antagonist protocol (Kansal Kalra et
2902 al., 2008). Another very small RCT (40 women) reported similar findings in the short GnRH agonist
2903 protocol, with median number of oocytes collected: 4.5 (2-12) in the experimental group vs. 6 (1-10) in
2904 the control group (Rombauts et al., 1998). However, another very small RCT (42 women) reported an
2905 increased number of mature oocytes (mean number: 6.8 vs. 3.2) with luteal gonadotropin pre-
2906 treatment as compared to the normal-start stimulation in the long GnRH agonist protocol (Kucuk et al.,
2907 2008).

2908 Luteal phase stimulation without fresh transfer

2909 A small RCT compared luteal phase stimulation (n=31) with follicular phase stimulation (n=33) in women
2910 with a poor ovarian response to stimulation (Dastjerdi et al., 2024). Significantly more MII oocytes were
2911 retrieved with luteal stimulation (3 (0-8)) compared to follicular stimulation (2 (0-5)). Eleven women in
2912 both groups proceeded with embryo transfer resulting in 1 clinical pregnancy in the study group and
2913 none in the control group.

2914 Another RCT investigated the effect of luteal phase stimulation in women with a poor ovarian response
2915 to stimulation (Suñol et al., 2023). In the ITT analysis, the mean number of MII oocytes retrieved was
2916 not different between the FPS and LPS groups (5.4±3.6 vs. 5.2±2.8).

2917 Follicular versus luteal phase stimulation in double ovarian stimulation

2918 An RCT compared double stimulation in one menstrual cycle (n=23) with one conventional ovarian
2919 stimulation cycle (n=23) (Boudry et al., 2024). The mean number of MII oocytes retrieved after follicular
2920 stimulation was 3.0±2.2 compared to 2.4±2.2 after luteal stimulation. However, the cancellation rate
2921 due to insufficient response for the second oocyte retrieval was 39.1% (9/23).

2922 An RCT compared double stimulation in one menstrual cycle (n=21) with one conventional ovarian
2923 stimulation cycle (n=21) (Saharkhiz et al., 2024). The mean number of MII oocytes retrieved after
2924 follicular stimulation was 1.63 ± 1.40 compared to 1.72 ± 1.72 after luteal stimulation.

2925 An RCT compared double stimulation (n=44) with 2 conventional ovarian stimulation cycles (n=44) using
2926 a GnRH antagonist protocol in women experiencing a poor ovarian response to stimulation (Massin et
2927 al., 2023). There was no significant difference in the mean number of MII oocytes retrieved after
2928 follicular and luteal stimulation (2.4±2.3 vs. 2.2±1.7). The cumulative cycle cancellation rate was similar
2929 in both groups 7.7% vs. 4.9%.

2930 An RCT investigated the efficacy of double stimulation (n=28) in PGT-A cycles, compared to two
2931 conventional stimulation cycles (n=28) in women with a poor prognosis (Cerrillo et al., 2023). There was
2932 no significant difference in the mean number of MII oocytes retrieved after follicular and luteal
2933 stimulation (3.3±1.0 vs. 3.6±1.2).

2934 Recommendations

Luteal start ovarian stimulation could be used when a fresh transfer is not intended and there is no possibility of natural conception. [updated]

Conditional ⊕○○○

2935



Late luteal phase start of gonadotropins with fresh transfer is probably not recommended for low responders. [updated]

Conditional ⊕○○○

2936

2937 Justification

2938 Mention should be made about late luteal gonadotropin start protocol (before menstruation), that can
2939 also be considered as gonadotropin pre-treatment. It has been used with intention of fresh transfer.
2940 Results are inconclusive and based on very little studies with very small study populations.

2941 The quality of evidence is very low and controversial regarding the luteal start of FSH in normal and low
2942 responders, and there are no data for PCOS patients. However, the oocyte competence is probably not
2943 impacted by its luteal phase origin compared to follicular phase. Absence of adverse effects on neonatal
2944 outcomes and long-term child health needs to be evaluated on a larger scale.

2945 An potential disadvantage of the luteal start stimulation is the mandatory freeze-all of oocytes or
2946 embryos. One study reported on neonatal outcomes comparing frozen/thawed from follicular and
2947 luteal phase stimulation (Chen et al., 2015). Therefore, luteal phase stimulation could be considered as
2948 an option in specific cases, for organization and shortened time to oocyte retrieval, for example in
2949 urgent oncologic fertility preservation, as well as in freeze-all policy programs.

2950 Also, the drug marketing approval for gonadotropin use in luteal phase needs to be considered.

2951 DOUBLE STIMULATION

2952 Double stimulation or “dual stimulation” or “duostim” (Vaiarelli et al., 2018) or “Shanghai protocol”
2953 (Kuang et al., 2014) is experimented in low responder patients or in urgent oncologic fertility
2954 preservation. It corresponds to the sequencing of 2 stimulation protocols within the same menstrual
2955 cycle: first in the follicular phase then second, immediately after the oocyte pick up, in the luteal phase
2956 of the same cycle. So, two oocyte pick-ups are performed at approximately 2 weeks apart. This protocol
2957 uses the physiological principles of multiple waves of folliculogenesis within one cycle (Baerwald et al.,
2958 2003). It allows to recover more oocytes in a shorter time period. As shown in luteal phase stimulation
2959 protocols, the quality of oocytes retrieved in the second stimulation seems as good as the ones
2960 retrieved in the first stimulation (same euploid embryo rate) (Vaiarelli et al., 2018).

2961 Evidence

2962 An RCT compared double stimulation in one menstrual cycle (n=44) with 2 conventional ovarian
2963 stimulation cycles (n=44) using a GnRH antagonist protocol in women experiencing a poor ovarian
2964 response to stimulation (Massin et al., 2023). No significant difference between double stimulation and
2965 2 conventional stimulations for cumulative live birth rate (17.9% (7/39) vs. 34.1% (14/41)) or number
2966 of MII oocytes (2.4±2.3 vs. 2.5±2.7). No serious adverse events were reported.

2967 Another RCT investigated the efficacy of double stimulation (n=28) in PGT-A cycles, compared to two
2968 conventional stimulation cycles (n=28) in women with a poor prognosis (Cerrillo et al., 2023). No
2969 significant differences were found with double stimulation or two conventional stimulations for live
2970 birth rate (19.5% (8/41) vs. 23.1% (9/39)), pregnancy rate (24.4% (10/41) vs. 23.1% (9/39)) or MII



2971 oocytes (6.8±1.7 vs. 8.7±1.8). The study was ended prematurely because of a high probability that no
2972 statistical differences would be confirmed at the end of study.

2973 **Recommendation**

Double stimulation can be considered for urgent fertility preservation cycles [2019]

GPP

2974

Double stimulation can be used with the intention to accumulate oocytes or embryos when fresh transfer is not planned. [updated]

Strong

⊕⊕○○

2975

2976 **Justification**

2977 Two RCTs show that there is no benefit of double stimulation over two conventional stimulation cycles.
2978 There is a chance of pregnancy after the first ovarian stimulation as shown by the RCT by Boudry *et al.*,
2979 (Boudry *et al.*, 2024). In double stimulation, this would lead to an unnecessary second ovarian
2980 stimulation cycle.

2981

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PART D: Fertility preservation and oocyte donation

10. Ovarian stimulation for fertility preservation

PICO QUESTION: WHAT IS THE PREFERRED STIMULATION PROTOCOL FOR FERTILITY PRESERVATION IN PATIENTS FACING GONADOTOXIC TREATMENT?

Fertility preservation represents a major issue for young women suffering from diseases that might impact their reproductive potential (Recommendations ASCO, ISFP). OS followed by oocyte or embryo vitrification constitutes the best option. Collecting as much oocytes as possible, sometimes in an extremely reduced time frame represents an important issue. Fertility preservation has emerged relatively recently in the field of reproductive medicine. Therefore, many questions raised, in particular regarding the preferred protocol and the feasibility of random-start ovarian stimulation. In addition, the specificity of OS performed in contexts of oestrogen-sensitive diseases has led, in the name of the precautionary principle, to the development of protocols using anti-oestrogen therapies. Considering the motivation for this treatment, critical and important outcomes in this chapter are different from the rest of this guideline. Critical outcomes for fertility preservation in this guideline are the number of oocytes/embryo's and preventing OHSS and other complications.

More information and recommendations on female fertility preservation for women with cancer, benign diseases, and also transgender patients and women undergoing elective freezing, is covered in the ESHRE guideline on female fertility preservation (www.eshre.eu/FFPguideline).

INITIATION OF STIMULATION

Evidence

Random-start

A systematic review and meta-analysis²¹, including 2 prospective observational and 9 retrospective studies, compared random (688 cycles) and conventional start (1076 cycles) protocols for ovarian stimulation in cancer patients seeking fertility preservation (Sönmezer et al., 2023). No significant difference was found between random and conventional start of stimulation for number of MII oocytes retrieved (SMD -0.11, 95% CI -0.44 to 0.21, 6 studies, 787 cycles) and number of embryos frozen (SMD -0.04, 95% CI -0.28 to 0.20, 5 studies, 673 cycles).

In a prospective cohort study, ovarian stimulation was started irrespective of the menstrual cycle (early follicular, n=43; late follicular, n=17; or luteal, n=35) (Dezellus et al., 2024). The number of MII oocytes cryopreserved was not statistically different irrespective of the menstrual cycle phase (early follicular

²¹ The cohort study by Muteshi et al., 2018 is included in the new meta-analysis and therefore no longer mentioned separately.



3081 10.0±7.3 vs. late follicular 7.7±4.0 vs. luteal 10.4±5.3). Seven embryo transfers with frozen-thawed
3082 oocytes were performed among five patients, none resulted in pregnancy.

3083 In a retrospective cohort study, conventional start stimulation in the early follicular phase (n=176) was
3084 compared with the late follicular phase (n=8) start of ovarian stimulation for fertility preservation (Baig
3085 et al., 2023). No significant difference was found comparing early with late follicular phase start of
3086 stimulation for number of MII oocytes retrieved (9.0 (6.0-13.0) vs. (7.0 (2.3-13.3)).

3087 In a retrospective cohort study, the cycle characteristics and outcomes of random-start ovarian
3088 stimulation (n=39) protocols were compared to the outcomes of conventional-start ovarian stimulation
3089 (n=117) cycles for women with breast cancer undergoing fertility preservation (Turan et al., 2023). The
3090 mean number of MII oocytes retrieved (10.9±4.2 vs. 10.1±5.8) and number of embryo's cryopreserved
3091 (77±4.0 vs. 7.7±4.8) was similar with random-start and conventional start stimulation. To date, seven
3092 women returned to utilize their cryopreserved embryos after RSCOS. Of those, six were conceived after
3093 the first single embryo transfer.

3094 In a retrospective cohort study, random-start (n=36) was compared to conventional follicular start
3095 (n=25) in breast cancer patients undergoing ovarian stimulation for fertility preservation (Sahin et al.,
3096 2022). All patients received letrozole during ovarian stimulation independent of oestrogen receptor
3097 status. Random or follicular start of the ovarian stimulation did not significantly influence the total
3098 number of oocytes retrieved (10.9±6.9 vs. 11.5±9.3) or the number of MII oocytes retrieved (8.5±5.7
3099 vs. 7.0±5.6).

3100 Luteal start

3101 A systematic review of 8 (non-randomized) studies of which 6 were performed in context of fertility
3102 preservation, showed in 251 women, that number of oocytes recovered (WMD -0.6 oocytes, 95 % CI
3103 -2.8 to 1.6) did not differ whatever the phase of the cycle at which FSH was started. Interestingly,
3104 oocytes obtained in cycles initiated in the luteal phase fertilized more efficiently (WMD 0.16, 95 % CI
3105 0.13 to 0.19). No conclusion can be drawn on pregnancy and live birth rates regarding the very small
3106 number of patients and the extremely low re-utilization rates of cryopreserved oocytes and embryo in
3107 cancer patients (Boots et al., 2016).

3108 In a retrospective cohort study, conventional start stimulation in the early follicular phase (n=176) was
3109 compared with the luteal phase start (n=52) of ovarian stimulation for fertility preservation (Baig et al.,
3110 2023). No significant difference was found comparing early follicular with luteal phase start of
3111 stimulation for number of MII oocytes retrieved (9.0 (6.0-13.0) vs. 11.5 (7.0-16.0)).

3112 In a retrospective cohort study, conventional follicular ovarian stimulation (n=80) was compared to
3113 luteal phase ovarian stimulation (n=20) in women requiring gonadotoxic treatment (Jochum et al.,
3114 2019). Significantly more MII oocytes were retrieved after luteal phase ovarian stimulation compared
3115 to follicular phase (13.1±8.0 vs. 9.2±5.8).

3116 In a prospective cohort study, the effectiveness of controlled ovarian stimulation in the follicular (n=68)
3117 and luteal phase (n=72) of the menstrual cycle in cancer patients for the preservation of reproductive
3118 material before gonadotoxic therapy was evaluated (Nazarenko et al., 2021). No significant difference
3119 was reported in the total number of oocytes retrieved (715 vs. 766) or the proportion of MII oocytes
3120 (520 (72.8%) vs. 557 (72.6%)).



3121 Duostim or dual stimulation

3122 A systematic review and meta-analysis including cohort studies compared the outcomes of single or
3123 double ovarian stimulation cycles for fertility preservation (Chen et al., 2022). As expected, a double
3124 ovarian stimulation significantly increased the total number of retrieved oocytes available for fertility
3125 preservation in comparison to one cycle (MD 7.91, 95% CI 3.42 to 12.40).

3126 In a case series, the effectiveness of duostim was evaluated in 36 female oncology patients for fertility
3127 preservation (Puthur et al., 2023). A total of 324 oocytes were retrieved in the follicular phase
3128 stimulation, of which 184 were MII oocytes. A total of 337 oocytes were obtained after the luteal phase
3129 stimulation, of which 184 were MII oocytes. None of the thirty-six patients reported any symptoms of
3130 OHSS or delays to any previously planned cancer therapy.

3131 In a retrospective cohort study, the optimal timing of second ovarian stimulation using the dual
3132 stimulation method in 69 good ovarian responders with cancer undergoing oocyte retrieval for fertility
3133 preservation was evaluated (Takeuchi et al., 2023). In the first (follicular) stimulation, the numbers of
3134 retrieved and matured oocytes were 7.5 ± 5.6 and 5.3 ± 3.9 , respectively; in the second stimulation,
3135 these numbers were significantly higher (9.9 ± 6.6 and 9.4 ± 6.1 , respectively). Based on their data, they
3136 advise an 8-day waiting interval for a stable retrieval in the second cycle for cases where >5 oocytes
3137 were retrieved in the first oocyte retrieval because of ovarian enlargement resulting in a poor response
3138 to stimulation and delayed follicular development.

3139 Recommendation

For patients facing gonadotoxic treatment, ovarian stimulation for fertility preservation should be started irrespective of the menstrual cycle phase. [Updated]	Strong ⊕○○○
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3140

3141 Justification

3142 For fertility preservation for patients facing gonadotoxic treatment, ovarian stimulation should be
3143 started as soon as possible, also in view of double stimulation. Solid evidence for the optimal waiting
3144 time in between ovarian stimulation cycles is currently lacking.

3145 The systematic review and meta-analysis by Chen et al. shows that despite longer duration of
3146 stimulation and higher total gonadotropin consumption, the random-start stimulation finally led to
3147 similar number of oocytes retrieved, and metaphase II oocyte yield when compared with conventional
3148 start protocol. Therefore, random-start seems to be viable strategy in the setting of fertility
3149 preservation for cancer patients, although additional pregnancy rate data are needed. These findings,
3150 however, should be interpreted with caution given the limitations of the analysis. These include lack of
3151 randomized controlled trials, small sample sizes, retrospective nature of most studies, lack of detailed
3152 information on gonadotropin and trigger types and heterogeneity among the studies included.

3153 Current evidence indicates that oocyte competence is probably not impacted by its luteal phase origin
3154 compared to follicular phase. Absence of adverse effects on neonatal outcomes and long-term child
3155 health need to be evaluated on a larger scale, especially in cancer patients.

3156 The drug marketing approval for gonadotropin use in luteal phase needs to be considered.



3157 PITUITARY SUPPRESSION PROTOCOL

3158 PITUITARY SUPPRESSION

3159 Evidence

3160 Only one retrospective analysis, including 24 women, compared the long GnRH agonist and GnRH
3161 antagonist protocols in women with breast cancer who were treated with FSH plus letrozole (Ben-
3162 Haroush et al., 2011). The number oocyte recovered was higher with GnRH agonist protocol (24.8 ± 24.6
3163 vs. 12.0 ± 8.8), however this difference was not statistically significant. Furthermore, one patient had 82
3164 oocytes retrieved after long GnRH agonist protocol. When this patient is excluded, the mean of oocytes
3165 was 9.6 oocytes (range 0–30) (Ben-Haroush et al., 2011).

3166 Two systematic reviews including a total of 33 studies (Boots et al., 2016; Rodgers et al., 2017) and 14
3167 other investigations (Alvarez and Ramanathan, 2016, Cardozo et al., 2015, Chan et al., 2015, Das et al.,
3168 2011, Devesa et al., 2014, Druckenmiller et al., 2016, Garcia-Velasco et al., 2013, Johnson et al., 2013,
3169 Lawrenz et al., 2010, Lee et al., 2010, Muteshi et al., 2018, Pereira et al., 2016, Shapira et al., 2015)
3170 reported data of cancer patients having undergone ovarian stimulation for oocyte and/or embryo
3171 cryopreservation. More than 2200 cycles were described, most of them (>90%) with GnRH antagonist
3172 protocols. Among them, random-start ovarian stimulation or protocols using aromatase inhibitors or
3173 tamoxifen were considered. In addition, different methods of final oocyte maturation were used. The
3174 main outcome measure was usually the overall number of oocytes recovered and the number of
3175 mature oocytes obtained.

3176 In a retrospective cohort study, ovarian stimulation with progestins for pituitary suppression combined
3177 with hMG and double trigger (n=46) was compared to pituitary suppression with a GnRH antagonist
3178 protocol combined with rFSH and trigger with hCG or GnRH agonist alone (n=78) (Filippi et al., 2023).
3179 Significantly more oocytes were retrieved with the GnRH antagonist protocol compared to the
3180 progestins (16 (10-21) vs. 10 (5-17)), however, the number of MII oocytes frozen was similar in both
3181 groups (10 (6-18) vs. 9 (4-14)). The number of cancelled cycles was also similar in both groups (3 (7%)
3182 vs. 5 (6%)).

3183 In a retrospective cohort study, ovarian stimulation with progestins for pituitary suppression (n=20)
3184 was compared to GnRH antagonist (n=20) in patients with breast cancer for fertility preservation
3185 (Oliveira et al., 2021). No significant difference was noted comparing the GnRH antagonist protocol with
3186 progestins for the number of oocytes retrieved (4.5 (3-10.7) vs. 9 (4.1-12.8) or the proportion of MII
3187 oocytes (4 (2.1-9.8) vs. 7.5 (3.1-10)). There were 2 cases of OHSS in the GnRH antagonist group and 5
3188 in the progestins group.

3189 OVARIAN STIMULATION

3190 Evidence

3191 *Fertility preservation in breast cancer represents a complex issue since this disease is considered as*
3192 *oestrogen sensitive. Indeed, ovarian stimulation for the purpose of freezing oocytes or embryos is*
3193 *associated with supra-physiological serum oestradiol levels that could theoretically result in the*
3194 *proliferation of malignant cells.*



3195 *Therefore, innovative stimulation protocols have been developed in an effort to reduce potential harm*
3196 *associated with high oestradiol levels. Co-administration of either aromatase inhibitors or selective*
3197 *oestrogen receptor modulators during ovarian stimulation is used frequently.*

3198 A systematic review and meta-analysis²², including 16 cohort studies, compared the outcomes of
3199 coadministration of aromatase inhibitors or tamoxifen cycles during ovarian stimulation for fertility
3200 preservation (Chen et al., 2022). No significant differences in the numbers of retrieved oocytes were
3201 observed between those using and not using letrozole regardless of ovarian stimulation protocol (mean
3202 difference -0.55; 95% CI -2.01 to 0.91 and similar results were observed with the used of tamoxifen
3203 (mean difference 0.67; 95% CI -1.29 to 2.64). A significantly lower peak serum oestradiol concentration
3204 was observed in letrozole-based groups than in letrozole-free groups (mean difference -1.22; 95% CI
3205 -1.42 to -1.02).

3206 A systematic review and meta-analysis investigated the effect of tamoxifen supplementation compared
3207 to letrozole for patients with oestrogen-sensitive breast cancer undergoing ovarian stimulation for
3208 fertility preservation (Yoshida et al., 2023). No significant difference was reported for the number of
3209 oocytes retrieved (MD -0.47, 95% CI -3.84 to 2.90, 2 RCT) or MII oocytes (MD 0.22, 95% CI -2.20 to 2.64,
3210 2 RCT).

3211 In a retrospective cohort study, the outcomes of women with oestrogen-sensitive breast cancer
3212 undergoing ovarian stimulation with tamoxifen supplementation (n=154) were compared to women
3213 with non-oestrogen-sensitive breast cancer having ovarian stimulation without tamoxifen (n=60) (Sii et
3214 al., 2023). No significant difference was noted between ovarian stimulation with or without tamoxifen
3215 for the total number of oocytes retrieved (13.8 (12.1-15.4) vs. 12.0 (9.7-14.3)) or number of MII oocytes
3216 retrieved (10.5 (9.1-12.0) vs. (8.9 (7.3-10.5)).

3217 In a retrospective cohort study, the impact of letrozole use in oocyte cryopreservation (n=48, 55 cycles)
3218 among adolescent and young adult cancer patients for fertility preservation was investigated and
3219 compared to conventional gonadotropin stimulation (n=25, 26 cycles) (Suzuki et al., 2023). There was
3220 no significant difference between ovarian stimulation with or without letrozole for the total number of
3221 oocytes retrieved (10.4±6.4 vs. 9.3±5.7) or their maturation rate (69.6±25.8% vs. 68.6±25.8%).

3222 A retrospective cohort study included women undergoing ovarian stimulation for fertility preservation
3223 with the GnRH antagonist protocol, with (n=84) or without the use of supplemental letrozole (n=162)
3224 (Lalami et al., 2022). There was no significant difference in the number of oocytes retrieved (14.2±0.7
3225 vs. 14.0±0.8) nor number of embryos cryopreserved (7.0±4.3 vs. 4.2±2.9) with or without letrozole
3226 supplementation during ovarian stimulation.

3227 In a retrospective cohort study, the effects of letrozole (n=36) or tamoxifen (n=30) coadministration on
3228 the outcomes of ovarian stimulation for fertility preservation were assessed and compared to
3229 conventional gonadotropin stimulation (n=52) (Shulman et al., 2021). There was no significant
3230 difference in number of oocytes retrieved or maturation rate with letrozole or tamoxifen
3231 coadministration compared to conventional stimulation (12 (7.5-18.5) and 78.6% MII vs. 12 (8-20.3)

²² The meta-analysis by Rodgers et al., 2017 cited here in the 2018 version of the guideline was replaced by a more recent meta-analysis. The retrospective cohort study by Pereira et al., 2017 is included in the new meta-analysis and therefore no longer mentioned separately.



3232 and 79% MII vs. (10.5±6-18) and 81.5% MII). The number of cryopreserved embryo's was also similar
3233 between groups (7 (2-10) vs. 5 (3-12.5) vs. 5 (3-7.5)).

3234 In a retrospective cohort study, the effect of letrozole supplementation (n=94) during ovarian
3235 stimulation for fertility preservation was compared to conventional gonadotropin stimulation (n=83)
3236 (Sonigo et al., 2019). There was no significant difference noted for the number of oocytes retrieved
3237 with or without letrozole supplementation (12.2±8.3 vs. 13.1±10.0), however, the maturation rate was
3238 significantly lower with letrozole supplementation during ovarian stimulation (64.9±22.8% vs.
3239 77.4±19.3%).

3240 Recommendation

For ovarian stimulation in women seeking fertility preservation for medical reasons the GnRH antagonist protocol is probably recommended. [2019]	Conditional ⊕○○○
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In ovarian stimulation for fertility preservation in oestrogen sensitive diseases the concomitant use of anti-oestrogen therapy, such as letrozole or tamoxifen, can be considered. [2019]	GPP
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3242

3243 Justification

3244 There is low-quality evidence of the necessity of considering a specific GnRH analogue protocol. GnRH
3245 antagonist protocols are preferred since they shorten the duration of OS, offer the possibility of
3246 triggering final oocyte maturation with GnRH agonist in case of high ovarian response, and reduce the
3247 risk of OHSS. Moreover, especially in cancer patients, who are at higher risk of thrombosis due to their
3248 oncologic status, GnRH antagonist protocols seem to be preferred since they enable GnRH agonist
3249 trigger, therefore reducing the risk of OHSS. Melo et al. reported that 3.1% of study participants had a
3250 thromboembolic event at a median of 0.25 years from oocyte aspiration for fertility preservation and
3251 0.33 year from their cancer diagnosis (Melo et al., 2022).

3252 RCTs aiming to compare GnRH agonist and GnRH antagonist protocols for fertility preservation may
3253 be interesting. However, considering such studies may be difficult since GnRH agonist trigger
3254 represents an important advantage in this field.

3255 Data on live births are dramatically lacking, in particular in cancer patients having vitrified oocytes.

3256 The quality of evidence is still low given the number and quality of studies available. The existing
3257 literature concerning ovarian stimulation for fertility preservation in women with oestrogen sensitive
3258 cancer is limited by its observational nature, small patient numbers and relatively short duration of
3259 follow-up. Definitive statements regarding the safety of OS in women with a recent diagnosis of breast
3260 cancer would require long-term and large-scale studies, and these do not yet exist. A recent
3261 retrospective cohort study reported that the 5-year invasive disease-free survival was not statistically
3262 different between the fertility preservation recipients and a subgroup of patients cotreated with



3263 tamoxifen during stimulation because of oestrogen-receptor positive disease (HR 1.66, 95% CI 0.67-
3264 3.49) (Shapira et al., 2025).

3265 Undertaking RCTs in this patient population represents a major limitation. It is not known whether the
3266 transient period of raised oestrogen concentrations during ovarian stimulation is harmful to women
3267 with breast cancer. A study aiming to compare the short- and long-term effects of ovarian stimulation
3268 with or without letrozole co-administration is ongoing. However, the use of letrozole is off-label for OS
3269 and safety concerns have been raised regarding possible teratogenicity associated with letrozole.

3270 Aromatase inhibitors protocols enable GnRH agonist trigger (Oktay et al., 2010, Reddy et al., 2014).

3271 FINAL OOCYTE MATURATION PROTOCOL

3272 Evidence

3273 In a retrospective cohort study, patients undergoing ovarian stimulation for fertility preservation
3274 received depot GnRH agonist (n=22), short-acting GnRH agonist (n=26) or hCG (10,000 IU; n=34) for
3275 final oocyte maturation (Massarotti et al., 2023). There was no significant difference between long-
3276 acting, short-acting or hCG triggering for the number of oocytes retrieved or MII rate (13.9±5.2 (80%
3277 mature) vs. 18±10.3 (80% mature) vs. 11.5±6.7 (74% mature)). No cases of OHSS were reported after
3278 the long-acting GnRH agonist trigger, and 1 case after the short-acting GnRH agonist trigger and the
3279 hCG trigger.

3280 In a retrospective cohort study, 293 patients (373 cycles) underwent ovarian stimulation for fertility
3281 preservation with the GnRH antagonist protocol. Final oocyte maturation was triggered with double
3282 trigger (n=148) in the study group compared to rhCG triggering in the control group (n=225) (Hong et
3283 al., 2022). No significant difference was found when comparing rhCG to double triggering for the
3284 number of oocytes retrieved (7.9±5.7 vs. 8.8±7.2), the proportion of MII oocytes (4.8±3.8 vs. 5.7±4.9)
3285 or the number of OHSS cases (5/225 (2.2%) vs. 7/148 (4.7%)).

3286 Recommendation

For final oocyte maturation, hCG is preferred, unless the patient is at risk of early OHSS, in which case GnRH agonist triggering is advised. [2025]

GPP

3287

3288 Justification

3289 Final oocyte maturation is a key step for fertility preservation. hCG has been the conventional strategy
3290 to induce final oocyte trigger. However, GnRH agonist trigger in antagonist protocols represents a safe
3291 option to limit the risk of ovarian hyperstimulation syndrome. Current evidence regarding the best
3292 trigger option is of low-quality, only based on retrospective studies. Therefore, hCG still appear to be
3293 the preferred strategy for inducing final oocyte maturation in case of normal ovarian response to
3294 stimulation.

3295



PICO QUESTION: WHAT IS THE PREFERRED STIMULATION PROTOCOL FOR ELECTIVE OOCYTE CRYOPRESERVATION?

3296 INITIATION OF STIMULATION

3297 Evidence

3298 In a prospective cohort study, patients presenting for elective oocyte preservation were offered the
3299 choice for either random-start (n=443) or conventional day 2/3 start (n=859) stimulation (Pereira et al.,
3300 2017). No significant difference was observed for number of MII oocytes retrieved with either random-
3301 start (early follicular, late follicular or luteal start) or conventional day 2/3 start stimulation (10.8±2.7
3302 vs. 11.1±3.0 vs. 10.9±3.2 vs. 13.1±2.3).

3303 Recommendation

Ovarian stimulation for elective oocyte preservation can be started irrespective of the menstrual cycle phase. [2025]

Conditional ⊕○○○

3304

3305 Justification

3306 Since in elective oocyte freezing cycles all oocytes will be cryopreserved, ovarian stimulation can be
3307 started irrespective of the menstrual cycle phase.

3308 PITUITARY SUPPRESSION PROTOCOL

3309 PITUITARY SUPPRESSION

3310 Evidence

3311 In a retrospective cohort study, including women of advanced maternal age undergoing elective oocyte
3312 cryopreservation, the use of a progestin protocol (n=89) was compared to a GnRH antagonist protocol
3313 (n=178) (Vaiarelli et al., 2024). No significant difference in the number of MII oocytes retrieved was
3314 reported between the progestin and GnRH antagonist protocol (6.8±5.6 vs. 6.2±4.1). A total of 61 and
3315 107 vitrified-warmed euploid SETs were performed. No significant difference was observed for
3316 cumulative LBR (24.7% (21/85) vs. 21.9% (39/178)) or live birth rate/transfer ((37.7% (23/61) vs. (39.3%
3317 (42/107)) between progestin and GnRH antagonist protocol.

3318 OVARIAN STIMULATION

3319 Evidence

3320 In a retrospective cohort study, 217 patients presenting for elective oocyte cryopreservation
3321 underwent a first IVF cycle with 300 IU rFSH and a second IVF cycle with a an adjusted rFSH dosage
3322 (increased, decreased or no change) (Orvieto et al., 2022). Comparing the first to the second ovarian
3323 stimulation cycle, significantly more MII oocytes were retrieved in the second cycle (8.96±5.19 vs.
3324 8.04±4.7). In the second ovarian stimulation cycle, 23 (10.6%) women received a lower daily
3325 gonadotropin dose, 60 (27.6%) received the same dose and 134 (61.7%) an increased daily dose. Those



3326 who achieved a lower oocyte yield in the second cycle received significantly higher daily dose of
3327 gonadotropins (415±88 IU vs. 369±106 IU).

3328 Recommendation

GnRH antagonist or progestin protocol are probably recommended over GnRH agonist protocols for pituitary suppression in elective oocyte cryopreservation.

Conditional ⊕○○○

3329

3330 Justification

3331 Only low-quality evidence from one retrospective cohort study was available in the elective oocyte
3332 cryopreservation population. However, data from the general infertility population showed that GnRH
3333 antagonist and progestin protocol are preferred over GnRH agonist protocol for elective
3334 cryopreservation. The reader is referred to the chapter 6 for information on the choice of
3335 gonadotropins for ovarian stimulation for elective oocyte cryopreservation.

3336 FINAL OOCYTE MATURATION PROTOCOL

3337 Evidence

3338 In a retrospective cohort study, reproductive outcomes were compared after GnRH agonist (n=40) or
3339 hCG (n=29) for the final oocyte maturation trigger (Herzberger et al., 2021). The decision was made
3340 according to laboratory and sonographic results on the day of triggering, with the risk of OHSS
3341 considered. Patients included in the GnRH agonist trigger group were significantly younger compared
3342 to the hCG group. Significantly more oocytes were retrieved after GnRH agonist trigger compared to
3343 hCG (16.5 (8.0-25.0) vs. 6.0 (2.5-11.0)). However, the maturation rate was comparable (0.8 (0.7-0.9 vs.
3344 0.8 (0.7-1.0)).

3345 In a retrospective cohort study, it was examined whether GnRH agonist trigger (n=959) for final oocyte
3346 maturation can be reliably used and was compared to hCG (n=671) and dual trigger (n=50) (Maslow et
3347 al., 2020). Cycles using hCG trigger were characterised by significantly higher age and lower AMH and
3348 LH. Significantly less MII oocytes were retrieved with hCG trigger compared to GnRH agonist and dual
3349 trigger (8.4±5.9 vs. 13.3±9.1 vs. 13.0±7.8). There were no cases of severe OHSS requiring
3350 hospitalisation, medical or surgical intervention.

3351 In a retrospective cohort study, dual trigger (n=40) was compared to hCG trigger (n=36) for final oocyte
3352 maturation in patients with diminished ovarian reserve undergoing elective cryopreservation (Kim et
3353 al., 2020). Significantly more MII oocytes were retrieved with dual trigger compared to hCG trigger
3354 (3.7±2.7 vs. 2.3±1.7). furthermore, the oocyte maturation rate was significantly higher after dual trigger
3355 compared to hCG trigger (68.5% (146/213) vs. 45.6% (82/180)).



3356 Recommendation

For final oocyte maturation in elective oocyte cryopreservation, hCG is preferred, unless the patient is at risk of early OHSS, in which case GnRH agonist trigger is advised. [2025]

GPP

3357

3358 Justification

3359 hCG and GnRH agonist for final oocyte maturation result in similar numbers of mature oocytes. If a
3360 patient is at risk of early OHSS, a GnRH agonist trigger is advised.

3361

3362

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3506

Draft for review



3507 11. Ovarian stimulation for oocyte donation

PICO QUESTION: WHAT IS THE PREFERRED STIMULATION PROTOCOL FOR OOCYTE DONATION?

3508 Considering the motivation for ovarian stimulation for oocyte donation, critical and important
3509 outcomes in this chapter are different from the rest of this guideline. Critical outcomes for ovarian
3510 stimulation for oocyte donation in this guideline are the number of oocytes/embryo's and preventing
3511 OHSS and other complications.

3512 INITIATION OF STIMULATION

3513 Ovarian stimulation in the luteal phase, several case reports have described spontaneous pregnancies
3514 that were undetectable at the time of starting ovarian stimulation. This carries the risk of early
3515 pregnancy exposure to medications used during ovarian stimulation, as well as the risk of ovarian
3516 hyperstimulation syndrome due to endogenous hCG production. It is therefore essential to inform
3517 oocyte donors about the risk of natural conception prior to commencing random-start ovarian
3518 stimulation (Lawrenz et al., 2024, Semrl et al., 2024).

3519 Evidence

3520 In an RCT, 67 oocyte donors were randomised to receive ovarian stimulation starting either in the early
3521 (n=35) or late (n=32) follicular phase in a GnRH antagonist protocol (De Rijdt et al., 2024). There was no
3522 significant difference in the number of MII oocytes retrieved when comparing early to late follicular
3523 stimulation (14.1±8.1 vs. 12.7±8.5). No cases of OHSS were reported in either group.

3524 In a prospective cohort study, oocyte donors underwent two consecutive ovarian stimulation protocols
3525 with at least one month in between both cycles. The cycles were identical, aside from the start of
3526 stimulation, follicular phase in the first cycle and luteal phase in the second cycle (Martinez et al., 2022).
3527 There was no significant difference for number of MII oocytes with follicular or luteal start stimulation
3528 (20.27±9.60 vs. 20.73±8.65). The mean number of euploid embryos was equivalent between the
3529 follicular and the luteal start groups (1.59±1.30 vs. 1.61±1.17). At the time of publication, 42 recipients
3530 have undergone at least one FET, with a total of 68 FET being performed. Clinical pregnancy rate was
3531 42.9% from the follicular phase stimulation and 59.0% from the luteal phase stimulation.

3532 In a retrospective cohort study, live birth rates were investigated in recipients matched with donors
3533 using random-start or conventional follicular start ovarian stimulation in (Guerrero et al., 2024). There
3534 were no significant differences in the total number of oocytes retrieved (17.2±8.5 vs. 17.6±8.8) or MII
3535 oocytes retrieved (13.5±7.0 vs. 13.8±7.1) between random and conventional start ovarian stimulation.
3536 There was no significant difference in live birth rate in recipients with oocytes retrieved after random
3537 start or conventional start ovarian stimulation (46.6% (201/537) vs. 47.7% (62/173); OR 0.88, 95% CI
3538 0.48-1.58).

3539 Recommendation

Conventional follicular start or random-start ovarian stimulation are equally recommended for oocyte donation cycles. [2025]

Strong ⊕○○○



3540

If random-start ovarian stimulation is used, oocyte donors need to adopt contraceptive measures to prevent the possibility of a natural pregnancy. [2025]

GPP

3541

3542 Justification

3543 Current evidence in oocyte donors reports no difference in efficacy in terms of the number of oocytes
3544 or the number of mature oocytes retrieved, and no difference in the live birth rate in oocyte recipients
3545 when stimulation is initiated in the early follicular or luteal phase. This supports the option of random-
3546 start ovarian stimulation protocols for oocyte donors. Some studies have reported unexpected
3547 spontaneous pregnancies during ovarian stimulation with random-start protocols, the possibility of
3548 which must be carefully excluded before commencing ovarian stimulation.

3549 PITUITARY SUPPRESSION PROTOCOL

3550 CONTRACEPTIVE PRE-TREATMENT

3551 Evidence

3552 *No randomised controlled studies were identified for this section.*

3553 In a retrospective study, including 491 consecutive cycles of vitrified oocyte donation undergoing
3554 ovarian stimulation using GnRH antagonist co-treatment and GnRH agonist trigger, the use of
3555 contraceptive pre-treatment with an IUD (n=103 cycles) was compared to no pre-treatment (n=388
3556 cycles) (Galvão et al., 2019). Comparing contraceptive pre-treatment to no pre-treatment, no
3557 significant differences were found for the number of MII oocytes retrieved (14.5 ± 6.9 vs. 14.2 ± 7.3) and
3558 number of top quality embryos (2.3 ± 1.2 vs. 2.3 ± 1.3). Cumulative live birth rate per embryo transfer in
3559 oocyte recipients was also similar between groups (49% (47/96) vs. 45.3% (162/358)).

3560 In a prospective cohort study oocyte donors were assigned to receive ovarian stimulation after 5 days
3561 (n=42), or after 7 days of pill discontinuation (n=50) in a GnRH antagonist protocol (Pérez-Calvo et al.,
3562 2017). Extended pill-free interval of 7 days did not significantly influence the number of MII oocytes
3563 retrieved (12.4 ± 7.4 vs. 10.6 ± 4.9).

3564 Recommendation

The use of any type of contraception (hormonal, non-hormonal, oral, vaginal or intrauterine) before or during ovarian stimulation is not a contraindication in oocyte donors. [2025]

GPP

3565



3566 Justification

3567 Current evidence in oocyte donors reports no difference in efficacy in terms of the number of oocytes
3568 or the number of mature oocytes retrieved, when comparing pre-treatment with OCP or IUD to no pre-
3569 treatment. Furthermore, no differences were observed in the cumulative live birth rates in oocyte
3570 recipients. An extended pill free interval of 5 or 7 days is usually recommended prior to initiation of
3571 stimulation.

3572 PITUITARY SUPPRESSION

3573 Evidence

3574 GnRH analogues

3575 In a systematic review and meta-analysis, clinical outcomes were compared between the use of GnRH
3576 agonists and antagonists in oocyte-donation cycles (Bodri et al., 2011). Comparing GnRH agonist and
3577 GnRH antagonist protocols, no significant difference was found for the number of retrieved oocytes
3578 (WMD 0.60, 95% CI 2.26 to 1.07, 7 RCT, 932 donors). Although OHSS incidence was not different
3579 between treatment groups (RR 0.61, 95% CI 0.18-2.15, 4 RCT), results should be interpreted with
3580 caution, since this might be related to the small sample size, unable to detect any significant differences.

3581 A prospective cohort study investigated the use of a long-acting GnRH antagonist in comparison to the
3582 use of a conventional GnRH antagonist in historic controls (Boniface et al., 2023). The average number
3583 of total oocytes retrieved was similar between the control and study group (30.55 vs. 30.31). The
3584 average number of mature oocytes was similar as well between the control and study group (25.42
3585 vs.24.73).

3586 In a retrospective cohort study, the clinical outcomes were compared between clomiphene-citrate
3587 (n=133) and GnRH antagonist-based protocols (n=100) in donor cycles (Singh et al., 2016). The number
3588 of MII oocytes retrieved (13.04±5.73 vs. 12.96±6.08) and the number of grade I embryos (8.32±5.09 vs.
3589 7.95±4.77) was similar in the clomiphene citrate and the GnRH antagonist groups. The number of OHSS
3590 cases was also similar between groups (10 vs. 9). No significant difference was reported in live birth
3591 rate per started cycle: 47.8% in the clomiphene group and 39.55% in the GnRH antagonist group.

3592 Progestins

3593 In a systematic review and meta-analysis, clinical outcomes were compared between the use of
3594 progestins and GnRH antagonist protocols for pituitary suppression in oocyte donors (Martinez et al.,
3595 2021). Meta-analysis of the 2 RCTs comparing PPOS with GnRH antagonist protocols for the treatment
3596 in 490 oocyte donors showed no differences in mean number of retrieved oocytes (MD 0.33, 95% CI -
3597 1.30 to 1.96) and in clinical pregnancy rate among 625 recipients (OR 0.83, 95% CI 0.33-2.06).

3598 In an RCT, reproductive outcomes were compared in oocyte donors undergoing pituitary suppression
3599 with progestins (n=161) versus conventional treatment with a GnRH antagonist (n=157) (Giles et al.,
3600 2021). No significant difference was found between the study and control group for the number of MII
3601 oocytes retrieved (16.7±9 vs. 16.9±7.7). Cumulative live birth rate (70.6% (130/175) vs. 68.7%
3602 (121/171)) and live birth rate (52.7% (90/175) vs. 47.1% (83/171)) were not significantly different in
3603 recipients of the oocytes after the use of the progestin or GnRH antagonist protocol.



3604 Recommendation

For pituitary suppression in oocyte donors the GnRH antagonist and progestin protocol are probably equally recommended. [2025]

Strong ⊕⊕○○

3605

A GnRH agonist protocol is not recommended in oocyte donors. [2025]

GPP

3606

3607 Justification

3608 Although GnRH agonist and GnRH antagonist protocols in oocyte donors results in comparable numbers
3609 of oocytes and mature oocytes, and result in similar live birth, the use of GnRH agonists is associated
3610 with higher risk of OHSS. There is low-quality evidence that ovarian stimulation in oocyte donors using
3611 the GnRH antagonist protocol or progestin-primed ovarian stimulation yield similar numbers of oocytes
3612 and mature oocytes, and result in similar live birth and cumulative live birth rates in oocyte recipients.
3613 Both offer the possibility of triggering final oocyte maturation with a GnRH agonist, minimising the risk
3614 of OHSS and optimising safety for oocyte donors.

3615 The use of a long-acting GnRH antagonist has been studied only in a cohort study design, without
3616 reporting the effects on the outcome live birth. The same is reported for the use of Clomiphene citrate
3617 and recommendations are therefore not given. Given the high risk of OHSS long-acting agonist should
3618 not be recommended in oocyte donors.

3619 OVARIAN STIMULATION

3620 Evidence

3621 Type of stimulation drug

3622 In an RCT, healthy oocyte donors were randomly assigned to start ovarian stimulation with a single dose
3623 of long-acting rFSH 7 days after OCP discontinuation (n=90), compared to a conventional protocol
3624 where ovarian stimulation is started 5 days after OCP discontinuation with a single dose of long-acting
3625 rFSH followed by additional 225 IU rFSH starting on day 8 (n=90) in the GnRH antagonist protocol
3626 (Alvarado Franco et al., 2023). The number of MII oocytes retrieved was significantly lower in the study
3627 group compared to the control group (10 (6-14) vs. 12 (9-17.25)).

3628 In an RCT, three types of gonadotropins were compared in an oocyte donor programme: long-acting
3629 rFSH (n=68), rFSH (150 IU, n=69) and hMG (225 IU, n=71) (Cruz et al., 2017). Comparing long-acting
3630 rFSH to rFSH and hMG, no significant difference was observed for the number of MII oocytes retrieved
3631 (12.2±1.1 vs. 12.1±1.4 vs. 12.3±2.1) and cycle cancellation for poor response (2/68 vs. 2/69 vs. 5/71).
3632 Clinical pregnancy rates in oocyte recipients were similar: 60.5% for the long-acting rFSH group; 59.5%
3633 for the rFSH group; and 63.2% for the hMG group.

3634 In an RCT, participants were randomly assigned to one of three gonadotropin regimes: rFSH only
3635 (n=346), hMG only (n=333) or rFSH and hMG combination (n=349) in a GnRH agonist protocol (Melo et



3636 al., 2010). When comparing rFSH only to hMG only and the rFSH and hMG combination, there was no
3637 significant difference reported for risk of mild and moderate OHSS (7.04% (20/284) vs. 6.78% (19/280)
3638 vs. 5.52% (16/290)), number of top quality embryos (3.4 ± 0.4 vs. 3.5 ± 0.5 vs. 3.6 ± 0.4) or cycle
3639 cancellation rate (18% (62/346) vs. 16% (53/333) vs. 17% (59/349)). No cases of severe OHSS were
3640 observed. No significant differences were observed in clinical pregnancy rates in oocyte recipients after
3641 rFSH (56.7% (199/351)), hMG (57% (207/363)) or rFSH and hMG combination (59.2% (216/365)) for
3642 ovarian stimulation.

3643 In an RCT, oocyte donors were randomly assigned to received either rFSH alone (n=127) or rFSH with
3644 LH supplementation by hMG on stimulation days 5-7 (n=126) in a long GnRH agonist protocol. The
3645 groups were further stratified based on their baseline LH levels: baseline LH < 1 IU/L (groups 1 and 2,
3646 without and with supplemental LH activity, respectively) and baseline LH > 1 IU/L (groups 3 and 4,
3647 without and with supplemental LH activity respectively). On stimulation day 5, the groups were further
3648 stratified based on their oestradiol levels: <100 pg/ml (a) and ≥ 100 pg/ml (b) (Tesarik and Mendoza,
3649 2002). The number of MII oocytes per donor was significantly higher in all groups co-stimulated with
3650 LH when compared with corresponding groups stimulated with FSH alone. In women with baseline LH
3651 < 1 IU/L, the number of good-quality cleavage-stage embryos was significantly higher with LH activity
3652 supplementation. No differences in pregnancy rates were detected between any comparable groups
3653 with and without the inclusion of exogenous LH to the stimulation protocol.

3654 In an RCT, participants were randomly assigned to receive either hp-FSH (n=20) or hMG (n=21) for
3655 ovarian stimulation in an oocyte donation programme (Söderström-Anttila et al., 1996). One donor in
3656 each group developed moderate OHSS after oocyte retrieval. Two cycles were cancelled, one in each
3657 group. 53% of the donors in the hp-FSH group (10/19) and 42% in the hMG group (8/19) had com-
3658 plaints about side-effects and discomfort (headache, tiredness, abdominal swelling and pain, nausea
3659 and irritability). One donor in the hp-FSH group and two donors in the hMG group experienced a mild
3660 fever reaction.

3661 In an RCT, ovarian stimulation with rFSH alone (225IU, n=20) was compared to rFSH (225 IU) combined
3662 with LH (75 IU) from day 6 of stimulation (n=22) in a short GnRH agonist protocol for oocyte donors
3663 (Acevedo et al., 2004). The number of MII oocytes retrieved (80 vs 71) and the number of grade I
3664 embryos (17 vs. 3) was significantly higher with LH supplementation compared to no supplementation.
3665 None of the donors developed severe OHSS. No significant difference was reported in clinical pregnancy
3666 rate (51% vs. 30%) in oocyte recipients.

3667 Dosing and formulation

3668 In a prospective cohort study, clinical outcomes were compared between two ovarian stimulation
3669 cycles in the same high responder oocyte donors: a dose of 225 IU (n=32) stimulation protocol, followed
3670 by a dose of 150 IU (n=32) stimulation protocol (Rubio et al., 2010). The number of MII oocytes retrieved
3671 was significantly lower with the lower dose compared to the higher dose (262 vs. 428). Only 22 donors
3672 completed both cycles, for 10 donors, the reduced-dose cycle was cancelled for low response. The
3673 number of live births was similar after 150 IU or 225 IU for ovarian stimulation (13 vs. 11).

3674 In a retrospective cohort study, clinical outcomes were compared between rFSH filled by mass (n=12
3675 cycles) compared to rFSH filled by conventional bioassay (n=11 cycles) in the same oocyte donors
3676 (Martinez et al., 2007). The number of oocytes retrieved was significantly higher with rFSH filled by mass



3677 compared to rFSH filled by bioassay (23.8±8.7 vs. 17.1±8.5). The number of day-5 embryos was similar
3678 in both groups (5.4±3.1 vs. 5.1±3.0). There were no cases of OHSS reported in either group.

3679 In a retrospective cohort study, clinical outcomes and patient satisfaction were compared between
3680 reconstituted rFSH (n=19 cycles) or a cartridge pen system (n=79 cycles) in oocyte donors (Christianson
3681 et al., 2007). The number of MII oocytes retrieved was not significantly different with the reconstituted
3682 rFSH or the pen system (23.7±3 vs. 23.1±1.3). Donors scored significantly higher medication tolerance
3683 scores using the cartridge pen device (3.9 ± 0.4 vs. 3.1 ± 0.6, p < 0.05). Five donors who had used both
3684 formulations also noted greater satisfaction using the cartridge pen device rFSH compared to
3685 reconstituted rFSH (3.7 ± 0.2 vs. 3.1 ± 0.4, p < 0.01, respectively). No significant difference was reported
3686 in clinical pregnancy rate per embryo transfer in oocyte recipients (45% (8/18) vs. 61% (55/90).

3687 Recommendation

The use of recombinant FSH, purified FSH, long-acting FSH or hMG is probably equally recommended in oocyte donors undergoing ovarian stimulation. [2025]	Conditional ⊕○○○
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3688

Gonadotropin dose should be individualised based on ovarian reserve with the goal to maintain donors' safety and also obtain an optimal number of oocytes. [2025]	GPP
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3689

3690 Justification

3691 Several randomised, controlled trials have shown no difference in the number of oocytes or number of
3692 embryos obtained using different FSH preparations in oocyte donors. One RCT reported a high cycle
3693 cancellation rate due to low response in donors receiving 150 IU FSH/day compared to 225 IU FSH/day.
3694 No studies have reported on live birth outcomes, and further clinical research is required.

3695 The issue of dosing in oocyte donation cannot be answered with the current evidence.

3696 FINAL OOCYTE MATURATION PROTOCOL

3697 Evidence

3698 A Cochrane systematic review and meta-analysis, including three RCTs and 372 donors, compared hCG
3699 trigger with GnRH agonist for final oocyte maturation in oocyte donors (Youssef et al., 2014). The
3700 incidence of OHSS was lower with GnRH agonist compared to hCG for final oocyte maturation (OR 0.05,
3701 95% CI 0.01-0.28, 3 RCT, 372 donors) and mild-moderate OHSS was observed only after hCG triggering.
3702 No significant difference was found for the number of retrieved oocytes between GnRH agonist and
3703 hCG for final oocyte maturation. Live birth rate was similar between hCG and GnRH agonist trigger (OR
3704 0.92, 95% CI 0.53-1.61, 1 RCT, 212 women).



3705 A meta-analysis found no differences in the CPR among the corresponding recipients after ovulation
3706 triggering with GnRH agonist or hCG (OR 0.86, 95% CI 0.58-1.26, 4 RCT, 460 donors) (Martinez et al.,
3707 2021).

3708 An RCT compared two different recombinant hCG (r-hCG) doses, 250 µg (n=57) and 500 µg (n=55), for
3709 final oocyte maturation in a GnRH antagonist protocol in oocyte donors (Clua et al., 2012). Comparing
3710 the lower to the higher rhCG dose, no significant difference was noted in the number of MII oocytes
3711 retrieved (10.1±3.2 vs. 9.2±3.4). Mild OHSS was observed in 17 donors (29%) of the 250 rhCG dose
3712 group and in 23 (39%) of the 500 lg r-hCG dose group. Clinical pregnancy rate was similar in oocyte
3713 recipients (56.1% (32/57) vs. 58.2% (32/55)).

3714 In a retrospective cohort study, clinical outcomes were compared after hCG (42 cycles), GnRH agonist
3715 (232 cycles) and dual (59 cycles) trigger for final oocyte maturation in oocyte donor cycles (Jones et al.,
3716 2021). The number of MII oocytes retrieved was significantly lower after hCG trigger compared to GnRH
3717 agonist and dual trigger (7.1±3.4 vs. 11.2±5.5 vs. 11±6.0). Significantly more cases of OHSS were
3718 reported after dual trigger compared to hCG and GnRH agonist trigger (8.5% (5/59) vs. 0% vs. 0.4%
3719 (1/232)).

3720 Recommendation

The routine use of a GnRH agonist trigger is recommended in oocyte donors using the GnRH antagonist or progestin protocols for pituitary suppression. [2025]

Strong ⊕⊕○○

3721

The use of a hCG trigger is not routinely recommended in oocyte donation cycles. [2025]

Strong ⊕⊕○○

3722

3723 Justification

3724 Two systematic reviews and meta-analysis reported similar oocyte and mature oocyte yield between
3725 GnRH agonist triggering and hCG triggering, while no differences in the CPR among the corresponding
3726 recipients were observed. Owing to the risk of ovarian hyperstimulation syndrome when using an hCG
3727 trigger compared to a GnRH agonist trigger hCG trigger should not be recommended in oocyte donation
3728 cycles. The GDG has not considered the rare occasions where the donor has gonadotropin insufficiency.
3729 In most oocyte donors, GnRH agonist should be used for safety (OHSS). However, there may be cases
3730 where hCG is needed.

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3732

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Draft for review



PART E: Monitoring

3808

12. Hormonal assessment during ovarian stimulation

3809

PICO QUESTION: WHEN TO START MONITORING OF FOLLICULAR DEVELOPMENT?

3810 Monitoring the response of the ovaries to the gonadotropin stimulation serves the purpose of knowing
3811 the number and size of follicles that is growing and are expected to deliver a useful oocyte after the
3812 follicle aspiration. In addition, the size and number of follicles with a certain diameter can be assessed
3813 in order to time the moment of the ovulation trigger. Although usual practice consists of a baseline
3814 ultrasound scan, with follow up ultrasound monitoring from day 8 of the stimulation onwards, quite
3815 some practice variation exists. The same is true for hormonal assessments that mainly focus on the
3816 degree of pituitary suppression, the development of early progesterone rises and the measurement of
3817 oestradiol as an indicator of follicle numbers. For none of these markers scientific studies exist to
3818 demonstrate a benefit of certain starting moments.

PICO QUESTION: IS THE ADDITION OF HORMONAL ASSESSMENT (OESTRADIOL/PROGESTERONE/LH TO ULTRASOUND MONITORING IMPROVING EFFICACY AND SAFETY?)

3819 A survey was conducted to understand the global practice of routine hormone monitoring during
3820 ovarian stimulation for IVF/ICSI (Sachs-Guedj et al., 2023). Most respondents (98.9%) used ultrasound
3821 for monitoring ovarian stimulation cycles. Hormonal monitoring was widely accepted and used by 420
3822 (79.5%) of participants during any of the cycle monitoring visits. Oestradiol was the most frequently
3823 monitored hormone during the first and second/third clinic visit after the first gonadotropin injection.
3824 Hormone monitoring was most commonly performed on the day of, or day prior to final oocyte
3825 maturation, with 71% of respondents measuring oestradiol. The number of respondents who measured
3826 P4 (67.7%) was twice that during the second/third visit. There was also an increase in the proportion of
3827 respondents measuring LH, from 27.3% in the second/third visit, to 31.5% in the visit on the day of, or
3828 day prior to ovulation triggering. Oestradiol monitoring was used by 74% of respondents for the
3829 prediction of ovarian hyperstimulation syndrome (OHSS). Among the respondents, 23.5% measured
3830 progesterone in all patients or nearly all patients, and 21.1% measured it in some patients. Most
3831 respondents (60.7%) believed that hormones play an important role in monitoring ovarian response
3832 during OS, and 56% considered that HA is important to guide decision-making for the prevention of
3833 OHSS.

3834 ULTRASOUND AND OESTRADIOL MEASUREMENTS

3835 Evidence

3836 A Cochrane meta-analysis on monitoring of ovarian stimulation in IVF/ICSI with ultrasound alone
3837 compared to ultrasound plus serum oestradiol concentration combined 6 RCTs including 781 women
3838 (Kwan et al., 2014). Monitoring of the stimulation phase by using serum oestradiol measurements and
3839 ultrasound did not appear to decrease the probability of OHSS (6 RCT, OR 1.03, 95% CI 0.48-2.20, 781
3840 women), nor increase the probability of clinical pregnancy (4 RCT, OR 1.10, 95% CI 0.79-1.54, 617



3841 women), or the number of oocytes retrieved (5 RCT, WMD 0.32, 95% CI -0.60 to 1.24, 596 women)
3842 (Kwan et al., 2014).

3843 Recommendation

The addition of oestradiol measurements to ultrasound monitoring is probably not recommended. [2019]

Conditional ⊕⊕○○

3844 Justification

3845 On the basis of the currently published evidence, monitoring of the stimulation phase by serum
3846 oestradiol measurements and ultrasound is not superior to monitoring by ultrasound alone in terms
3847 of efficacy and safety. The addition of oestradiol in the monitoring does not appear to increase the
3848 probability of pregnancy, the number of oocytes retrieved, or to decrease the probability of OHSS.

3849 From the six studies included in the meta-analysis, a GnRH agonist protocol was used exclusively in
3850 four of them, while in the remaining two both GnRH agonists and antagonists were used (Kwan et al.,
3851 2014). Thus, it is not known whether the recommendation is valid in patients treated exclusively with
3852 GnRH antagonists.

3853 The Cochrane meta-analysis was updated in 2021 (Kwan et al., 2021), however, no new studies were
3854 identified. The evidence based on the six trials identified in 2014 remained unchanged.

3855 **ULTRASOUND AND PROGESTERONE MEASUREMENTS OR ULTRASOUND AND LH MEASUREMENTS.**

3856 Currently no published evidence exists to allow for a recommendation to be formulated answering
3857 these questions.

3858 **ULTRASOUND AND COMBINATION OF HORMONAL MEASUREMENTS**

3859 Evidence

3860 One RCT (114 women) reported no difference in OHSS (5.3% (3/57) vs. 7.0% (4/57)), pregnancy rate
3861 (22.2% vs. 25%), or number of oocytes retrieved (11.7±8.4 vs. 13.4±7.5) when monitoring was
3862 performed with ultrasound with or without hormonal measurements (Golan et al., 1994). Similarly, a
3863 more recent RCT (63 women) reported no difference in clinical pregnancy rate (40.0% (12/30)) vs.
3864 57.5% (19/33) or number of oocytes retrieved (10.0±5.5 vs. 11.7±8.0) with ultrasound and hormone
3865 panel monitoring compared with ultrasound only (Wiser et al., 2012). Furthermore, no cases of OHSS
3866 were reported in either the study or control group (Wiser et al., 2012).

3867 Recommendation

The addition of a hormonal panel consisting of a combination of oestradiol, progesterone and LH measurements to ultrasound monitoring is probably not recommended. [2019]

Conditional ⊕○○○

3868



3869 **Justification**

3870 According to one RCT, monitoring of the stimulation phase by using hormonal panel assessments
3871 (oestradiol, LH, progesterone) and ultrasound is not beneficial in terms of efficacy and safety over
3872 monitoring by ultrasound alone in terms of efficacy and safety. The addition of hormonal assessments
3873 in the monitoring does not appear to increase the probability of pregnancy, the number of COCs
3874 retrieved, or to decrease the probability of OHSS or cycle cancellation for high response.

3875 In the two studies, pituitary suppression was performed with GnRH agonists (Golan et al., 1994) or
3876 either GnRH agonists/antagonists (Wiser et al., 2012). Thus, it is not known whether the
3877 recommendation is valid in patients treated exclusively with GnRH antagonists.

3878

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Draft



3895 13. Endometrial thickness

PICO QUESTION: DOES MONITORING OF ENDOMETRIAL THICKNESS AFFECT THE EFFICACY AND SAFETY?

3896 Human endometrium has a key role in implantation process. Adequate endometrial development is
3897 required for pregnancy to occur. Thin endometrium on ultrasound during ovarian stimulation has been
3898 thought to be associated with poor success rates after IVF, even in the absence of prior intrauterine
3899 surgery or infection. At present, results from studies that investigated the relationship between
3900 endometrial thickness (EMT) and IVF outcomes are conflicting (Kasius et al., 2014). A meta-analysis by
3901 Kasius et al. reported a thin endometrium (≤ 7 mm) in 2.4% (260/10,724) of patients (Kasius et al., 2014).
3902 A more recent retrospective study reported 11% (57/517) of patients presenting with thin
3903 endometrium in ICSI cycles (Coelho Neto et al., 2015). However, in a large retrospective study by Holden
3904 et al. the proportion of patients with thin endometrium < 7 mm was 5.5% (347/6331) in IVF cycles
3905 (Holden et al., 2017).

3906 Evidence

3907 *There are no studies comparing monitoring endometrial thickness compared to no monitoring, which*
3908 *would be the ideal study to answer this question. Alternatively, we looked at studies investigating*
3909 *whether endometrial thickness is predictive for implantation and live birth.*

3910 A meta-analysis combining 22 prospective and retrospective studies (10,724 patients and cycles) and
3911 several more recent studies found EMT having little to no discriminatory capacity for clinical pregnancy
3912 (Table 9) (Griesinger et al., 2018, Kasius et al., 2014, Lamanna et al., 2008, Rehman et al., 2015,
3913 Shakerian et al., 2021, Zhao et al., 2014). In addition, the study by Griesinger et al. reported that the
3914 independent contribution of EMT (assessed on day of embryo transfer) to live birth likelihood is small
3915 and may result from (undetermined) confounding factors. If EMT indeed is an independent factor
3916 affecting outcome, this finding implies that at a baseline live birth rate of 20% an increase of 2 mm in
3917 EMT should result in an increase of the live birth rate of $\sim 1.6\%$ (Griesinger et al., 2018). In contrast, a
3918 recent retrospective cohort study including 5133 fresh ET cycles reported that EMT was found to be a
3919 significant independent predictor of LBR (OR 0.935, 95% CI 0.908-0.962), in addition to age, previous
3920 parity, ovarian response and number of available embryos (Simeonov et al., 2020). Similarly, a large
3921 retrospective cohort study, including 42132 fresh ET cycles reported significant associations between
3922 EMT and clinical pregnancy rate (adjusted OR 1.05, 95% CI 1.06-1.08) and EMT and live birth rate
3923 (adjusted OR 1.04, 95% CI 1.03-1.05) (Xu et al., 2021). Meanwhile, the miscarriage rate was significantly
3924 declined by 8% (OR 0.92, 95% CI 0.90-0.95) with each mm increment of EMT. These results were
3925 confirmed in a recent large retrospective cohort study, including 11,738 fresh IVF/ICSI cycles. After
3926 controlling for potential confounding factors, EMT had a significant, but small, effect on the clinical
3927 pregnancy rate (adjusted OR 1.07, 95% CI 1.05-1.08) (Wang et al., 2023). In addition, EMT resulted
3928 inversely proportional to ectopic pregnancy rate.



3929 Table 9: Accuracy of EMT in predicting pregnancy outcome

Predictive power of EMT on pregnancy outcome		
Study	Cohort (n)	ROC-AUC
Kasius 2014	10.724 women and cycles	0.56
Other studies:		
Lamanna 2008	685 women	<0.70
Zhao 2014	3319 women	0.60
Rehman 2015	282 women	0.88
Griesinger 2018	1483 women	0.53

3930

3931 A meta-analysis²³ combining 30 cohort studies (9 prospective and 21 retrospective) including 88,056
 3932 cycles reported that women with lower EMT had a lower chance of clinical pregnancy than those with
 3933 a higher EMT (OR 0.61, 95% CI 0.52-0.70) irrespective of fresh or frozen embryo transfer (Gao et al.,
 3934 2020). When looking only at the prospective studies with fresh transfer and a cutoff value of >8 mm,
 3935 no significant association between EMT and pregnancy rates were found. Similar results were found
 3936 when pooling the 11 studies reporting on live birth rate/ongoing pregnancy rates, with a lower chance
 3937 of live birth/ongoing pregnancy with lower EMT versus higher EMT (OR 0.60, 95% CI 0.48-0.73). Again,
 3938 no association was found when only including prospective studies with fresh embryo transfer.
 3939 Furthermore, there was no significant association between EMT and incidence of abortion rate (OR
 3940 1.33; 95% CI 0.98-1.80).

3941 Several more recent studies and studies not included in the meta-analysis also reported a significantly
 3942 lower probability of conceiving with EMT <8 mm as compared to EMT >8 mm (Table 10) (Aydin et al.,
 3943 2013, Gallos et al., 2018, Rehman, et al., 2015). A large retrospective cohort study reporting on the
 3944 results of 5546 fresh embryo transfers also found a higher rate of obstetric complications, such as
 3945 preeclampsia, placental abruption, placenta previa, small for gestational age and preterm delivery with
 3946 EMT <7.5 mm (adjusted OR 1.53, 95% CI 1.03-2.42) (Oron et al., 2018).

3947 Table 10: Probability of pregnancy with thin endometrium.

Probability of pregnancy with EMT			
Study	Cohort (n)	<8 mm	>8 mm
Gao 2020	88.056 cycles	OR 0.61, 95% CI 0.52-0.70	
Gallos 2018	45.279 cycles	15.6%	33.1%
Rehman 2015	282 women	5%	57.2%
Aydin 2013	593 women	7.1%	35.5%-43.9%

3948

3949 A large retrospective cohort study (3319 women) reported significant thicker EMT on the hCG day in
 3950 the clinical pregnancy group compared with the not pregnant group (11.0±2.2 vs. 10.3±2.2 mm) (Zhao,
 3951 et al., 2014). In contrast, a large prospective study in 435 women reported no difference in endometrial

²³ The meta-analysis from Kasius 2014 cited here in the 2019 version was replaced by a more recent meta-analysis. Data from the studies by Wu et al., 2014, Yuan et al., 2016, Ribeiro et al., 2018 - previously cited in table 9 - are included in the meta-analysis by Gao et al. 2020 and therefore not mentioned separately anymore.



3952 thickness between pregnant and non-pregnant patients (11.2 mm (9.8-12.7) vs. 11.1 mm (9.5-12.9)
3953 (Zhang et al., 2016).

3954 The thinnest endometrial thickness at which pregnancy occurred was 3.7 mm, in the study by Holden
3955 et al. and 5.6 mm in the study by Coelho Neto et al. Both pregnancies resulted in a live birth (Coelho
3956 Neto et al., 2015, Holden et al., 2017).

3957 Recommendations

Routine monitoring of endometrial thickness during
ovarian stimulation is probably not recommended. [2019]

Conditional ⊕○○○

3958

The guideline group suggests performing a single
measurement of the endometrium during ultrasound
assessment on the day of triggering or oocyte pick-up to
counsel patients on potentially lower pregnancy chance.
[2019]

GPP

3959

3960 Justification

3961 There are indications that thin endometrium is related to lower ongoing/clinical pregnancy chances as
3962 an independent factor. This condition of thin endometrium occurs infrequent (2-5%). Interventions to
3963 correct thin EMT have little rational basis and should be abandoned until contrary evidence arises.

3964 There are indications that thin endometrium is also associated with obstetric complications, even
3965 though rare (Lai et al., 2024, Oron et al., 2018). These observations, however, are only supported by a
3966 few retrospective cohort studies and the evidence is not solid.

3967 A single ultrasound assessment is necessary to identify patients with very thin or very thick EMT, and
3968 appropriate diagnostic work-up should be done.

3969

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- 4019



4020 14. Criteria for final oocyte maturation

PICO QUESTION: IS THE OUTCOME OF OVARIAN STIMULATION DEPENDENT ON THE CRITERIA FOR FINAL OOCYTE MATURATION?

4021 FOLLICLE SIZE

4022 Evidence

4023 A meta-analysis including 7 RCTs investigating the effect of postponing final oocyte maturation by 24-
4024 48 hours. There was no significant difference in live birth rate (3 RCT, RR 1.14, 0.46-2.83, 354 women)
4025 or ongoing pregnancy rate per oocyte pick-up (4 RCT, RR 0.97, 95% CI 0.54–1.74, 743 women) between
4026 early hCG and the late hCG group. However, significantly more oocytes were retrieved in late hCG group
4027 than in early hCG group (4 RCT, MD 1.2, 95% CI 1.11–1.30, 743 women) (Chen et al., 2014).

4028 In the meta-analysis there was one study comparing triggering at different follicular sizes, the only trial
4029 identified by the literature search investigating this research question. In this RCT (190 women),
4030 triggering was performed when the leading follicle reached either 18 or 22 mm. There was no significant
4031 difference in live birth rate when trigger was administered when the leading follicle was 22 mm (35%
4032 (34/97)) compared to 18 mm (23% (21/93)) (RR 1.6 (0.98–2.47)). However, more women reached an
4033 ongoing pregnancy (38% (37/97)) compared with the 18-mm group (24% (22/93)) (RR 1.6, 95% CI: 1.03–
4034 2.5) and significantly more oocytes were retrieved (11.7 ± 5.7 vs. 9.7 ± 4.1) (Mochtar et al., 2011).

4035 Recommendations

The association of follicle size as a triggering criterion with outcome has not been sufficiently studied. Physicians may choose the follicle size upon which final oocyte maturation is triggered on a case to case basis. [2019]

Conditional ⊕⊕○○

4036

The decision on timing of triggering in relation to follicle size is multi-factorial, taking into account the size of the growing follicle cohort, the hormonal data on day of pursued trigger, duration of stimulation, patient burden, financial costs, experience of previous cycles and organizational factors for the centre. Most often, final oocyte maturation is triggered at sizes of several of the leading follicles between 16-22 mm. [2019]

GPP

4037

4038 Justification

4039 The available studies have compared, except for one (Mochtar et al., 2011), not different follicle sizes
4040 as trigger criteria but postponing hCG administration after a given sonographic follicular criterion had



4041 been reached. Later hCG administration is associated with the retrieval of more oocytes. An effect on
4042 any other efficacy or safety or patient-related outcome was either not studied or not demonstrated in
4043 a consistent (e.g. homogenous) way across studies.

4044 OESTRADIOL LEVEL

4045 Evidence

4046 There are no interventional studies investigating triggering based on oestradiol levels.

4047 Recommendations

The GDG does not recommend to base timing of final oocyte maturation triggering on oestradiol levels alone. [2019]

GPP

4048

4049 Justification

4050 No interventional study has been performed assessing the use of serum oestradiol as a criterion for
4051 when to trigger final oocyte maturation. Serum oestradiol levels during ovarian stimulation vary
4052 depending on the size of the growing follicular cohort, the distribution of follicles between different
4053 size classes within the growing cohort as well as the endocrine situation of the patient and the
4054 endocrine milieu of the stimulation cycle. The association of the serum oestradiol levels with clinical
4055 outcomes and OHSS risk has been studied in several observational studies, but management
4056 recommendations cannot be derived from these observational data.

4057 OESTRADIOL/FOLLICLE RATIO

4058 Evidence

4059 There are no interventional studies investigating triggering based on the oestradiol/follicle ratio.

4060 Recommendations

The GDG does not recommend to base timing of final oocyte maturation on oestradiol/follicle ratio alone. [2019]

GPP

4061

4062 Justification

4063 No interventional study has been performed assessing the use of serum oestradiol-to-follicle ratio as a
4064 criterion for when to trigger final oocyte maturation. The oestradiol-to-follicle ratio will vary depending
4065 on the size of the growing follicular cohort, the distribution of follicles between different size classes
4066 within the growing cohort as well as the endocrine situation of the patient and the endocrine milieu of
4067 the stimulation cycle. The association of the oestradiol-to-follicle ratio with clinical outcomes has been
4068 studied in several observational studies, but management recommendations cannot be derived from
4069 these observational data.



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4078

Draft for review



4079 15. Hormonal assessment on the day of final oocyte 4080 maturation

PICO QUESTION: IS HORMONAL ASSESSMENT ON THE DAY OF FINAL OOCYTE MATURATION RECOMMENDED?

4081 HCG TRIGGERED CYCLES

4082 PROGESTERONE

4083 Evidence

4084 A systematic review and meta-analysis, including 55,199 fresh embryo transfer cycles from 63
4085 prospective and retrospective studies, reported that serum progesterone levels above 0.8 ng/ml on the
4086 day of hCG administration was associated with significantly decreased odds of live birth/ongoing
4087 pregnancy rate (OR 0.72, 95% CI 0.56-0.94; OR 0.64, 95% CI 0.53-0.77; OR 0.62, 95% CI 0.57-0.69; OR
4088 0.67, 95% CI 0.55-0.81 for serum progesterone levels of 0.8-1.1 ng/mL, 1.2–1.4 ng/mL, 1.5–1.75 ng/mL,
4089 and 1.9–3.0 ng/mL, respectively) (Venetis et al., 2013). A meta-regression analyses suggested that the
4090 type of patient population (i.e., low responders, normal responders, high responders), the
4091 developmental stage of embryo at transfer (cleavage versus blastocyst stage), or the study design
4092 (retrospective vs prospective) did not modulate the conclusions. Based on an analysis of 37 studies
4093 reporting the number of oocytes collected, the mean number of cumulus oocyte complexes retrieved
4094 was significantly increased in patients with progesterone elevation compared with those without
4095 progesterone elevation. This finding was consistent across all progesterone elevation threshold groups,
4096 ranging from +1.9 in the 1.2–1.4 ng/mL to +3.1 COCs in the 1.5–1.75 ng/mL group (Venetis et al., 2013).

4097 A retrospective study including 4,651 patients undergoing their first IVF cycles reported significantly
4098 lower cumulative live birth rates in patients with low ovarian response (≤ 5 oocytes collected),
4099 intermediate ovarian response (6-19 oocytes collected) and high ovarian response (>19 oocytes
4100 collected), when serum progesterone levels on the day of HCG trigger was >1.5 ng/mL, 2.24 ng/mL and
4101 2.5 ng/mL, respectively. Adjusted analyses demonstrated an inverse relationship between serum
4102 progesterone levels on the day of HCG trigger and cumulative live birth rates in all groups (Bu et al.,
4103 2014).

4104 Based on sixteen studies, the same meta-analysis reported that serum progesterone elevation on the
4105 day of HCG trigger in the stimulation cycle was not associated with the probability of pregnancy
4106 achievement in a subsequent frozen–thawed cycle. This finding was consistent across all progesterone
4107 threshold groups (Venetis et al., 2013).

4108 A multicentre retrospective study compared cumulative live birth rate over 24 months following a
4109 freeze all approach between patients with serum progesterone levels <1.50 ng/mL and >1.50 ng/mL on
4110 the day of hCG trigger. There were 471 patients in each group, who were matched for age and oocyte
4111 yield. Cumulative LBR was similar the two study groups (29.3% and 28.2%) (Racca et al., 2021).



4112 Recommendations

It is probably recommended to measure serum progesterone levels on the day of final oocyte maturation in cycles aimed for a fresh embryo transfer. [2025]	Conditional ⊕○○○
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4113

If serum progesterone levels are high, the patient should be counselled about potentially lower ongoing pregnancy/live birth rates. The decision to defer embryo transfer should include other factors (number of oocytes, number of embryos, and embryo quality). [2025]	GPP
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4114

4115 Justification

4116 Patients cannot be randomized to have different serum progesterone levels on the day of HCG trigger
4117 so decisions have to be based on observational studies. Observational studies consistently report
4118 decreased live birth/ongoing pregnancy rate and even suggest a gradient effect, i.e., higher
4119 progesterone levels are associated with lower ongoing pregnancy/live birth rates, supporting a causal
4120 relationship. While a 2024 systematic review and meta-analysis reported similar results for day 3
4121 transfers, live birth and clinical pregnancy rates were not significantly affected by elevated
4122 progesterone in a subgroup analysis limited to day 5 transfers (Lim et al., 2024). However, the pooled
4123 analysis result seems to be possibly shifted by one retrospective study, which has a small sample size,
4124 an unusually high rate of progesterone elevation and unusually low rate of live birth and clinical
4125 pregnancy rate in the non-progesterone elevation group (Huang et al., 2015) Thus, the GDG
4126 acknowledges some controversy but still holds the opinion that elevated progesterone would affect
4127 fresh day 3 and day 5 similarly. An indirect study suggest that effect of elevated progesterone levels is
4128 mediated through endometrial advancement and resultant embryo – endometrium asynchrony, not
4129 through a perceivable effect on oocyte developmental potential (Racca et al., 2021). Thus, deferring
4130 embryo transfer to a frozen embryo transfer cycle without endometrial advancement seems to provide
4131 similar live birth rates with non-progesterone elevated cycles. Given that a solution exists for the
4132 problem it is justifiable to diagnose progesterone elevation and forfeit a fresh embryo transfer as
4133 necessary.

4134 OESTRADIOL

4135 Evidence

4136 A systematic review and meta-analysis, including 3 cohort studies and 641 cycles, investigated whether
4137 the probability of live birth/ongoing pregnancy (≥ 12 weeks of gestation) or clinical pregnancy (up to 6–
4138 8 weeks of gestation) after ovarian stimulation for IVF, using gonadotropin-releasing hormone (GnRH)
4139 analogues and gonadotrophins is associated with serum oestradiol levels on the day of triggering final
4140 oocyte maturation with hCG (Karatasiou et al., 2020). While the odds of achieving a clinical pregnancy



4141 gradually declined with higher oestradiol levels, demonstrating a gradient effect, the difference was
4142 not statistically significant.

4143 A retrospective study including 1,141 non-PCOS patients with an AFC of >7 who underwent a long luteal
4144 GnRH agonist or a flexible GnRH antagonist protocol reported that peak serum oestradiol level on the
4145 day of hCG administration was not associated with cumulative live birth rate in a multivariable analysis
4146 (OR 0.995, 95% CI 0.98-1.01) (Zhang et al., 2019). A quantitative analysis suggested that until a peak
4147 oestradiol level of <2,185 pg/ml, the cumulative LBR statistically significantly increased by about 12%
4148 with every 100 pg/ml increase of the peak oestradiol level. Between peak oestradiol levels of 2,185 and
4149 6,136 pg/ml, the cumulative LBR only slightly decreased (0.4% per 100 pg/mL increase in peak
4150 oestradiol). When the peak oestradiol level that was higher than 6,136 pg/mL, a more prominent
4151 decrease in cumulative LBR was observed (10% per 100 pg/ml increase in peak E2), but this was short
4152 of statistical significance (Zhang et al., 2019).

4153 A retrospective study divided 1,771 fresh embryo transfer cycle following ovarian stimulation with a
4154 long luteal GnRH agonist or a GnRH antagonist protocol into six groups based on peak oestradiol levels
4155 on the day of hCG administration as the following; ≤1000 pg/mL, 1001–2000 pg/mL, 2001–3000 pg/mL,
4156 3001–4000 pg/mL, 4001–5000 pg/mL, and > 5000 pg/mL (Li et al., 2019). Clinical pregnancy rate
4157 gradually increased from <100 pg/mL group to 4001–5,000 pg/ml and declined in the >5,000 pg/mL
4158 group. Similar pattern was observed for number of MII oocyte counts.

4159 A retrospective study included 3,393 patients younger than 40 years undergoing IVF with a long luteal
4160 GnRH agonist protocol (Wang et al., 2017). Cycles with a serum oestradiol level >3,757 pg/mL on the
4161 day of HCG trigger were reported to have a significantly higher mean number of oocytes (14.4±5.3 vs.
4162 7.4±3.9), 2PN oocytes (9.56±4.18 vs. 4.98±2.97), good-quality embryos (5.69±3.45 vs. 2.96±2.27), as
4163 well as higher risk of OHSS (3.9% vs 0.6%). Live birth (47.4% vs. 43%) and clinical pregnancy (57.2% vs.
4164 52.1%), were significantly higher in the high oestradiol group (Wang et al., 2017).

4165 Recommendations

It is not recommended to routinely measure serum oestradiol levels on the day of HCG trigger in ovarian stimulation cycles with an intent for a fresh embryo transfer.

Strong ⊕○○○

4166

4167 Justification

4168 Patients cannot be randomized to have different serum oestradiol levels on the day of hCG trigger,
4169 therefore decisions have to be based on observational studies. Observational studies consistently
4170 suggest that serum oestradiol levels are poor predictors of live birth/ongoing pregnancy rate beyond
4171 an association between serum oestradiol levels and oocyte yield. Serum oestradiol levels are poor
4172 predictors of obstetric and neonatal adverse events. While serum oestradiol level is strongly correlated
4173 with follicle count, serum oestradiol levels considerably overlap between patients who develop
4174 moderate severe OHSS following a hCG trigger and fresh embryo transfer.



4175 LH

4176 Evidence

4177 A retrospective study including 3,059 patients who underwent a fresh embryo transfer following
4178 ovarian stimulation with an hCG triggered GnRH antagonist protocol, divided patients in three
4179 categories of anticipated ovarian response (low: AMH <1.1 ng/mL or AFC <5 or previous low response;
4180 normal: AMH >1.1 ng/mL or AFC >5 and regular menstrual cycles) and PCOS (as per Rotterdam criteria)).
4181 Patients in each anticipated ovarian response category were categorized according to quartiles of
4182 serum LH levels on the day of the trigger (<25th percentile, 25th to 75th percentile and >75th percentile).
4183 Compared to patients with anticipated normal ovarian response and LH levels >75th percentile, patients
4184 in <25th percentile (adjusted OR 0.662, 95%CI 0.508-0.863) and 25th-75th percentile categories (adjusted
4185 OR 0.791, 95% CI 0.633-0.988) had significantly lower live birth rates than those in the >75th percentile
4186 category. Likewise, patients with PCOS and LH levels <25th percentile also had significantly lower live
4187 birth rates in comparison to patients with LH levels >75th percentile (adjusted OR 0.479, 95% CI 0.277-
4188 0.828). Live birth rates were not correlated with LH quartiles in patients with an anticipated low ovarian
4189 response (Zhou et al., 2023).

4190 A retrospective study including 4,502 fresh embryo transfers following ovarian stimulation with an hCG
4191 triggered short GnRH agonist protocol, divided patients in five categories based on serum LH levels on
4192 the day of HCG trigger (Group A: LH ≤0.5 IU/L, Group B: 0.5 IU/L < LH ≤1.2 IU/L, Group C: 1.2 IU/L < LH
4193 ≤2.0 IU/L, Group D: 2.0 IU/L < LH ≤5.0 IU/L, Group E: LH >5 IU/L). Regression analyses showed that each
4194 unit increase in LH levels on the day of HCG trigger was inversely correlated with the number of oocytes
4195 retrieved (adjusted OR -0.351, 95% CI -0.453 to -0.249). However LH levels were not associated with
4196 live birth rates (Zhang et al., 2022).

4197 A retrospective study included 9,334 a fresh ART cycles following ovarian stimulation with an hCG
4198 triggered long luteal GnRH agonist or a flexible GnRH antagonist (Luo et al., 2023). Cycles were divided
4199 in three categories based on tertiles of serum LH levels on the day of hCG trigger. Multivariable
4200 regression analysis suggested that higher LH levels were associated with significantly higher live birth
4201 and clinical pregnancy rates with both protocols. However, in GnRH antagonist cycles, the difference
4202 was only significant for when comparing the third tertile with the first tertile (Luo et al., 2023).

4203 Recommendation

It is not recommended to measure serum LH levels on the day of HCG trigger in ovarian stimulation cycles aimed for a fresh embryo transfer.

Conditional ⊕○○○

4204

4205 Justification

4206 The available studies are limited by retrospective design and complicated analytical approach using
4207 different thresholds. Their results are inconsistent and do not provide actionable conclusions.

4208



4209 GnRH AGONIST TRIGGERED CYCLES

4210 PROGESTERONE

4211 Evidence

4212 A retrospective study including 1,484 GnRH agonist triggered PPOS cycles reported that serum
4213 progesterone levels on the day of trigger were not associated with the risk of inadequate response to
4214 the agonist trigger defined as a serum LH level <15 IU/L, 12 h after the agonist trigger (Lu et al., 2016).

4215 A retrospective study including 3,334 agonist triggered GnRH antagonist cycles reported similar serum
4216 progesterone levels on the day of agonist trigger between cycles with an adequate and with an
4217 inadequate response, defined as the ratio between the total number of oocytes retrieved and the
4218 number of follicles with a mean diameter >10 mm on the day of/prior to the trigger <45% (1.3±0.8 vs.
4219 1.4±0.9 ng/ml, respectively) (Popovic-Todorovic et al., 2019).

4220 OESTRADIOL

4221 Evidence

4222 A retrospective study including 1,484 GnRH agonist triggered PPOS cycles reported significantly
4223 different serum oestradiol levels on the day of trigger between cycles with an adequate and inadequate
4224 response to the GnRH agonist trigger defined as a serum LH level <15 IU/L, 12 h after the agonist trigger
4225 (2,753.23 ± 1,616.34 vs. 1,906.41 ± 1,656.87) (Lu et al., 2016).

4226 A retrospective study including 3,334 GnRH agonist triggered GnRH antagonist cycles reported
4227 significantly different serum oestradiol levels on the day of trigger between cycles with an adequate
4228 and with an inadequate response, defined as the ratio between the total number of oocytes retrieved
4229 and the number of follicles with a mean diameter >10 mm on the day of/prior to the trigger <45%
4230 (2796.2±1752.6 vs. 2277.5±1728.1 pg/mL, respectively) (Popovic-Todorovic et al., 2019).

4231 A retrospective study including 502 GnRH agonist triggered GnRH antagonist cycles reported that
4232 serum oestradiol levels on the day of trigger were significantly different between cycles with and
4233 without an adequate post-trigger LH response defined as serum LH level >15 IU/L 12 hours after the
4234 GnRH agonist trigger (3242 ± 1233 vs. 2564 ± 1257 pg/ml, respectively) (Kummer et al., 2013).

4235 LH

4236 Evidence

4237 A retrospective study including 1,747 GnRH agonist triggered GnRH antagonist cycles reported that
4238 serum LH level on the day of trigger was not associated the risk of low oocyte maturation rate, defined
4239 as <75% of all oocytes collected being at MII stage, or the risk of having a low oocyte recuperation rate,
4240 defined as the ratio of collected oocytes over the number of follicles measuring ≥12 mm on the day of
4241 trigger below the 10th percentile (Gambini et al., 2024).

4242 A retrospective study including 1,484 GnRH agonist triggered PPOS cycles reported that serum LH levels
4243 on the day of trigger were not associated the risk of inadequate response to the agonist trigger defined
4244 as a serum LH level <15 IU/L 12 h after the agonist trigger (Lu et al., 2016).



4245 A retrospective study including 3,334 GnRH agonist triggered GnRH antagonist cycles reported similar
4246 serum LH levels on the day of agonist trigger between cycles with an adequate and with an inadequate
4247 response, defined as the ratio between the total number of oocytes retrieved and the number of
4248 follicles with a mean diameter >10 mm on the day of/prior to the trigger <45% (Popovic-Todorovic et
4249 al., 2019).

4250 A retrospective study including 502 GnRH agonist triggered GnRH antagonist cycles reported that
4251 serum LH levels on the day of trigger were significantly different between cycles with and without an
4252 adequate post trigger LH response defined as serum LH level >15 IU/L 12 hours after the agonist trigger
4253 (2.1±1.9 vs 1±1.4 IU/L, respectively) (Kummer et al., 2013).

4254 15.2.4 OVERALL RECOMMENDATION

4255 Recommendation

It is not recommended to measure serum oestradiol, progesterone or luteinizing hormone levels on the day of a GnRH agonist trigger in freeze all cycles. Conditional ⊕○○○

4256

4257 Justification

4258 Serum levels of oestradiol, progesterone and luteinizing hormone levels largely overlap in cycles with
4259 and without an adequate response to a GnRH agonist trigger, hence they do not have a discriminatory
4260 value. Patients at risk of inadequate response, e.g. patients with hypogonadotropic hypogonadism,
4261 prolonged combined contraceptive use, etc. can be identified at the beginning of the stimulation cycle.
4262 When a fresh embryo transfer is not intended serum progesterone levels on the day of trigger would
4263 not affect live birth rates with a subsequent frozen embryo transfer. While the studies reporting similar
4264 cumulative live birth rates and live birth rates after the first frozen embryo transfer between stimulation
4265 cycles with and without progesterone elevation were not performed exclusively GnRH agonist triggered
4266 cycles, available evidence does not support a carryover effect of endometrial advancement due to
4267 progesterone elevation in the stimulation cycle.

4268

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4325 16. Criteria for cycle cancellation

PICO QUESTION: WHICH CRITERIA FOR CYCLE CANCELLATION ARE MEANINGFUL REGARDING PREDICTED LOW/HIGH OOCYTE YIELD?

4326 Since the year 1983 –when the term „poor responder” was described for the first time (Garcia et al.,
4327 1983), no international consensus regarding the definition of a poor response was available and
4328 different definitions were used. In 2011, the European Society of Human Reproduction and
4329 Endocrinology (ESHRE) defined poor response as: ‘cycle cancellation or retrieval of fewer than four
4330 oocytes with a conventional ovarian stimulation protocol’ (Ferraretti et al., 2011).

4331 Similarly, there is no international consensus definition for high response, which would help to identify
4332 women who can develop OHSS and allow undertaking interventions to avoid developing the condition.

4333 LOW OOCYTE YIELD

4334 Evidence

4335 *The occurrence of poor response is reported to vary between 5.6% and 35.1% or 9% to 24 % depending*
4336 *on the definition of poor response (Oudendijk et al., 2012). The decision making to stop the treatment,*
4337 *or to encourage to start another cycle is always difficult in respect to low number of oocytes and should*
4338 *be individually taken. Other factors, which influence pregnancy rate (e.g. age of patient) and burden of*
4339 *therapy, should be taken into account. The data also demonstrated that the pregnancy could still occur*
4340 *even in the first cycle the women is defined as poor responder (Baka et al., 2006).*

4341 In a meta-analysis combining prospective and retrospective cohort studies, the pooled estimate of
4342 pregnancy rate for poor responders was 14.8%, compared with 34.5% for normal responders (6 cohort
4343 studies, n=14338 women/cycles) (Oudendijk et al., 2012). The chance of pregnancy in respect to
4344 number of oocytes varied across studies. Women with 1 oocyte retrieved had 0-7%, 2 oocytes 4.3-
4345 15.2%, 3 oocytes 8.7-15.6%, and 4 oocytes 11.5–18.6% (4 cohort studies, 8744 women/cycles)
4346 (Oudendijk et al., 2012). Finally, in one study where 5 oocytes were obtained, pregnancy rate was up
4347 to 22 % (Oudendijk et al., 2012, Timeva et al., 2006). A more recent, large retrospective study reported
4348 a predicted live birth rate of 2% (n=541 cycles, 95% CI 2-3%) in women >40 years of age with one oocyte
4349 retrieved (Sunkara et al., 2011).

4350 In a retrospective study, it was examined whether IVF stimulation that results in one or two mature
4351 follicles should proceed to oocyte retrieval. The treatment outcomes were stratified in age groups (≤ 34 ,
4352 35-39, ≥ 40 years) (Shrem et al., 2022). The number of MII oocytes retrieved was 1.7 ± 0.9 , which did not
4353 differ between the age groups (<34: 1.8 ± 0.7 , 35-39: 1.7 ± 1.0 , ≥ 40 : 1.7 ± 0.8). There was however a
4354 significant difference in live birth rate per cycle between women ≤ 34 years (15.6%) and 35-39 years
4355 (6.5%) and ≥ 40 years (2.7%). In regression models, for LB, age was the only significant predictor. The
4356 change in pregnancy rate or LB as a function of age is dependent on AFC, suggesting that AFC is an
4357 important independent predictor which is more significant as age decrease.

4358 A large prospective study (1012 women, long GnRH agonist protocol) reported no live birth in women
4359 with AFC <4 (0%), but a live birth rate of 5% with an AFC of 4 (Jayaprakasan et al., 2012). The presence
4360 of one or two follicles in poor responders still could lead to obtain pregnancy. A large retrospective



4361 study (800 cycles, long GnRH agonist/GnRH antagonist protocols) in poor responders with 1 or 2 follicles
4362 >12 mm after ovarian stimulation, reported a clinical pregnancy rate of resp. 5.4% (12/223) and 9.2%
4363 (53/577) and an ongoing pregnancy rate of resp. 4.5% (10/223) and 7.6% (44/577) (Nicopoullos and
4364 Abdalla, 2011). A more recent, large retrospective study (256.381 cycles) reported a live birth rate of
4365 17% when the number of retrieved oocytes was between 0-5 (Steward et al., 2014).

4366 HIGH OOCYTE YIELD

4367 Evidence

4368 *The incidence of severe OHSS reported in clinical studies varies from 2% (Papanikolaou et al., 2006) to*
4369 *almost 9% (Toftager et al., 2016). The incidence of high response varied from >14 to >16 retrieved*
4370 *oocytes (Broer et al., 2013). It has been demonstrated in several prospective studies that a high number*
4371 *of growing follicles is an independent predictor of OHSS (Jayaprakasan, et al., 2012, Papanikolaou, et*
4372 *al., 2006).*

4373 A large prospective study with 2362 women advised cycle cancellation with >30 follicles of 12 mm
4374 during OS with long GnRH agonist protocol (Mathur et al., 2000). In a large prospective cohort study
4375 with 1801 women (2524 cycles), the threshold of ≥ 18 follicles ≥ 11 mm during OS with GnRH antagonist
4376 protocol predicted severe OHSS with 83% sensitivity rate with a specificity as high as 84% (Papanikolaou
4377 et al., 2006). According to the SART registry, analysis of 256.381 cycles revealed that retrieval of >15
4378 oocytes significantly increases the risk of OHSS and does not lead to an increased live-birth rate in fresh
4379 cycles (Steward et al., 2014). A recent large retrospective analysis of the Engage, Ensure and Trust trials
4380 found that the threshold of 19 follicles of ≥ 11 mm on hCG day predicted moderate to severe OHSS with
4381 62.3% sensitivity and 75.6% specificity (ROC-AUC 0.73), and predicted severe OHSS with 74.3%
4382 sensitivity and 75.3% specificity (ROC-AUC 0.77) in GnRH antagonist protocol (Griesinger et al., 2016).

4383 There was a strong association between the number of oocytes and LBR; LBR rose with an increasing
4384 number of oocytes up to 15, plateaued between 15 and 20 oocytes and steadily declined beyond 20
4385 oocytes. The LBR for women with 15 oocytes retrieved in age groups 18–34, 35–37, 38–39 and 40 years
4386 and over was 40, 36, 27 and 16% respectively (Sunkara et al., 2011).

4387 Recommendations

A low response to ovarian stimulation alone is not a reason
to cancel a cycle. [2019]

Strong ⊕○○○

4388

The physician should counsel the individual unexpected low
responder regarding pregnancy prospects and decide
individually whether to continue this cycle. [updated]

GPP

4389

In GnRH agonist cycles with an ovarian response of ≥ 19
follicles of ≥ 11 mm, there is an increased risk of OHSS and

Strong ⊕○○○



preventative measures are recommended, which should include primarily cancelling final oocyte maturation trigger. [updated]

4390

In GnRH antagonist cycles, withholding GnRH agonist triggering may still be considered in women with extremely high ovarian response. [2025]

GPP

4391

4392 Justification

4393 Reported pregnancy rates among low responders to ovarian stimulation differ between O-max reported
4394 18%. These differences could be explained by the exact number of oocytes retrieved, as well as the age
4395 of the patient and indication for treatment. Although pregnancy rates may be low, they are not absent
4396 per se.

4397 For an expected low responder, a cycle should not be cancelled due to low response. The GDG assumes
4398 that pregnancy prospects, costs etc. have been considered before starting the ovarian stimulation cycle.

4399 For an unexpected low responder, the GDG recommends the physician to counsel patients individually
4400 regarding pregnancy prospects and the decision to continue this cycle.

4401 Regarding a high response there are also no solid criteria to cancel a cycle. A high response identifies
4402 women most at risk for OHSS. The risk of OHSS and the number of growing follicles, is not a linear
4403 connection. There is probably a threshold effect, however, this is currently unknown. The current
4404 evidence comes from studies in GnRH antagonist cycles. The study by Griesinger et al. did not include
4405 PCOS patients, in contrast, the study by Papanikolaou did, explaining the lower threshold used in that
4406 study. Therefore, preventive measures are recommended which should include cycle cancellation.

4407 In GnRH antagonist cycles, withholding GnRH agonist triggering may still be considered in women with
4408 extremely high ovarian response (Berkovitz-Shperling et al., 2024). The GDG could not provide a
4409 threshold for this extremely high ovarian response, because the significance of this response could vary
4410 based on individual patient clinical characteristics .

4411

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PART F: Triggering ovulation and luteal support

17. Triggering of final oocyte maturation

PICO QUESTION: WHAT IS THE PREFERRED DRUG FOR TRIGGERING OF FINAL OOCYTE MATURATION IN TERMS OF EFFICACY AND SAFETY IN THE OVERALL IVF/ICSI POPULATION?

URINARY (UHCG) vs RECOMBINANT HUMAN CHORIONIC GONADOTROPHIN (RHCG)

Evidence

A Cochrane systematic review and meta-analysis found no difference in live birth/ongoing pregnancy rate (7 RCT, OR 1.15, 95% CI 0.89-1.49, 1136 women), moderate to severe OHSS (3 RCT, OR 1.76, 95%CI 0.37-8.45, 417 women), moderate OHSS (1 RCT, OR 0.78, 95% CI 0.27-2.27, 243 women), mild to moderate OHSS (2 RCT, OR 1.00, 95%CI 0.42-2.38, 320 women), undefined OHSS (3 RCT, OR 1.18, 95%CI 0.50-2.78, 495 women) or number of oocytes (12 RCT, MD-0.11, 95% CI -0.70 to 0.47, 1744 women) between recombinant and urinary hCG when used for triggering final oocyte maturation (Youssef et al., 2016).

One RCT including 100 women compared 10.000 IU with 5000 IU of urinary hCG for triggering final oocyte maturation in the long GnRH agonist protocol (Shaltout et al., 2006). There was no significant difference in pregnancy rate (not specified) (35.4% vs. 33.3%, incidence of OHSS (8.3% (4/48) vs. 2% (1/50)) or number of oocytes retrieved (7.4±3 vs. 7±3.5) between 10.000 IU and 5000 IU of uhCG for final oocyte maturation (Shaltout, et al., 2006).

One RCT including 80 PCOS patients randomized to receive 10.000 IU, 5000 IU, or 2500 IU of uhCG for triggering final oocyte maturation in the GnRH antagonist protocol as soon as 3 or more follicles of 17 mm or larger were present at ultrasound (Kolibianakis et al., 2007). There was no significant difference in ongoing pregnancy rate ((25.0% (7/28) vs. 30.8% (8/26) vs. 30.8% (8/26)), severe OHSS (1/28 vs. 1/26 vs. 0/26) or number of oocytes retrieved (median 14 vs. 11.5 vs. 9) between 10.000 IU, 5000 IU and 2500 IU uhCG (Kolibianakis et al., 2007).

One RCT including 180 women compared 500 µg with 250 µg recombinant hCG for triggering final oocyte maturation in the long GnRH agonist protocol (Madani et al., 2013). There was no significant difference in clinical pregnancy rate (34.5% (19/55) vs. 42.2% (19/45)), occurrence of OHSS (10% (6/60) vs. 6.7% (4/60)) or number of oocytes retrieved (12.25±5.30 vs. 12.40±6.44) between 500 µg and 250 µg rhCG (Madani et al., 2013).



4494 Recommendation

The use of recombinant hCG and urinary hCG is equally recommended for triggering final oocyte maturation in ovarian stimulation protocols. [2019]

Strong ⊕⊕○○

4495

A reduced-dose of 5.000 IU urinary hCG for final oocyte maturation is probably recommended over a 10.000 IU dose in GnRH agonist protocols, as it may improve safety. [2019]

Conditional ⊕○○○

4496

4497 Justification

4498 The Cochrane meta-analysis shows equal efficacy and safety for urinary and recombinant hCG. The
4499 grand majority of the trials (17 out of 18) included in the meta-analysis by Youssef et al. 2016,
4500 performed pituitary downregulation using a long GnRH agonist protocol, only one trial was performed
4501 using a GnRH antagonist protocol (Youssef et al., 2016). The evidence regarding antagonist protocol is
4502 inconclusive so the recommendation might not be applicable for GnRH antagonist cycles, although
4503 there is no evidence to suggest a difference in safety and efficacy.

4504 Different doses of uhCG have been described in the literature ranging from 2.000 IU to 10.000 IU.
4505 According to 2 RCTs, a reduced-dose of urinary hCG (5.000 IU) does not appear to affect the probability
4506 of pregnancy compared to conventional dose (10.000 IU). Similarly, data from 1 RCT suggests that a
4507 low dose (250µg) of recombinant hCG does not appear to influence the probability of pregnancy as
4508 compared to a higher dose (500 µg). The probability of OHSS was reduced when lower doses of hCG
4509 were administered but this did not reach statistical significance in any of the 3 RCTs. Lower doses of
4510 hCG could be considered when an unpredicted high response has occurred, and GnRH long agonist
4511 protocol is applied.

4512 **RECOMBINANT LH (rLH) VS URINARY HCG (uHCG)**

4513 Evidence

4514 The trials had administered different dosages of rLH which varied from 5000 IU (Manau et al., 2002)
4515 to 15000 IU and an additional 10000 IU three days post the first injection (2001).

4516 The Cochrane meta-analysis, mentioned before, reported no difference in live birth/ongoing
4517 pregnancy rate (2 RCT, OR 0.95, 95% CI 0.51-1.78, 289 women), moderate OHSS (2 RCT, OR 0.83, 95%
4518 CI 0.40-1.70, 289 women) or number of oocytes retrieved (2 RCT, MD-1.33, 95%CI -3.26 to 0.60, 103
4519 women) between rLH and uHCG when used for triggering final oocyte maturation (Youssef et al.,
4520 2016).



4521 Recommendation

It is not recommended to administer recombinant LH for triggering final oocyte maturation. [2019]

Strong ⊕○○○

4522

4523 Justification

4524 The available evidence is currently very limited to allow solid conclusions to be drawn. There was
4525 large heterogeneity between the three trials included with respect to study methods. Therefore, we
4526 cannot recommend the use of rLH to trigger final oocyte maturation.

4527 GnRH AGONIST TRIGGER VS HCG

4528 Evidence

4529 A systematic review and meta-analysis, including 9 RCTs and 1277 women compared GnRH agonist to
4530 hCG for final oocyte maturation (Beebeejaun et al., 2024). There was no significant difference
4531 observed between hCG and GnRH agonist trigger for live birth rate (RR 0.82; 95% CI 0.59–1.13, 3 RCT,
4532 723 women) or clinical pregnancy rate (RR 1.15; 95% CI 0.81–1.63; 3 RCT, 687 women).

4533 Recommendation

The use of GnRH agonist for final oocyte maturation is not recommended in the general IVF/ICSI population with fresh transfer, regardless of luteal phase support (with or without LH-activity). [updated]

Strong ⊕⊕○○

4534

4535 Justification

4536 Current evidence shows a disadvantage in ongoing/clinical pregnancy rate with GnRH agonist and
4537 conventional luteal support as compared to hCG in normal responders.

4538 Recent evidence shows that this disadvantage could be overcome by adding LH-activity to the LPS,
4539 however, this effect needs to be studied in a large RCT. Thus, with the current knowledge we cannot
4540 recommend GnRH agonist triggering with modified LPS for the overall IVF/ICSI population.

4541 There were no RCTs comparing GnRH agonist to hCG triggering in PPOS protocol. The only available
4542 evidence was a retrospective cohort study, which is insufficient evidence to formulate a
4543 recommendation.

4544 Although GnRH agonist trigger is associated with decreased OHSS rates, it is associated with low levels
4545 of endogenous LH secretion after triggering. In a retrospective cohort study, including 1747 patients,
4546 patients were divided into <10th percentile of oocyte recuperation rate (n=139) and >10th percentile
4547 oocyte recuperation rate (1281). Lower ovarian reserve and lower LH level 12-h post-triggering were
4548 predictive of lower ORR (OR 0.80 [95% CI 0.68–0.94]) and 0.80 [0.73–0.89], respectively (Gambini et
4549 al., 2024). In another retrospective cohort study, including 14066 patients, 51 patients were found to



4550 have empty follicle syndrome. After adjusting for confounding factors, PCOS was found to be a
4551 significant risk factor for EFS (aOR = 2.67; 95% CI 1.47-4.83) (Luo et al., 2024).

4552 GnRH agonist triggering for (predicted) high responder is discussed further in the guideline (chapter
4553 19).

4554 **TRIPTORELIN 0.1 MG VS HIGHER DOSAGES**

4555 **Evidence**

4556 One RCT including 165 oocyte donors compared different dosages (0.2 mg vs. 0.3 mg vs. 0.4 mg) of
4557 triptorelin for final oocyte maturation in GnRH antagonist protocol and reported no significant
4558 differences in number of oocytes retrieved (18.4±8.8 vs. 18.7±8.9 vs. 17.8±10.7) or mature oocytes
4559 (16.0±8.5 vs. 15.9±7.8 vs. 14.7±8.4). One case of OHSS in the 0.3 mg group (Vuong et al., 2016).

4560 **Recommendation**

If the GnRH agonist trigger with triptorelin is applied,
dosages ranging of 0.1-0.4 mg can be chosen.

GPP

4561

4562 **Justification**

4563 There are no studies investigating the direct comparison of hCG with different dosages of GnRH agonist
4564 trigger with triptorelin. Current evidence is derived from an RCT in oocyte donors, however, the
4565 guideline group thinks that the findings can be extrapolated to the general IVF population.

4566 **BUSERELIN 0.2 MG VS 0.5 – 1 – 2 MG**

4567 **Evidence**

4568 There are no studies investigating the direct comparison of hCG with different dosages of GnRH
4569 agonist trigger with buserelin. No controlled studies or RCT could be found comparing different
4570 dosages of Buserelin for final oocyte maturation. Therefore, no recommendation can be formulated
4571 regarding optimal dosage.

4572 **LEUPROLIDE 0.15 MG VS 0.5 – 1 – 2 - 4 MG**

4573 **Evidence**

4574 There are no studies investigating the direct comparison of hCG with different dosages of GnRH
4575 agonist trigger with leuprolide. No controlled studies or RCT could be found comparing different
4576 dosages of Leuprolide for final oocyte maturation. Therefore, no recommendation can be formulated
4577 regarding optimal dosage.



4578 DUAL AND DOUBLE TRIGGER

4579 Although GnRH agonist trigger is associated with decreased OHSS rates, it is associated with low levels
4580 of endogenous LH secretion after triggering, resulting in lower progesterone levels during the luteal
4581 phase. Several concepts of intensified luteal phase support have been formulated, among which the
4582 concept of dual and dual trigger. Dual trigger is defined as the simultaneous administration of hCG and
4583 GnRH agonist for final oocyte maturation. Staggered coadministration of GnRH agonist and hCG for
4584 final oocyte maturation, the double trigger, was proposed as another trigger option.

4585 DUAL TRIGGER

4586 Evidence

4587 A systematic review and meta-analysis²⁴ investigated the use of hCG and GnRH agonist (dual trigger)
4588 for final oocyte maturation and compared its efficacy to hCG in normal responders (Beebeejaun et al.,
4589 2024). Higher live birth rates were found with dual trigger (RR 1.31, 95% CI 1.00–1.70, 1 RCT, 496
4590 women) (Beebeejaun et al., 2024, Zhou et al., 2022). No significant difference was found between dual
4591 trigger and hCG trigger for final oocyte maturation for clinical pregnancy rate (RR 1.20, 95% CI 0.89–
4592 1.60, 3 RCT, 613 participants).

4593 In an RCT, participants were randomised to receive dual trigger (n=56) or hCG (n=57) for final oocyte
4594 maturation in normal responders (Keskin et al., 2023). No significant difference was observed for live
4595 birth rate (48.2% (27/56) vs. 31.5% (18/57)), however, clinical pregnancy rate was significantly higher
4596 with dual trigger compared to hCG alone (57.1% (32/56) vs. 38.5% (22/57)).

4597 In an RCT, participants with a normal ovarian reserve underwent ovarian stimulation for IVF/ICSI with
4598 final oocyte maturation triggered by either dual trigger (n=50) or hCG only (n=50) (Singh et al., 2023).
4599 No significant difference was observed in clinical pregnancy rate between dual trigger and hCG for final
4600 oocyte maturation (21% vs. 19.6%). No cases of OHSS were observed in either group.

4601 An RCT compared hCG 6500 IU with dual trigger (6500 IU hCG+0.2 mg GnRH agonist) in 192 normal
4602 responder women (Eftekhar et al., 2017). There was no significant difference in ongoing pregnancy rate
4603 (22.9% (20/93) vs. 24.2% (24/99)) between hCG and dual trigger. However, significantly more oocytes
4604 with dual trigger compared to hCG trigger (10.85± 4.71 vs. 9.35 ±4.35) (Eftekhar et al., 2017).

4605 In a retrospective cohort study one complete oocyte retrieval cycle (fresh+frozen) was compared for
4606 dual trigger and hCG trigger in the PPOS protocol in normal responders (Li et al., 2022). No significant
4607 difference was observed in cumulative live birth rate between dual trigger and hCG trigger only (40.72%
4608 (204/501) vs. 43.72% (247/565)).

4609 One RCT, compared dual trigger (n=168) to GnRH agonist (n=164) for final oocyte maturation in women
4610 of advanced age (Zhou et al., 2022). Comparing dual trigger to GnRH agonist for final oocyte maturation
4611 in women having fresh embryo transfer, no significant difference was observed for live birth rate (36.8%
4612 (7/19) vs. 20% (1/5)). No cases of moderate or severe OHSS were observed in either group.

²⁴ The systematic review by Ding et al., 2017 cited here in the 2019 version of the guideline was replaced by a more recent systematic review.



4613 Low responders

4614 A sub-analysis of a systematic review and meta-analysis investigated the use of hCG and GnRH agonist
4615 (dual trigger) for final oocyte maturation and compared its efficacy to hCG in poor responders (He et
4616 al., 2023). A significantly higher clinical pregnancy rate was observed (RR 2.2, 95% CI 1.05–4.61, 2 RCT,
4617 36 patients).

4618 In an RCT, women with a poor response to ovarian stimulation were randomised to receive dual trigger
4619 (n=57) or hCG (n=55) for final oocyte maturation (Keskin et al., 2023). Live birth per oocyte pick-up
4620 (17.5% (10/57) vs. 36.3% (20/55)) and clinical pregnancy rate per oocyte pick-up (26.3% (15/57) vs.
4621 52.7% (29/55)) was significantly lower with dual trigger compared to hCG trigger only.

4622 **Recommendation**

The addition of a GnRH agonist to hCG as a dual trigger for final oocyte maturation is probably not recommended for predicted normal responders. [2019]	Conditional ⊕⊕○○
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4623

The addition of a GnRH agonist to hCG as a dual trigger for final oocyte maturation is probably not recommended for low responders. [2025]	Conditional ⊕⊕○○
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4624

4625 **Justification**

4626 Available evidence has been rated of low quality. Current evidence in the form of RCT performed in
4627 normal responders suggests no improvement in the number of oocytes retrieved, with an improvement
4628 in pregnancy rate, but this finding needs to be further evaluated in well-designed RCTs. The additional
4629 intervention has not been shown to improve clinical outcomes in terms of live birth/ongoing pregnancy
4630 rate.

4631 Evidence in low responders is very poor. The evidence comes from three very small RCT reporting
4632 conflicting results.

4633 Regarding patients with history of low fertilization rate or high number of immature oocytes, the
4634 existing literature is limited by its observational nature. In addition, large differences are observed in
4635 the definition of low maturity rate, low fertilization rate, dose of hCG administered and most
4636 importantly lack of LBR and OHSS rate as an outcome. The dual trigger in this subgroup of patients,
4637 cannot be recommended until data on its efficacy and safety from RCT's are available.

4638 Dual triggering for (predicted) high responder is discussed further in the guideline (chapter 19).

4639 **DOUBLE TRIGGER**

4640 **Evidence**

4641 In an RCT, women with a normal response to ovarian stimulation and low oocyte maturation rate were
4642 randomised to receive either double trigger (40 and 36 hours before oocyte pick-up) or hCG only for



4643 final oocyte maturation (Yan et al., 2023). Cumulative live birth rate was significantly higher after double
4644 trigger compared to hCG only for final oocyte maturation (66.7% (24/36) vs. 36.0% (9/25)). Comparing
4645 double trigger to hCG for final oocyte maturation in women having fresh embryo transfer, no significant
4646 difference was observed for live birth rate (50% (2/4) vs. 36.4% (4/11)).

4647 In an RCT, poor responder patients were randomised to receive either double trigger, GnRH agonist
4648 trigger with hCG bolus on day of oocyte pick-up or hCG trigger for final oocyte maturation (Haas et al.,
4649 2019). There was no significant difference in ongoing pregnancy (18.2% (2/11) vs. 0 vs. 9.1% (1/11)) or
4650 number of MII oocytes retrieved (1.8 ± 1.4 vs. 2.1 ± 1.6 vs. 1.4 ± 1.5) between double trigger, GnRH
4651 agonist trigger or hCG trigger for final oocyte maturation.

4652 Conclusion

4653 There is too limited evidence to draw conclusions on the use of double trigger for final oocyte
4654 maturation for IVF/ICSI.

4655

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- 4716



4717 18. Luteal phase support (LPS)

4718 [KEY QUESTION: WHAT IS THE EFFICACY AND SAFETY OF LUTEAL SUPPORT PROTOCOLS?](#)

4719 18.1 PROGESTERONE

4720 Evidence

4721 A Cochrane meta-analysis reported a higher live birth/ongoing pregnancy rate with progestins
4722 compared to placebo/no treatment for luteal phase support (LPS) (5 RCT, OR 1.77, 95% CI 1.09-2.86,
4723 642 women) (van der Linden et al., 2015).

4724 Dosing

4725 The Cochrane meta-analysis also investigated the dosage of vaginal progesterone. Five studies
4726 compared a low dose (≤ 100 mg) with a high dose (≥ 100 mg) and reported no difference in live
4727 birth/ongoing pregnancy rate (5 RCT, OR 0.97, 95% CI 0.84-1.11, 3720 women) (van der Linden et al.,
4728 2015). After the publication of the Cochrane review, a small pilot study was conducted including 146
4729 women, investigating the effect of increasing the progesterone dosage in the mid-luteal phase in
4730 patients with progesterone levels below 15 ng/mL. There was no significant difference in live birth rate
4731 with increased progesterone dosage compared to original dosage (25% (9/36) vs. 17.1% (6/35)) (Aslih
4732 et al., 2017). Another small RCT including 111 women compared 600 mg vaginal progesterone
4733 (capsules) with 90 mg vaginal progesterone (gel) and reported no difference in live birth rate (52.8%
4734 (28/53) vs. 42.6% (20/47)) (Michnova et al., 2017).

4735 Administration route

4736 Several studies compared the efficacy of different administration routes for progesterone as LPS. An
4737 IPD meta-analysis compared the subcutaneous with the vaginal route (2 RCT, 1435 women) (Doblinger
4738 et al., 2016). Live birth rate was 35.3% (252/714) with subcutaneous progesterone vs. 37.6% (271/721)
4739 with vaginal progesterone (risk difference -0.02, 95% CI -0.07 to 0.03). There was no difference in
4740 incidence of OHSS between both groups (27/714 vs. 26/721; OR 1.04, 95% CI 0.60-1.81) (Doblinger, et
4741 al., 2016).

4742 Two newer RCTs also compared the efficacy of the subcutaneous and vaginal administration of
4743 progesterone for LPS (Moini et al., 2022, Salehpour et al., 2021). In the RCT by Moini *et al.*, patients
4744 undergoing their first IVF cycle were randomised to receive either subcutaneous (n=40) or vaginal
4745 progesterone (n=40) (Moini et al., 2022). The clinical pregnancy rate was significantly higher with the
4746 use of subcutaneous progesterone compared to vaginal (57.5% (23/40) vs. 32.5% (13/40)). In the RCT
4747 by Salehpour *et al.*, patients undergoing ICSI were randomised to receive either subcutaneous (n=100)
4748 or vaginal progesterone (n=100) (Salehpour et al., 2021). No significant difference in ongoing pregnancy
4749 rate was reported comparing subcutaneous with vaginal progesterone (37.1% (36/97) vs. 36%
4750 (36/100)).

4751 The Cochrane meta-analysis investigated vaginal/rectal compared to the oral route and reported no
4752 difference between groups for live birth/ongoing pregnancy rate (4 RCT, OR 1.19, 95% CI 0.83-1.69, 857
4753 women) (van der Linden et al., 2015). In a more recent RCT, infertile women were randomised on the



4754 day of final oocyte maturation trigger to receive either 400 mg/day oral micronised progesterone
4755 (n=430), 600 mg/day oral micronised progesterone (n=440) or vaginal progesterone (90 mg/day,
4756 n=440) (Niu et al., 2023). Comparing oral micronised progesterone at a dose of 400 or 600 mg/day with
4757 vaginal progesterone for LPS, no significant difference was observed for live birth rate (33.5% (144/430
4758 vs. 29.8% (131/440) vs. 35.5% (156/440). The number of adverse events was similar in the three groups:
4759 56 (13.0%) in the oral micronised progesterone 400 mg/day group, 60 (13.6%) in the oral micronized
4760 progesterone 600 mg/day group and 40 (9.1%) in the vaginal progesterone group.

4761 The Cochrane meta-analysis also investigated the vaginal/rectal compared to the intramuscular route
4762 and reported no difference in live birth/ongoing pregnancy rate (7 RCT, OR 1.37, 95% CI 0.94 to 1.99,
4763 2039 women) (van der Linden et al., 2015). A more recent RCT including 400 women also investigated
4764 the intramuscular compared to vaginal route and reported no difference in clinical pregnancy rate
4765 (26.5% (53/200) vs. 26.5% (53/200)) (Zargar et al., 2016). One very small RCT including 40 women
4766 investigated the intramuscular compared to the oral route and reported no difference in live birth rate
4767 (OR 0.71, 95% CI 0.14-3.66) (Iwase et al., 2008, van der Linden et al., 2015).

4768 Timing

4769 Six RCTs investigated the timing of LPS initiation (Baruffi et al., 2003, Fanchin et al., 2001, Gao et al.,
4770 2018, Mochtar et al., 2006, Sohn et al., 1999, Williams et al., 2001). One RCT compared starting LPS
4771 with progesterone on the day of oocyte retrieval with the day after oocyte retrieval in 233 women and
4772 reported no significant difference in live birth rate (46.6% (48/103) vs. 45.7% (43/94)) (Gao et al., 2018).
4773 Three RCTs compared starting LPS with progesterone on the evening of oocyte retrieval with starting
4774 on the evening of embryo transfer in respectively 103, 84 and 255 women and reported no significant
4775 difference in clinical pregnancy rate (respectively 27.4% vs. 28.8%; 42% vs. 29%; 28.1% (36/128) vs.
4776 29.1% (37/127)) (Baruffi et al., 2003, Fanchin et al., 2001, Mochtar et al., 2006). Only one study reported
4777 live birth rate and found no significant difference between groups (21.1% (27/128) vs. 20.5% (26/127);
4778 RR 0.97, 95% CI 0.60-1.56) (Mochtar et al., 2006). One newer RCT compared starting LPS with
4779 progesterone on the day of oocyte retrieval (n=86) with the day of embryo transfer (n=85) (Ghanem et
4780 al., 2021). No significant difference was observed in ongoing pregnancy rate when LPS was started on
4781 the day of oocyte retrieval or embryo transfer (38.3% (33/86) vs. 44.7% (38/85)). Two RCTs (respectively
4782 314 cycles and 385 women) compared starting LPS with progesterone before oocyte retrieval
4783 (respectively 12h before oocyte retrieval and at the evening of hCG trigger) with starting LPS after
4784 oocyte retrieval (Mochtar et al., 2006, Sohn et al., 1999). Mochtar *et al.* reported no significant
4785 difference in live birth (20% (26/130) vs. 21.1% (27/128); RR 0.94, 95% CI 0.58-1.52) or clinical
4786 pregnancy rate (23.1% (30/130) vs. 28.1% (36/128); RR 0.82, 95% CI 0.54-1.24) between groups
4787 (Mochtar et al., 2006). However, Sohn *et al.* found a significantly lower clinical pregnancy rate when
4788 LPS was started before oocyte retrieval compared to after (12.9% vs. 24.6%) (Sohn, et al., 1999). One
4789 small RCT including 126 women compared starting LPS with progesterone on day 3 or day 6 after oocyte
4790 retrieval and found a significantly lower clinical pregnancy rate when LPS was started on day 6
4791 compared to day 3 (44.8% vs. 61.0%) (Williams et al., 2001).

4792 A systematic review and meta-analysis²⁵ including 7 RCTs compared early progesterone LPS cessation
4793 (at the 11th or 14th day post embryo transfer after a positive hCG test) with continuing progesterone
4794 until week 6/7 or 10 (Watters et al., 2020). No significant difference was found for the probability of

²⁵ The meta-analysis by Liu et al., 2012 cited here in the previous version of the guideline was replaced by an updated meta-analysis.



4795 the pregnancy continuing to a live birth when comparing early or late cessation of LPS (RR 0.94, 95% CI
4796 0.84-1.00, 3 RCT, 830 participants).

4797 Recommendations

Progesterone is recommended for luteal phase support after IVF/ICSI. [2019]	Strong ⊕○○○
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Any of the previously mentioned administration routes (non-oral) for natural progesterone as luteal phase support can be used. [2019]	GPP
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The dosing of natural progesterone has evolved empirically, usually dosages used include: 50 mg once daily for intramuscular progesterone 25 mg once daily for subcutaneous progesterone 90 mg once daily for vaginal progesterone gel 200 mg three times daily for micronized vaginal progesterone in-oil capsules 100 mg two or three times daily for micronized vaginal progesterone in starch suppositories 400 mg two times daily for vaginal pessary. [2019]	GPP
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Starting of progesterone for luteal phase support should be in the window between the evening of the day of oocyte retrieval and day 3 post oocyte retrieval. [2019]	GPP
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Progesterone support should be administered until at least the day of the pregnancy test. [updated]	GPP
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4802
4803 Justification

4804 There are only a few, very old, RCTs comparing the use of progestins to placebo for LPS. Still,
4805 progesterone is recommended for luteal phase support for IVF/ICSI. Despite that the RCTs comparing
4806 use of progestins to placebo are scarce and old, the evidence clearly supports the use of progestins in
4807 the luteal phase. Very likely there are no future RCTs planned to challenge or confirm the existing
4808 evidence that progestins are crucial for the LPS.



4809 Start of luteal support has not been studied properly. More studies are necessary to investigate the
4810 need of luteal support and the correct timing to support endogenous progesterone levels. Until studies
4811 have been performed, luteal support should be provided in the window between the evening of the
4812 day of oocyte retrieval and D3 post oocyte retrieval.

4813 With the current evidence available, no major differences in efficacy have been found comparing the
4814 different administration routes of progesterone or duration of progesterone LPS.

4815 Current RCT on oral micronised progesterone showed non-inferiority to vaginal micronised
4816 progesterone (Niu et al., 2023). Despite these promising results, more data are necessary to be able to
4817 formulate a recommendation. Long-term offspring health studies are currently lacking.

4818 18.2 DYDROGESTERONE

4819 Evidence

4820 Daily dosages of 30 mg dydrogesterone are most frequently used for LPS.

4821 An IPD meta-analysis²⁶, including 2 RCTs, compared the use of dydrogesterone to vaginal micronised
4822 progesterone for LPS after IVF (Griesinger et al., 2020). Meta-analysis of the two RCTs with available
4823 IPD comparing dydrogesterone and vaginal micronised progesterone for LPS showed a significant
4824 higher live birth rate (OR 1.28; 95% CI 1.04-1.57, 2 RCT, 2065 women) and ongoing pregnancy rate (OR
4825 1.32; 95% CI 1.08-1.61, 2 RCT, 2065 women) in favour of dydrogesterone. The same systematic review
4826 included a meta-analysis of the aggregate data of all eligible studies (9 RCT) and found no significant
4827 difference for live birth rate (OR 1.14; 95% CI 0.99-1.32, 5 RCT, 4470 women) or ongoing pregnancy
4828 rate (OR 1.13; 95% CI 1.00-1.28, 9 RCT, 6312 women).

4829 An RCT including 207 women compared the use of oral dydrogesterone to vaginal micronised
4830 progesterone for LPS (Atarieh et al., 2024). The live birth rate was significantly lower with
4831 dydrogesterone compared to vaginal micronised progesterone for LPS (17.6% (23/103) vs. 41.3%
4832 (43/104)). No significant difference was reported in clinical pregnancy rates between groups (30.1%
4833 (31/103) vs. 41.3% (43/104)).

4834 A small RCT including 105 women compared the use of oral dydrogesterone with placebo for LPS and
4835 found no statistical difference in clinical pregnancy rate (29.6% (16/54) vs. 27.4% (14/51)) (Kupferminc
4836 et al., 1990).

4837 Recommendations

Dydrogesterone is probably recommended for luteal phase support. [2019]

Conditional ⊕⊕⊕○

4838 *There are pharmacovigilance reports of association between dydrogesterone exposure and increased*
4839 *risk of congenital malformations, although the observed relations cannot necessarily be translated into*
4840 *a conclusion on causality.*

²⁶ The meta-analysis by Barbosa et al., 2018 cited here in the previous version of the guideline has been replaced by an updated meta-analysis. The RCT by Griesinger et al., 2018 is included in the new meta-analysis and therefore no longer mentioned separately.



4841 Justification

4842 When compared to progesterone, oral dydrogesterone has similar live/birth ongoing pregnancy rate.

4843 An older meta-analysis reported on patient dissatisfaction, including 3 RCTs, the oral administration
4844 route was preferred over the vaginal route of progesterone in 2/3 RCTs (women in the 3rd RCT showed
4845 no difference in dissatisfaction) (Barbosa et al., 2018).

4846 As dydrogesterone is a synthetic, orally-active progestogen, metabolised into 20-
4847 dihydrodydrogesterone, and different in structure from natural progesterone, safety for the offspring
4848 is of key importance. Evidence from the two RCTs by Tournaye et al. and Griesinger et al. reported no
4849 difference in the rate of congenital anomalies as compared to natural progesterone (Griesinger et al.,
4850 2018, Tournaye et al., 2017). A recent systematic review and meta-analysis, including 6 RCTs (mainly in
4851 couples with recurrent miscarriage), reported that the risk ratio for congenital malformations with the
4852 use of dydrogesterone was 0.92 (95% CI 0.55-1.55, 6 RCT, 1512 women) compared to placebo, no
4853 treatment or other interventions (Katalinic et al., 2024), so that offspring safety does not seem
4854 jeopardised.

4855 However, a recent pharmacovigilance study using the WHO global safety database reported that a
4856 significant disproportionate reporting of birth defects was found with dydrogesterone when compared
4857 to any other drug in the study cohort, including natural progesterone (reporting OR 5.4, 95% CI 3.9–
4858 7.5) and to any other ART drug (ROR 6.0, 95% CI, 4.2–8.5) (Henry et al., 2025). Also, from the China
4859 maternal drug exposure birth cohort (DEBC) (Li et al., 2024), dydrogesterone exposure during the first
4860 trimester was correlated with higher incidence of birth defects (adjusted RR 1.13, 95% CI 1.06-1.21)
4861 compared to first trimester use of natural progesterone (aRR 1.05, 95% CI 0.97-1.13). It needs to be
4862 pointed out here that in these two pharmaco-vigilance studies, the observed relations cannot be
4863 translated into a conclusion on causality.

4864 No full agreement was reached within the guideline group regarding the strength of the
4865 recommendation. Arguments in favour of a strong recommendation were based on the safety approval
4866 by ICH-GCP standard and the historical use of dydrogesterone for early miscarriage prevention.
4867 However, the final recommendation was formulated as conditional, reflecting concerns about potential
4868 safety signals from recent pharmacovigilance data.

4869 18.3 OESTRADIOL SUPPLEMENTATION

4870 Evidence

4871 The Cochrane meta-analysis, mentioned before, reported no difference in live birth/ongoing pregnancy
4872 rate (9 RCT, OR 1.12, 95% CI 0.91-1.38, 1651 women) or OHSS (2 RCT, OR 0.58, 95% CI 0.20-1.68, 461
4873 women) between progesterone with oestradiol supplementation and progesterone alone (van der
4874 Linden et al., 2015). An RCT, more recent than the meta-analysis, including 220 women comparing
4875 progesterone and progesterone with oestradiol for LPS reported no significant difference in ongoing
4876 pregnancy rate (32.7% (36/110) vs. 36.3% (40/110)) (Ismail Madkour et al., 2016).

4877 In contrast, a RCT not included in the meta-analysis investigated the effect of adding oestradiol to a
4878 high dose of progesterone (200 mg vaginal capsules 3x/day + 100 mg intramuscular daily) for LPS in 240
4879 women and reported a significant higher clinical pregnancy rate with oestradiol supplementation in
4880 women undergoing the long GnRH agonist and short flexible GnRH antagonist protocol (43.3% vs. 35%



4881 and 60% vs. 36.6% resp.), but not with the short GnRH agonist protocol (43.3% vs. 40%) (Gizzo et al.,
4882 2014).

4883 Two RCTs compared different dosages of oestradiol in addition to progesterone for LPS (Kutlusoy et al.,
4884 2014, Tonguc et al., 2011). Tonguc et al. compared vaginal progesterone with 3 different dosages of
4885 oestradiol (2-4-6 mg) in 285 women and found no difference in clinical pregnancy rate between groups
4886 (31.6% (30/95) vs. 40% (38/95) vs. 32% (31/95) resp.) (Tonguc et al., 2011). Kutlusoy et al. compared
4887 vaginal progesterone with 2 mg oestradiol and 6 mg oestradiol in 62 women and found no significant
4888 difference in live birth rate between dosages (37% (10/27) vs. 22.9% (8/35)) (Kutlusoy et al., 2014).

4889 Recommendation

The addition of oestradiol to progesterone for luteal phase support is probably not recommended.

Conditional ⊕⊕○○

4890

4891 Justification

4892 The data suggests that oestradiol is not recommended for LPS, since it does not improve efficacy in
4893 terms of live birth/ongoing pregnancy rate, or safety in terms of OHSS.

4894 18.4 HUMAN CHORIONIC GONADOTROPHIN (HCG)

4895 Evidence

4896 The Cochrane meta-analysis, mentioned before, found a higher live birth/ongoing pregnancy rate with
4897 hCG for LPS compared to placebo/no treatment (3 RCT, OR 1.76, 95% CI 1.08-2.86, 527 women) (van
4898 der Linden et al., 2015). However, the OHSS rate was increased with hCG for LPS (1 RCT, OR 4.28, 95%
4899 CI 1.91-9.60, 387 women) (Belaisch-Allart et al., 1990, van der Linden et al., 2015).

4900 When compared to progesterone, hCG for LPS or supplementation of progesterone with hCG did not
4901 have a beneficial effect on live birth/ongoing pregnancy rate (5 RCT, OR 0.95, 95% CI 0.65-1.38, 833
4902 women). Furthermore, progesterone was associated with lower rates of OHSS rates than hCG with or
4903 without progesterone (5 RCT, OR 0.46, 95% CI 0.30-0.71, 1293 women) (van der Linden et al., 2015).

4904 Two pilot RCTs, one in women experiencing a normal response to ovarian stimulation with low risk of
4905 OHSS (≤ 13 follicles) and the second in women experiencing a normal response at risk of OHSS (14-25
4906 follicles). In both pilot studies, the study group received GnRH agonist for final oocyte maturation
4907 trigger, combined with two boluses of hCG after oocyte retrieval and on day 4 after oocyte retrieval
4908 ($n=50$ in RCT 1 and $n=46$ in RCT 2). The control group in both pilot studies received hCG for final oocyte
4909 maturation trigger and vaginal progesterone (3x daily) for luteal support ($n=54$ in RCT 1 and $n=52$ in
4910 RCT 2) (Humaidan et al., 2021). In women at low risk of OHSS, no cases of OHSS were reported. When
4911 comparing hCG and progesterone for LPS, there was no significant difference in live birth rate (40%
4912 (20/50 vs. 46% (25/54)) or ongoing pregnancy rate (44% (22/50) vs. 46% (25/54)). In women at risk of
4913 OHSS, two cases of OHSS were reported in the study group, compared to 4 in the control group (not
4914 statistically significant). No significant difference was observed with hCG compared to progesterone for
4915 LPS for live birth rate (51% (25/49) vs. 58% (30/52)), ongoing pregnancy (51% (25/49) vs. 60% (30/52))
4916 or number of MII oocytes retrieved (12.3 ± 4.4 vs. 12.2 ± 4.6).



4917 One small study including 91 women compared hCG with progesterone combined with oestradiol for
4918 LPS and found no difference in clinical pregnancy rate (RR 0.99, 95% CI 0.50-1.92) (Smitz et al., 1988).

4919 Recommendations

In hCG triggered ovarian stimulation cycles, hCG as luteal phase support in standard dosages of 1500 IU is not recommended. [updated]	Strong ⊕⊕○○
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4920

4921 Justification

4922 hCG is equal to progesterone protocols regarding efficacy. However, hCG increased the OHSS risk,
4923 specifically in high responders and with the dosages historically used (1500 IU).

4924 Studies comparing hCG and progesterone for luteal support have not been stratified according to
4925 ovarian response.

4926 18.5 GNRH AGONIST

4927 18.5.1 SINGLE GNRH AGONIST BOLUS SUPPLEMENTATION

4928 Evidence

4929 Most of the studies administered a single bolus of GnRH agonist for LPS on day 6 after oocyte pick-up
4930 at a dose of 0.1 mg for triptorelin 1 mg for leuprolide.

4931 A systematic review²⁷ and meta-analysis compared the use of a bolus GnRH agonist to the control LPS
4932 protocol (Liu et al., 2022). No significant difference was found between a single-dose GnRH agonist and
4933 control for LPS for live birth rate (OR 1.29, 95% CI 0.90-1.84, 6 RCT, 644 participants).

4934 Recommendation

A GnRH agonist bolus, in addition to progesterone for luteal phase support in hCG triggered cycles is probably not recommended.	Conditional ⊕⊕○○
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4935

4936 Justification

4937 The use of GnRH agonist for LPS needs further evaluation in well-designed RCTs, available studies in the
4938 meta-analysis have been rated as of very low quality. Current evidence indicates no significant
4939 difference in live birth/pregnancy rates with GnRH agonist bolus in addition to progesterone for LPS. It
4940 does not seem to increase the risk of OHSS (Yildiz et al., 2014).

²⁷ The meta-analysis by Van der Linden et al., 2015 cited here in the previous version of the guideline was replaced by an updated meta-analysis. The RCTs by Razieh et al., 2009 and Zafardoust et al., 2015 are included in the new meta-analysis and therefore no longer mentioned separately.



4941 Long-term health effects in the new-born have not been studied.

4942 18.5.2 REPEATED GnRH AGONIST

4943 Evidence

4944 Most of the studies administered GnRH agonist for LPS at dosages of 0.1 mg for triptorelin 1 mg for
4945 leuprolide.

4946 The Cochrane meta-analysis reported that multiple doses GnRH agonist added to progesterone for LPS
4947 significantly increased live birth/ongoing pregnancy rate compared to progesterone alone (5 RCT, OR
4948 0.64, 95% CI 0.42-0.98, 1325 women) (van der Linden et al., 2015). One RCT in the meta-analysis
4949 reported OHSS and showed no difference between the groups (OR 1.00, 95% CI 0.33-3.01, 300 women)
4950 (van der Linden et al., 2015, Yildiz et al., 2014).

4951 Recommendation

Repeated GnRH agonist injections, alone or in addition to progesterone for luteal phase support in hCG triggered cycles is probably not recommended. [reworded]

Conditional ⊕○○○

4952

4953 Justification

4954 Current evidence indicates higher live birth /pregnancy rates with GnRH agonist alone or in addition to
4955 progesterone for LPS. The evidence on safety of GnRH agonist for LPS is very limited (1 RCT), however,
4956 it does not seem to increase the risk of OHSS (Yildiz et al., 2014). The evidence on GnRH agonist for LPS
4957 in GnRH antagonist cycles is also limited.

4958 Long-term health effects in the new-born have not been studied. Until these data are available, the
4959 GDG recommends against using GnRH agonist for LPS.

4960 18.6 LH SUPPLEMENTATION

4961 Evidence

4962 One small RCT including 35 women reported no difference in live birth rate (22.2% (4/18) vs. 23.5%
4963 (4/17)) or number of oocytes retrieved (11.7±1.9 vs. 13.8±1.8) between the LH supplementation
4964 group and the progesterone alone group. No cases of OHSS were reported in either group
4965 (Papanikolaou et al., 2011).

4966 Recommendation

Addition of LH to progesterone for luteal phase support can only be used in the context of a clinical trial.

Research
only

4967



4968 **Justification**

4969 The available evidence consists of 1 very small pilot study, which has investigated the effect of adding
4970 LH to progesterone for LPS. However, the study and control group received different triggers for final
4971 oocyte maturation (rhCG compared to GnRH agonist). Therefore, no conclusions can be drawn on the
4972 effect of LH supplementation for LPS, and this intervention cannot be recommended.

4973
4974

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- 5090

PART G: Prevention of OHSS

5091

5092

5093 In previous sections, recommendations were formulated regarding the preferable protocol of ovarian
5094 stimulation for predicted high responders. In short, evidence indicates that GnRH antagonist protocol
5095 is as effective as the GnRH agonist protocol, and significantly reduces the risk of OHSS in PCOS women.
5096 Even though there is no specific evidence on predicted non-PCOS high responders or PCOM patients,
5097 consensus of the guideline group is that GnRH antagonist protocol should also be recommended in
5098 these patient groups (section 4A, page 47). Furthermore, evidence from one RCT indicated that in case
5099 an GnRH agonist protocol is used in high responders, a reduced gonadotropin dose may decrease the
5100 risk of OHSS. Progestin protocol stimulation allows the use of a GnRH agonist trigger and avoids a fresh
5101 embryo transfer. Given similar effectiveness to GnRH analogues for pituitary suppression progestin
5102 protocol can be considered a patient friendly and cost effective option for planned freeze all cycles in
5103 patients with an anticipated high response and risk of OHSS.

A reduced gonadotropin dose is probably recommended to decrease the risk of OHSS in predicted high responders. [2025]

Conditional ⊕○○○

5104

The GnRH antagonist protocol is recommended for predicted high responders. However, if GnRH agonist protocols are used, a reduced gonadotropin dose is recommended to decrease the risk of OHSS. [updated]

Strong ⊕○○○

5105

If freeze-all is planned, the use of progestin for pituitary suppression is probably equally recommended to GnRH analogues. [updated]

Conditional ⊕○○○

5106

PICO QUESTION: WHICH GnRH AGONIST MEDICATION AS A METHOD OF TRIGGERING WILL ADD TO THE PREVENTION OF THE OVARIAN HYPERSTIMULATION SYNDROME ALSO WITH REGARDS TO OVERALL EFFICACY?

5108 **GnRH AGONIST TRIGGER VS hCG TRIGGER IN (PREDICTED) HIGH RESPONDERS**

5109 **Evidence**

5110 *GnRH agonist vs hCG 10.000 IU trigger and fresh transfer*

5111 A Cochrane meta-analysis comparing GnRH agonist trigger with hCG trigger found that GnRH agonist
5112 trigger was associated with a significantly lower risk of moderate/severe OHSS when compared with
5113 hCG among women at high risk of OHSS (3 RCT, OR 0.09, 95%CI 0.02-0.52, 212 women) (Youssef et al.,
5114 2014).

5115 Due to technical limitations of the meta-analysis, all other outcomes were collected from individual
5116 studies. In an RCT including 28 PCO women, comparing GnRH agonist with hCG for final oocyte
5117 maturation, no significant difference was found for live birth rate (1/15 vs. 2/13) or number of oocytes
5118 retrieved (19.8 ± 2.5 vs. 19.5 ± 1.9) (Babayof et al., 2006). Similarly, in an RCT including 66 women with
5119 PCOS or previous high response, no significant difference was found in ongoing pregnancy rate (53.3%
5120 (16/30) vs. 48.3% (14/29)) or number of oocytes retrieved (20.2 ± 9.9 vs. 18.8 ± 10.4) between GnRH
5121 agonist and hCG for final oocyte maturation (Engmann et al., 2008). It is noted that the latter trial
5122 employed augmented luteal phase support protocols with additional oestrogen with intramuscular
5123 progesterone in the GnRH agonist triggered arm.

5124 *GnRH agonist trigger with fresh transfer vs freeze-all*

5125 An RCT including 212 women at risk of OHSS (>17 follicles of >11 mm on the day of trigger) compared
5126 GnRH agonist trigger in GnRH antagonist protocol with or without a freeze all (Santos-Ribeiro et al.,
5127 2020). While live birth rates were similar (39.4% (41/104) vs. 41.6% (42/101)), moderate-to-severe
5128 OHSS occurred only in the fresh transfer group that was given an additional single low-dose hCG on
5129 the day of the trigger (8.6% (9/105), 95% CI 3.2-13.9% vs. 0% (0/104), 95% CI 0-3.7%) (Santos-Ribeiro
5130 et al., 2020).

5131 An RCT including 280 women at risk of OHSS (number of follicles ≥ 12 mm between 14 and 25 on the
5132 day of trigger) compared GnRH agonist trigger with or without freeze-all (Aflatoonian et al., 2018).
5133 There was no significant difference in live birth rate (27.3% (33/121) vs. 26.9% (32/119); OR 1.02,
5134 0.57-1.80) or moderate OHSS (5.8% (7/121) vs. 5.9% (7/119)) between GnRH agonist trigger with
5135 freeze-all or fresh transfer. No cases of severe OHSS were reported in either group (Aflatoonian et al.,
5136 2018).

5137 *GnRH agonist vs hCG non-10.000 IU trigger and fresh transfer*

5138 One RCT including 118 patients at risk of OHSS (between 14 and 25 follicles ≥ 11 mm diameter on trigger
5139 day) reported no difference in OHSS between GnRH agonist trigger (0% (0/60)) compared to reduced
5140 hCG dose (3.4% (2/58)) in a GnRH antagonist protocol. No severe OHSS was reported in either group.
5141 Ongoing pregnancy rates were similar for GnRH agonist trigger (28.3% (17/60)) compared to reduced-
5142 dose hCG trigger (25.9% (15/58)) and also a similar number of oocytes was retrieved in both groups



5143 (13.7±5.9 vs. 13.5±5.7) (Humaidan et al., 2013). It is noted that augmented luteal phase support
5144 protocols with additional doses of hCG were employed in the GnRH agonist triggered arm.

5145 Recommendation

A GnRH agonist trigger is recommended for final oocyte maturation in women at risk of OHSS combined with a freeze-all strategy to minimise the risk of severe OHSS. [updated]

Strong ⊕○○○

5146

If a GnRH agonist protocol with hCG trigger is used in high responders, a freeze-all strategy is recommended to decrease the risk of late-onset OHSS. [updated]

GPP

5147

5148 Justification

5149 Triggering final oocyte maturation with GnRH agonist significantly reduces the risk of early-onset OHSS
5150 in patients at risk of OHSS.

5151 Limited evidence suggests that GnRH agonist trigger with fresh transfer is as efficient and safe as GnRH
5152 agonist trigger with freeze-all in patients at risk of OHSS with number of follicles ≥12 mm between 14
5153 and 25 on the day of trigger. Modified luteal support with LH-activity (hCG or LH) may overcome the
5154 reduction in clinical pregnancy rate after GnRH agonist trigger. However, its effectiveness of OHSS
5155 prevention is reduced.

5156 DUAL TRIGGER

5157 Evidence

5158 In a retrospective cohort study, dual trigger was compared to GnRH agonist for final oocyte maturation
5159 in PCOS patients undergoing ovarian stimulation for IVF/ICSI with freeze-all (Wang et al., 2024). No
5160 significant difference in live birth rate was observed when comparing dual trigger to GnRH agonist only
5161 for final oocyte maturation (56.2% (99/176) vs. 63.1% (111/176)). However, the total OHSS rate (14.8%
5162 (26/176) vs. 2.8% (5/176)) and the moderate/severe OHSS rate (11.4% (20/176) vs. 1.7% (3/176)) were
5163 significantly higher after dual trigger compared to GnRH agonist only.

5164 In a retrospective cohort study, dual trigger with 1000 IU (n=403) or 2000 IU hCG (n=363) was compared
5165 to GnRH agonist trigger only (n=577) in high responders to ovarian stimulation having freeze-all (He et
5166 al., 2022). Comparing GnRH agonist only to both groups of dual trigger (1000 IU and 2000 IU hCG,
5167 respectively), there was no significant difference for cumulative live birth rate (74.4% (429/577) vs.
5168 75.7% (305/403) vs. 69.7% (253/363)) or live birth rate (54.2% (302/577) vs. 54.5% (212/389) vs. 54.3%
5169 (191/352)). However, moderate to severe OHSS rate was significantly higher with dual trigger (1000 IU
5170 and 2000 IU hCG, respectively) compared to GnRH agonist trigger alone (1.5% (6/403) vs. 1.4% (5/363)
5171 vs. 0%).



5172 Recommendation

The addition of hCG to GnRH agonist as a dual trigger for final oocyte maturation is probably not recommended for high responders. [2025]

Conditional ⊕○○○

5173

5174 Justification

5175 The supporting evidence comes from retrospective cohort studies. No difference in efficacy was
5176 observed with dual trigger compared to GnRH agonist trigger. However, both studies reported
5177 significantly more cases of OHSS in the dual trigger group. Because of these safety concerns, adding
5178 hCG to GnRH agonist as dual trigger cannot be recommended in high responders.

5179 **GNRH AGONIST TRIGGER + FREEZE-ALL VS HCG TRIGGER+FREEZE-ALL IN (PREDICTED) HIGH RESPONDERS**

5180 Evidence

5181 A case-control study, including 248 women at risk of OHSS, compared GnRH agonist trigger and freeze-
5182 all to hCG trigger and freeze-all. There was no significant difference in cumulative pregnancy rate
5183 between GnRH agonist and hCG trigger with freeze-all (59.5% vs. 53.0%) (Borges et al., 2016).

5184 Similar results were found in a retrospective cohort study including 272 women at risk of OHSS, also
5185 comparing hCG trigger and freeze-all with GnRH agonist trigger and freeze-all. There was no difference
5186 in cumulative live birth rate between GnRH agonist and hCG for final oocyte maturation and freeze-all
5187 (48.15% vs. 48.08%) (Tannus et al., 2017).

5188 Recommendation

In patients at risk of OHSS, the use of a GnRH agonist for final oocyte maturation is probably recommended over hCG in cases where no fresh transfer is performed. [2019]

Conditional ⊕○○○

5189

5190 Justification

5191 Available evidence is derived from low-quality studies in patients at risk of OHSS. However, evidence
5192 from RCTs performed in oocyte donors indicates that GnRH agonist trigger is preferable over hCG when
5193 a freeze-all strategy is applied (Acevedo et al., 2006, Galindo et al., 2009, Melo et al., 2009, Sismanoglu
5194 et al., 2009). The guideline group thinks that the data can be extrapolated to GnRH agonist trigger
5195 compared to hCG with freeze-all in both arms for patients at risk of OHSS.

5196 **GNRH AGONIST TRIGGER VS COASTING+HCG TRIGGER IN (PREDICTED) HIGH RESPONDERS**

5197 Evidence

5198 A retrospective study including 94 women at risk of OHSS reported that 10/33 women in the coasting
5199 group had cycle cancellation because of the risk of development of OHSS vs. 0/61 in the GnRH agonist
5200 trigger group. No cases of OHSS occurred in either treatment group. Ongoing pregnancy rates (49.2%



5201 (30/61) vs. 24.2% (8/33)) and number of oocytes retrieved (26.9±9.5 vs. 17.7±9.3) were significantly
5202 higher in the GnRH agonist trigger group compared to the coasting group (DiLuigi et al., 2010).

5203 Another retrospective study including 248 women at risk of OHSS reported more cancelled cycles in
5204 the coasting group compared to the GnRH agonist trigger with freeze-all group (19.7% (30/152) vs.
5205 8.3% (8/96) because of poor embryo quality or risk of OHSS. The clinical pregnancy rate in the coasting
5206 group was 29.5% (36/122), which was significantly lower than the GnRH agonist trigger with freeze-all
5207 (50% (44/88)) (Herrero et al., 2011).

5208 Recommendation

A GnRH agonist trigger for final oocyte maturation with or without a freeze-all strategy is preferred over a coasting strategy in patients at risk of OHSS. [2019]

GPP

5209

5210 Justification

5211 The two most relevant studies were both on retrospective data, with inherent methodological and
5212 risk of bias problems. Therefore, the GDG cannot recommend coasting and hCG trigger over GnRH
5213 agonist trigger for final oocyte maturation in patients at risk of OHSS.

5214 DOPAMINE AGONISTS

5215 Evidence

5216 A systematic review and meta-analysis comparing a dopamine agonist to no intervention or placebo
5217 included 10 RCTs with 1202 participants and reported significantly lower risk of moderate or severe
5218 OHSS with the use of dopamine agonists (OR 0.32, 95% CI 0.23-0.44). Live birth rates were reported in
5219 only 3 RCTs, including 362 participants, and were similar in the two groups (OR 0.96, 95% CI 0.60-
5220 1.55) (Tang et al., 2021).

5221 A retrospective study, including 480 patients at risk of OHSS, compared GnRH agonist trigger alone,
5222 GnRH agonist trigger and a dopamine agonist from the day of trigger or oocyte retrieval for seven
5223 days, and GnRH agonist with dopamine agonist as described above in combination with daily GnRH
5224 antagonist for five days from oocyte retrieval day (Shrem et al., 2019). All embryos were frozen in the
5225 three groups. None of the patients developed severe OHSS, however, the incidence of mild or
5226 moderate OHSS was significantly higher in the GnRH agonist trigger only group than in the GnRH
5227 agonist trigger and dopamine agonist group (38% vs. 29%) and the GnRH agonist trigger, dopamine
5228 agonist and GnRH antagonist group (38% vs. 18%). The GnRH agonist trigger and dopamine agonist
5229 groups had a significantly higher risk of mild or moderate OHSS than the GnRH agonist trigger in
5230 combination with dopamine agonist and GnRH antagonist (29% vs. 18%).

5231



5232 Recommendation

Dopamine agonists are recommended to decrease the risk of early OHSS, particularly in patients receiving hCG for final oocyte maturation. [2025]	Strong ⊕⊕○○
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5233

5234 Justification

5235 The GDG recommends using GnRH agonist trigger combined with freeze-all for women at risk of
5236 OHSS. However, if the patient is deemed at risk of OHSS after an hCG trigger, dopamine agonist can
5237 be used as a preventive measure for early OHSS. Dopamine agonists inhibit endothelial VEGF
5238 receptors and decrease vascular permeability. However, rapid luteolysis with a GnRH agonist trigger
5239 combined with a freeze all strategy may render the addition of dopamine agonists obsolete or
5240 marginally effective with regard to clinically relevant OHSS in cycles with GnRH antagonist pituitary
5241 suppression.

5242

PICO QUESTION: IS THE FREEZE-ALL PROTOCOL MEANINGFUL IN THE PREVENTION OF OVARIAN HYPER-STIMULATION SYNDROME ALSO WITH REGARD TO EFFICACY?

5243 Ovarian hyperstimulation syndrome (OHSS) is a potential life-threatening condition. It implies
5244 hospitalization frequently, with health care additional costs and patient burden. However, it may be
5245 balanced to the possible negative effects of a freeze-all policy and the decline in live birth rates, due to
5246 eliminating the fresh transfer from the treatment scheme.

5247 Evidence

5248 A Cochrane systematic review and meta-analysis comparing freeze-all to conventional ovarian
5249 stimulation with fresh transfer reported a significantly lower incidence of OHSS (0.8% vs. 3.7% (Peto OR
5250 0.26, 95% CI 0.17-0.39; 6 RCTs, 4478 women)) with the freeze-all strategy compared to fresh transfer.
5251 Furthermore, they found no difference in cumulative live birth rate and pooled for all embryo stages at
5252 transfer (OR 1.08, 95% CI 0.95-1.22; 8 RCTs, 4712 women) (Zaat et al., 2021).

5253 Recommendation

A freeze-all strategy is recommended to minimise the risk of late-onset OHSS. [updated]	Strong ⊕⊕○○
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5254

Prior to start of ovarian stimulation, a risk assessment for high response is advised with the purpose of applying personalised treatment choices on pituitary suppression protocol, FSH dosage, final oocyte maturation trigger and embryo transfer strategy. [updated]	GPP
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5255 Justification

5256 The current evidence suggests that not performing a fresh embryo transfer lowers the OHSS risk for
5257 women at risk of OHSS, without completely eliminating the condition. The latter urges for follow up of
5258 haemo-concentration status even in cases with the freeze-all strategy applied.

5259 The conditions with a high prior risk of developing the OHSS comprise:

- 5260 • patients with the PCOS syndrome,
- 5261 • patients with an above average ovarian reserve status
- 5262 • patients exhibiting a high ovarian response as indicated by follicle number at ultrasound, high
5263 oestradiol levels, or high number of oocytes obtained

5264 Applying the freeze-all strategy implies the presence of a high-quality cryopreservation program.

5265

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- 5333

Glossary

5334

5335

Ovarian hyperstimulation syndrome (OHSS)	An exaggerated systemic response to ovarian stimulation characterized by a wide spectrum of clinical and laboratory manifestations. It may be classified as mild, moderate or severe according to the degree of abdominal distention, ovarian enlargement and respiratory, hemodynamic and metabolic complications.
Ovarian stimulation (OS)	Pharmacological treatment with the intention of inducing the development of ovarian follicles. It can be used for two purposes: 1) for timed intercourse or insemination; 2) in ART, to obtain multiple oocytes at follicular aspiration.
Low ovarian responder in assisted reproductive technology	A woman treated with ovarian stimulation for ART, in which at least two of the following features are present: (1) Advanced maternal age (≥ 40 years); (2) A previous low ovarian response (≤ 3 oocytes with a conventional stimulation protocol aimed at obtaining more than three oocytes); and, (3) An abnormal ovarian reserve test (i.e. antral follicle count 5–7 follicles or anti-Mullerian hormone 0.5–1.1 ng/ml (Bologna criteria); or other reference values obtained from a standardized reference population.)
Low ovarian response to ovarian stimulation	A condition in which fewer than four follicles and/or oocytes are developed/obtained following ovarian stimulation with the intention of obtaining more follicles and oocytes.
Mild ovarian stimulation	A protocol in which the ovaries are stimulated with gonadotropins, and/or other pharmacological compounds, with the intention of limiting the number of oocytes following stimulation for IVF.
Modified natural cycle	A procedure in which one or more oocytes are collected from the ovaries during a spontaneous menstrual cycle. Pharmacological compounds are administered with the sole purpose of blocking the spontaneous LH surge and/or inducing final oocyte maturation

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5342

Annexes

5343

5344

5345 [Annex 1: Guideline development group](#)

5346 [Annex 2: Abbreviations](#)

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5348 [Annex 4: Methodology](#)

5349 [Annex 5: Stakeholder consultation](#)

5350 [Annex 6: Summary of findings tables](#)

5351 [Annex 7: Literature study: flowcharts, list of excluded studies](#)

5352 [Annex 8: Evidence tables](#)

5353

Draft for review



5354

Annex 1: Guideline development group

5355 This guideline was developed by the ESHRE Reproductive Endocrinology Guideline Development
5356 Group (GDG). The GDG included gynaecologists with expertise in reproductive medicine and ovarian
5357 stimulation. We aimed for an equal distribution in gender, region and expertise.

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Methodological support

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5358 **DECLARATIONS OF INTEREST**

5359

5360 All members of the guideline development group were asked to declare possible conflicts of interest
5361 by means of the disclosure forms (see *ESHRE Manual for Guideline Development*).

Conflicts of interest	
Frank Broekmans	Research grants from Merck, Ferring, Besins Consulting fees from Ferring, Merck, Besins, Abbott Speaker's fees from Merck, Besins, Ferring
Nikolaos P. Polyzos	Research grants from Besins, Ferring, Merck, Organon, Roche Diagnostics and Theramex Consulting fees from Besins, Ferring, IBSA, Merck, Organon, and Abbott Speaker's fees from Besins, Ferring, Gedeon-Richter, IBSA, Merck, Organon and Theramex
Antonio La Marca	Research grants from Merck, Ferring, IBSA, Roche, Organon, Theramex, Beckman Coulter and Gedeon-Richter Consulting fees from Merck, Ferring, IBSA, Roche, Organon, Theramex, Beckman Coulter and Gedeon-Richter Speaker's fees from Merck, Ferring, IBSA, Roche, Organon, Theramex, Beckman Coulter and Gedeon-Richter
Georg Griesinger	Consulting fees from Organon, Ferring, Merck, Gedeon-Richter, Theramex, Abbott, ReproNovo, Igxos, OxoLife, Philipps, ReprodWissen, PregLem, Guerbet, Roche, IBSA, and Besins. Speaker's fees from Organon, Ferring, Merck, Gedeon-Richter, Theramex, Abbott, ReproNovo, Igxos, OxoLife, Philipps, ReprodWissen, PregLem, Guerbet, Roche, IBSA, and Besins. Research grants from Besin, Merck, Abbott, Ferring, Theramex.
Ernesto Bosch	Research grants from Gedeon-Richter Consulting fees from MSD, Ferring, Abbot, Gedeon-Richter, Merck, Roche Speaker's fees from MSD, Ferring, Abbot, Gedeon-Richter, Merck, Roche Ownership interest from IVI-RMS Valencia
Baris Ata	Speaker's fees from Gedeon-Richter, Ferring, IBSA, Intas, Merck, Organon. Consulting fees from Merck, Organon, Oxolife.
Janos Urbancsek	None declared.
Nathalie Massin	Research grants from IBSA, Organon Consulting fees from Organon, Merck, GE, Ferring Speaker's fees from Merck, Gedeon-Richter, Theramex
Töyli Mira	None declared.
Michael Grynberg	Speaker's fees from Merck Serono, Ferring, Gedeon Richter
Sesh Kamal Sunkara	Research grant from Ferring Consulting fees from Merck Speaker's fees from Merck and Ferring
Simone Broer	None declared.
George Lainas	Consulting and speaker's fees from Organon, Ferring, Merck, Gedeon- Richter, Cook, Vianex.
Estratios Kolibianakis	Travel/hotel expenses from Ferring, SERONO, Vianex



	Chair of the Greek Society of Fertility and Sterility
Michal Kunicki	Speaker's fees from Ferring
Tanya Timeva	Speaker's fees from Merck, Organon, MSD
Nathalie Le Clef	None declared.

5362

Draft for review



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Annex 2: Abbreviations

5364

AFC	Antral follicle count
AMH	Anti-Müllerian hormone
ART	Assisted reproductive technology
BMI	Body mass index
CC	Clomiphene citrate
CI	Confidence interval
COC	Cumulus-oocyte complex
COCP	Combined oral contraceptive pill
DHEA	Dehydroepiandrosterone
Duostim	Double stimulation, ovarian stimulation during the follicular and luteal phase of the same cycle
EFORT	Exogenous follicle stimulating hormone ovarian reserve test
EMT	Endometrial thickness
FSH	Follicle stimulating hormone
GDG	Guideline development group
GH	Growth hormone
GnRH	Gonadotropin-releasing hormone
GPP	Good practice point
hCG	Human chorionic gonadotrophin
hMG	Human menopausal gonadotropin
hp-FSH	Highly purified follicle stimulating hormone
ICSI	Intracytoplasmic sperm injection
IPD	Individual patient data
IU	International unit
IUI	Intra-uterine insemination
IVF	In vitro fertilization
LBR	Live birth rate
LH	Luteinizing hormone
LPS	Luteal phase support
LR	Logistic regression
MD	Mean difference
MNC	Modified natural cycle
MPA	Medroxy progesterone acetate
OHSS	Ovarian hyperstimulation syndrome
OPU	Oocyte pick-up
OR	Odds ratio
OS	Ovarian stimulation
PCOM	Polycystic ovary morphology
PCOS	Polycystic ovary syndrome
p-FSH	Purified follicle stimulating hormone
POI	Premature ovarian insufficiency
PR	Pregnancy rate
RCT	Randomized controlled trial
rFSH	Recombinant follicle stimulating hormone
rLH	Recombinant luteinizing hormone
ROC-AUC	Receiver operating characteristic – area under the curve
RR	Relative risk/risk ratio
SMD	Standardized mean difference
WMD	Weighted mean difference



5365 Annex 3: Recommendations for 5366 research in OS for IVF/ICSI

5367 From the literature and discussion of the available evidence, several topics were identified for which
5368 evidence is inconsistent, insufficient or non-existing. For the benefit of couples with RPL, the GDG
5369 recommends that future research, where possible in well-designed RCTs, should focus on these
5370 research gaps.

5371 Considered are:

- 5372 • Gonadotropin dose reduction in predicted high responders as a tool for normalization of
5373 ovarian response (GnRH agonist or antagonist) compared to a standard dosage with option
5374 GnRH agonist trigger and/or a freeze-all strategy (in GnRH antagonist protocol).
- 5375 • The effect on live birth rates of deferring embryo transfer in situations with elevated
5376 Progesterone on the day of the trigger, compared to standard scheduling the fresh transfer in
5377 day 5 transfer programmes.
- 5378 • Changing from rFSH stimulation to hMG stimulation or vice versa in cases with a high rate of
5379 immature oocytes (M1 and/or GV) after a standard stimulation phase and 10.000 IU hCG
5380 trigger: will it affect the immature oocyte rate and live birth rate?
- 5381 • Comparing the use of the PPOS scheme in predicted high responders to the use of a standard
5382 antagonist stimulation scheme, with respect to live birth, safety for the female and safety for
5383 the offspring
- 5384 • The effect of applying a FSH dose adaptation on day 5-6 of the stimulation versus continuing
5385 the same FSH dose from the start, provided that the FSH dose has been chosen based on prior
5386 identification of the predicted ovarian response, on FSH consumption and live birth prospects.

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Annex 4: Methodology

5389 GUIDELINE DEVELOPMENT

5390 European Society of Human Reproduction and Embryology (ESHRE) guidelines are developed based on
5391 the Manual for ESHRE guideline development (Vermeulen et al., 2017), which can be consulted at the
5392 ESHRE website (www.eshre.eu/guidelines). The principal aim of this manual is to provide stepwise
5393 advice on ESHRE guideline development for members of ESHRE guideline development groups. The
5394 manual describes a 12-step procedure for writing clinical management guidelines by the guideline
5395 development group, supported by the ESHRE methodological expert:

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|----------------------|------------------------|
| 1 TOPIC SELECTION | 7 RECOMMENDATIONS |
| 2 GDG FORMATION | 8 DRAFT FOR REVIEW |
| 3 SCOPING | 9 STAKEHOLDER REVIEW |
| 4 KEY QUESTIONS | 10 EXCO APPROVAL |
| 5 EVIDENCE SEARCH | 11 PUBLICATION |
| 6 EVIDENCE SYNTHESIS | 12 UPDATING / REVISING |

5397 The two versions of this guideline (2019 and 2025) were developed and funded by ESHRE, which
5398 covered expenses associated with the guideline meetings (travel, hotel and catering expenses)
5399 associated with the literature searches (library costs, costs associated with the retrieval of papers) and
5400 with the implementation of the guideline (printing, publication costs). Except for reimbursement of
5401 their travel expenses, GDG members did not receive any payment for their participation in the guideline
5402 development process.

5403 For the 2019 version of the guideline, the scope of the guideline and first version of the key questions
5404 were drafted by the coordinator and deputies of the ESHRE Special Interest Group Reproductive
5405 Endocrinology. A call was launched for experts in the field interested in joining the guideline
5406 development group. All applications were reviewed, and experts were selected based on expertise and
5407 geographical location. We strived towards a balance in gender and location within Europe. A meeting
5408 of the guideline development group was organized to discuss the key questions and redefine them
5409 through the PICO process (patients – interventions – comparison – outcome). This resulted in a final list
5410 of 18 key questions. Based on the defined key words, literature searches were performed by the
5411 methodological expert (Dr. N. Le Clef). Key words were sorted to importance and used for searches in
5412 PUBMED/MEDLINE and the Cochrane library. We searched the databases from inception up to 8
5413 November 2018. For the 2025 update of the guideline, all guideline group members of the 2019 were
5414 contacted to be part of the guideline development group, one member declined and was replaced. The
5415 key questions of the 2019 version were reviewed and refined, and new interventions were added were
5416 relevant. An update of the literature searches was performed by the methodological expert (Dr. N. Le
5417 Clef). We searched the databases for literature published between 1 November 2018 and 2 February
5418 2025.

5419 Literature searches were performed as an iterative process. In a first step, systematic reviews and meta-
5420 analyses were collected. If no results were found, the search was extended to randomized controlled



5421 trials, and further to cohort studies and case reports, following the hierarchy of the levels of evidence.
 5422 Reference were selected or excluded by the methodological expert and expert GDG member based on
 5423 title and abstract and knowledge of the existing literature. If necessary, additional searches were
 5424 performed in order to get the final list of papers. For interventional questions, focus was on prospective
 5425 (randomized) controlled studies. . It is not within ESHRE's remit to conduct a formal investigation or to
 5426 draw formal conclusions regarding the misconduct of an individual or group of individuals or to
 5427 determine whether a published article should be retracted. However, papers that are withdrawn, have
 5428 a published editorial note of concern or a published expression of concern have been excluded from
 5429 the guideline. In future revision or update of the guideline, the GDG will actively verify the status of all
 5430 the referenced studies.

5431 The quality of the selected papers was assessed by means of the quality assessment checklist, defined
 5432 in the ESHRE guideline manual. Furthermore, the evidence was collected and summarized in an
 5433 evidence table according to GIN format (<http://www.g-i-n.net/activities/etwg>). The quality assessment
 5434 and evidence tables were constructed by the expert GDG members.

5435 Summary of findings tables (Annex 6) were prepared following the GRADE approach for randomized
 5436 controlled intervention studies which reported the critical outcomes, i.e. cumulative live birth rate, live
 5437 birth rate and OHSS rate. Where available, summary of findings tables were based on existing up-to-
 5438 date well-executed systematic reviews, if necessary supplemented with additional recent RCTs. When
 5439 there was no recent valid systematic review available, we systematically searched for relevant studies,
 5440 as described above, with focus on prospective (randomized) studies.

5441 GDG meetings were organized to discuss the draft recommendations and the supporting evidence and
 5442 to reach consensus on the final formulation of the recommendations. In a final step, all evidence and
 5443 recommendations were combined in the ESHRE guideline: “Ovarian stimulation for IVF/ICSI”.

5444 FORMULATION OF RECOMMENDATIONS

5445 We labelled the recommendations as either “strong” or “conditional” according to the GRADE
 5446 approach. We used the words “we recommend” for strong recommendations and “we probably
 5447 recommend” for conditional recommendations. Suggested interpretation of strong and conditional
 5448 recommendations by patients, clinicians and health care policy makers is as follows:

Implications for	Strong recommendation	Conditional recommendation
Patients	Most individuals in this situation would want the recommended course of action, and only a small proportion would not	The majority of individuals in this situation would want the suggested course of action, but many would not
Clinicians	Most individuals should receive the intervention Adherence to this recommendation according to the guideline could be used as a quality criterion or performance indicator Formal decision aids are not likely to be needed to help individuals make decisions consistent with their values and preferences	Recognise that different choices will be appropriate for individual patients and that you must help each patient arrive at a management decision consistent with his or her values and preferences Decision aids may be useful in helping individuals to make decisions consistent with their values and preferences
Policy makers	The recommendation can be adopted as policy in most situations	Policy making will require substantial debate and involvement of various stakeholders



5449

5450 For each recommendation it is mentioned whether it is strong or conditional and what the quality of
5451 the supporting evidence was. In the justification section, more data are provided on the considerations
5452 taken into account when formulating the recommendations: balance between desirable and
5453 undesirable effects, certainty of the evidence of effects, certainty in how people value the outcome,
5454 acceptability and feasibility of the intervention. Impact on health equity and resource impact were only
5455 discussed where relevant.

5456

Draft for review



5457 STRATEGY FOR REVIEW OF THE GUIDELINE DRAFT

5458 After finalization of the guideline draft, the review process was initiated. The draft guideline was
5459 published on the ESHRE website, accompanied by the reviewers' comments form and a short
5460 explanation of the review process. The guideline was open for review between 6 May and 16 June 2025.

5461 To notify interested clinicians, we sent out an invitation to review the guideline by email to all members
5462 of ESHRE.

5463 Selected reviewers were invited personally by email. These reviewers included:

- 5464 • *Coordinators and deputies of the ESHRE SIG Reproductive Endocrinology and the ESHRE SIG*
5465 *Reproductive Endocrinology and the ESHRE SIG Quality and Safety in ART.*
- 5466 • *Contact persons of patient organizations across Europe.*
- 5467 • *Contact persons of international and national societies focused on IVF/ICSI across Europe.*

5468 All reviewers are listed in Annex 5. The Reviewer comments processing report, including further
5469 information on the review and a list of all comments per reviewer with the response formulated by the
5470 GDG will be published on the ESHRE website.

5471 GUIDELINE IMPLEMENTATION STRATEGY

5472 The standard dissemination procedure for all ESHRE guidelines comprises publishing and
5473 announcement.

5474 Each guideline is published on the ESHRE Website and in Human Reproduction Open. The
5475 announcement procedure includes a news item in "Focus on Reproduction", a newsflash on the ESHRE
5476 website homepage and a short presentation at the ESHRE Annual meeting. All participants in the annual
5477 ESHRE meeting will be informed about the development and release of new guidelines; all related
5478 national societies and patient organizations are informed about the guideline release. They are asked
5479 to encourage local implementation by, for instance, translations or condensed versions, but they are
5480 also offered a website link to the original document.

5481 Patient versions of the guideline will be developed by a subgroup of the GDG together with patient
5482 representatives. The patient version is a translation of the recommendations in everyday language, with
5483 emphasis on questions important to patients. It aims to help patients understand the guideline's
5484 recommendations and facilitates clinical decision-making.

5485 To further enhance implementation of the guideline, the members of the GDG, as experts in the field,
5486 will be asked to select recommendations for which they believe implementation will be difficult and
5487 make suggestions for tailor-made implementation interventions (e.g. option grids, flow-charts,
5488 additional recommendations, addition of graphic/visual material to the guideline).

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5490 **SCHEDULE FOR UPDATING THE GUIDELINE**

5491 The current guideline will be considered for revision in 2029 (four years after publication). An
5492 intermediate search for new evidence will be performed two years after publication, which will inform
5493 the GDG of the necessity of an update.

5494 Every care is taken to ensure that this publication is correct in every detail at the time of publication.
5495 However, in the event of errors or omissions, corrections will be published in the web version of this
5496 document, which is the definitive version at all times. This version can be found at
5497 www.eshre.eu/guidelines.

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5499 **For more details on the methodology of ESHRE guidelines, visit www.eshre.eu/guidelines**

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Draft for review



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Vermeulen N, Le Clef N, D'Angelo A, Tilleman K, Veleva Z, Nelen WL. Manual for ESHRE guideline development. 2017.