

Avoiding OHSS: Controlled Ovarian Low-Dose Stimulation in Women with PCOS

Vermeidung eines OHSS: kontrollierte ovarielle Low-Dose-Stimulation für IVF und ICSI bei Frauen mit polyzystischem Ovarsyndrom

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Abstract



The polycystic ovary syndrome is a common endocrine disorder which influences outcome and potential risks involved with controlled ovarian stimulation for artificial reproductive techniques (ART). Concrete practical recommendations for the dosage of gonadotropins, the preferred protocol and preventive methods to avoid ovarian hyperstimulation syndrome (OHSS) are lacking. We present retrospective data of 235 individually calculated gonadotropin low-dose stimulations for ART in a single center from 2012 to 2014. Clinical data and outcome parameter of patients diagnosed with PCOS according to Rotterdam criteria (n = 39) were compared with patients without PCOS (n = 196). The starting dose of gonadotropins was individually calculated depending on patients' age, BMI, ovarian reserve, ovarian response in previous cycles, and diagnostic criteria of PCOS. Mean age and duration of infertility did not differ between the groups, whereas mean BMI (p = 0.007) and AMH (p < 0.001) were higher in the PCOS-group. A lower mean FSH-starting and maximum dose was administered to women with PCOS (p < 0.001). The biochemical pregnancy rate of 42.4% and the clinical pregnancy rate of 32.2% for PCOS-patients did not differ from those of the control group (42.2% and 34.4% respectively). Neither mild, nor moderate or severe manifestation of OHSS occurred significantly more often in patients with PCOS. Our study supports the use of a calculated low-dose FSH-stimulation strategy in ART for patients with PCOS. Further randomized clinical trials should confirm this strategy and lead to define individual risk factors for OHSS, which can be used for recommendation of safer ART-techniques like in vitro maturation.

Zusammenfassung



Das polyzystische Ovarsyndrom (PCOS) ist eine häufige endokrinologische Störung der Frau, die erhebliche Auswirkungen auf Erfolg und potenzielle Risiken einer Gonadotropinstimulation bei künstlicher Befruchtung hat. Klare praktische Empfehlungen für die FSH-Dosierung und das optimale Stimulationsprotokoll zur Vermeidung eines OHSS fehlen weitgehend. Retrospektiv wurden Zyklen von 235 Patientinnen, die von 2012 bis 2014 für eine IVF/ICSI kontrolliert stimuliert wurden, analysiert. Klinische Daten der PCOS-Patientinnen (n = 39) wurden mit denen ohne PCOS (n = 196) verglichen. Die Gonadotropin-Startdosierung orientierte sich am Alter der Patientinnen, dem BMI, der ovariellen Reserve, Stimulationsverläufen in Vorzyklen und diagnostischen Kriterien des PCOS. Durchschnittliches Alter und Dauer der Infertilität unterschieden sich nicht signifikant zwischen den beiden Gruppen. Der durchschnittliche BMI (p = 0,007) und basale AMH-Wert (p < 0,001) lag bei Frauen mit PCOS höher. Bei ihnen fand sich eine signifikant niedrigere FSH-Start- und Maximaldosis (p < 0,001). Die Gesamtdosis verwendeter Gonadotropineinheiten sowie der maximale Östradiolspiegel jedoch unterschieden sich nicht. Auch die biochemischen und klinischen Schwangerschaftsraten von 42,2 und 32,2% bei PCOS-Patientinnen wichen nicht signifikant von den Ergebnissen der Kontrollgruppe ab (42,2 und 34,4%). Weder leichte noch moderate oder schwere Ausprägungen des OHSS kamen häufiger in einer der beiden Gruppen vor. Unsere Studie schlägt eine kalkulierte, niedrigdosierte Stimulationsstrategie mit Dosissteigerung in kleinen Schritten vor. Durch weitere randomisierte, kontrollierte klinische Studie sollte diese Strategie überprüft werden und individuelle Risikofaktoren erkannt werden, die in Einzelfällen als Indikation zu anderen sicheren Methoden der ART wie der In-vitro-Maturation genutzt werden können.

Introduction

The polycystic ovary syndrome (PCOS) is the most common endocrine disorder of women in reproductive age [1]. It causes chronic oligo- or anovulation and often leads to infertility. Inclusive diagnostic criteria were established 2003 in Rotterdam. The criteria provide opportunities to distinguish four clinical phenotypes using at least two of three criteria to define the syndrome: hyperandrogenism defined either as hyperandrogenaemia or clinically validated hyperandrogenism, oligo- or anovulation, and polycystic ovarian morphology in ultrasound [2]. Geisthövel proposed a further classification of women with PCOS mainly using the diagnostic aspect of hyperandrogenism. He defined five clinical subgroups: functional cutaneous androgenisation (FCA) and four typical manifestations of the female androgenisation syndrome (FAS) I–IV [3]. This approach includes a slightly different group of women than using the Rotterdam criteria for PCOS. Especially women with late onset adrenogenital syndrome or other adrenal origin of hyperandrogenaemia are included in this classification, which are explicitly excluded in the PCOS definition.

Controlled ovarian hyperstimulation (COS) with gonadotropins for artificial reproductive techniques (ART) leads to a higher risk of ovarian hyperstimulation syndrome (OHSS) for patients affected by PCOS, because of a higher sensibility and exaggerated response to gonadotropins [4].

Therefore, it is important to give recommendations for the dosage and the preferred stimulation-protocol to avoid ovarian hyperstimulation syndrome (OHSS). So far, clear guidelines from health institutes and PCOS societies or in scientific literature are lacking.

It is current practice to use the antagonist protocol to avoid OHSS in patients with elevated risk, mainly due to the possibility to use the GnRH-agonist for ovulation induction and cryopreserve all fertilised oocytes or embryos (“freeze all”) [5]. Most published studies use equal doses of gonadotropins for stimulation of patients with or without PCOS. Others propose fixed doses or lack to describe how to individualize dose and regimen. A biographical search of the MEDLINE database was performed in March 2014 and June 2015. The MESH words “PCO” and “IVF”, “controlled ovarian stimulation”, “FSH” were used. Articles prior to the year 2005, those without an abstract or those that were written in a language other than English, German or French were excluded (Table 1).

Clear recommendations for the FSH dosage and protocol should be established to improve outcome and security of controlled ovarian stimulation in a group of ART-patients with special demands.

A modern approach to classify the ovarian response or sensitivity to gonadotropin stimulation in PCOS patients is the evaluation of risks using the levels of AMH [6]. According to this classification, non-responders show basal AMH-values lower than 0.154 ng/ml, poor responders 0.154–0.7 ng/ml, normal responders 0.71–2.1 ng/ml and high responders above 2.11 ng/ml. Lee et al. further subdivided the group of high responders by introducing excessive responders with basal values above 3.35 ng/ml [7].

Aims of the study

The goal of this study was to compare pregnancy rates and complication rates, especially the occurrence of OHSS, under a lower, individualized gonadotrophin dosing mainly in the long agonist protocol in a clinical setting.

Materials und Methods

This study was designed as a single center retrospective clinical study. Data of cycles of controlled ovarian stimulation for IVF/ICSI were analyzed according to the criteria of PCOS.

Study population

We analyzed all stimulation cycles for IVF/ICSI of patients within three years, from 01.01.2012 to 31.12.2014, who were performed at the fertility center of the University Hospital, Department of Gynecology and Obstetrics of the Technical University of Dresden, Germany. With consent of the institutional ethics committee and written informed consent of the patients, 370 cycles of 235 women were reviewed, focusing on diagnosis of PCOS. Those patients with PCOS (n = 39) constitute the study population, while all other women (n = 196) serve as control group.

Inclusion criteria

According to the inclusive Rotterdam Criteria (Rotterdam ESHRE/ASRM-Sponsored PCOS consensus workshop group, 2004), we included women with at least two out of three following characteristics:

- ▶ Hyperandrogenemia (at least one of the androgens above the 95th percentile: free testosterone, androgen-index, androstenedione and DHEAS)
- ▶ Report of oligomenorrhea with cycle length of more than 35 days, amenorrhea with cycles lasting longer than three months or absence of a rise of progesterone before menstruation during cycle monitoring
- ▶ Documented polycystic ovarian morphology by an experienced sonographer or visualization of polycystic ovaries according to Rotterdam Criteria on printed pictures (more than 12 small antral follicles of 2–9 mm size in one ovary)

Exclusion criteria

All women stimulated were analyzed. The study group consisted of women with diagnosis of PCOS. Patients with clinical signs of PCOS but additional diagnosis of clinical thyroid dysfunction, early or late onset adrenogenital syndrome (AGS), androgen producing neoplasm, Cushing’s syndrome, hypogonadotrophic hypogonadism, premature ovarian failure, hyperprolactinaemia, HAIRAN-syndrome or intake of exogenous androgens were not regarded as PCOS according to Rotterdam criteria [2].

Six patients fulfilling diagnostic criteria of PCOS were not included in the study group, three because of the diagnosis of Hashimoto’s thyroiditis, two patients exhibited hypogonadotrophic hypogonadism and one woman was diagnosed with late onset AGS. One patient underwent stimulation without fertilization of oocytes for purpose of fertility preservation and two others had missing clinical data and were not included in the PCOS-study group.

Data collection

A total of 235 couples underwent ART in our center and were analyzed for the study. Prior to treatment three diagnostic appointments consisting of cycle monitoring, hormone profile and transvaginal ultrasound were used to exclude anatomical malformation, assess AFC and confirm ovulation. The medical history was taken separately for each partner and at least one sperm count was analyzed according to WHO criteria from 2010 [8]. OHSS was scored according to the classification of Aboulghar und Mansour [9]. A mild manifestation is characterized by mild

Table 1 Comparison of studies on controlled ovarian stimulation with FSH in recent studies of the last 10 years.

Reference	Study	FSH starting dose for PCOS	Recommendation for dose adjustment
Palep-Singh et al., 2007	Observational comparative study (PCOS in Asian women n = 104, Caucasian n = 220, controls n = 284)	Mean starting dose 150 IU for Asian PCOS 200 IU for Caucasian PCOS 225 IU for Asian and Caucasian tubal infertility	Adjustment according to age, basal FSH and BMI, after 7 days possible increase of dosage in case of suboptimal response
Weghofer et al., 2007	Retrospective cohort study of 47 women with PCOS compared to 100 controls	150–450 IU for PCOS and Non-PCOS	Adjustment to follicular response
Koundouros et al., 2008	Prospective randomized study with PCOS-patients (n = 225)	75 IU/d step up 225 IU/d step down 150 IU and individual adjustment	Step-up regimen: 75 IU/d for 6 days, then increase of 37.5 IU Step-down: 225 IU/d of FSH for the first 3 days followed, then decrease to 150 IU/d for the next 3 days, then decreased to 75 IU/d or sustain at 150 IU Step-up/Step-down: 150 IU on day 1, then decrease to 75 IU on day 2, then increase back to 150 IU and so on until day 6, then, sustain at 150 IU/d or 75 IU/d
Sahu et al., 2008	Retrospective analysis of 51 PCOS ART-cycles, compared to 50 cycles with ultrasound morphology of PCO, control group 104 cycles	300 IU	According ovarian reserve score (age, BMI, AFC, FSH, E2), control on day 4
Swanton et al., 2010	Prospective cohort study (n = 290 women, including PCOS n = 78, PCO n = 101 and control n = 101)	150–375 IU according to age, basal FSH and previous ovarian response to gonadotropins – no difference in PCOS, PCO	No adjustment described
Ashrafi et al., 2011	Controlled randomized prospective study with n = 90 women with PCOS – three protocol variations concerning type of gonadotropins given and dose adjustment	150 IU for PCOS in all groups	Fixed dose of 150 IE Step down protocol to 75 IU, when leading follicle reached 14 mm in diameter FSH discontinued and low dose HCG when leading follicle 14 mm
Decanter et al., 2013	Single center prospective non-randomized interventional study (n = 113). Intervention was pretreatment with oral contraceptives	100 to 200 IU, according to age, BMI and AFC	No adjustment described
Huber et al., 2013	Retrospective cross-sectional study with n = 7 520 cycles, mixed cohort, no recommendation for PCOS	Mixed cohort, 75–450 IU; 75–125 IE for expected high response, 150–225 IU for normal response 300–450 IU for poor response, according to age, markers of ovarian reserve, BMI, and previous response	Individual adjustment to response No recommendation for PCOS
Figen Turkcapar et al., 2013	Prospective randomized controlled study (n = 80 women with PCOS), HMG vs. FSH stimulation	150 IU	Adjustment according to E2, sonographical response
Akpınar et al., 2014	Retrospective observational study of n = 337 cycles of women with PCOS	75–300 IU according to BMI	No adjustment described
Shi et al., 2014	Multicenter prospective randomized controlled study of 1 180 women with PCOS. Intervention randomized in fresh ET at day 3 or freeze all	112.5 IU/day for patients ≤ 60 kg for PCOS 150 IU/day for patients > 60 kg for PCOS	Adjustment according to ovarian response

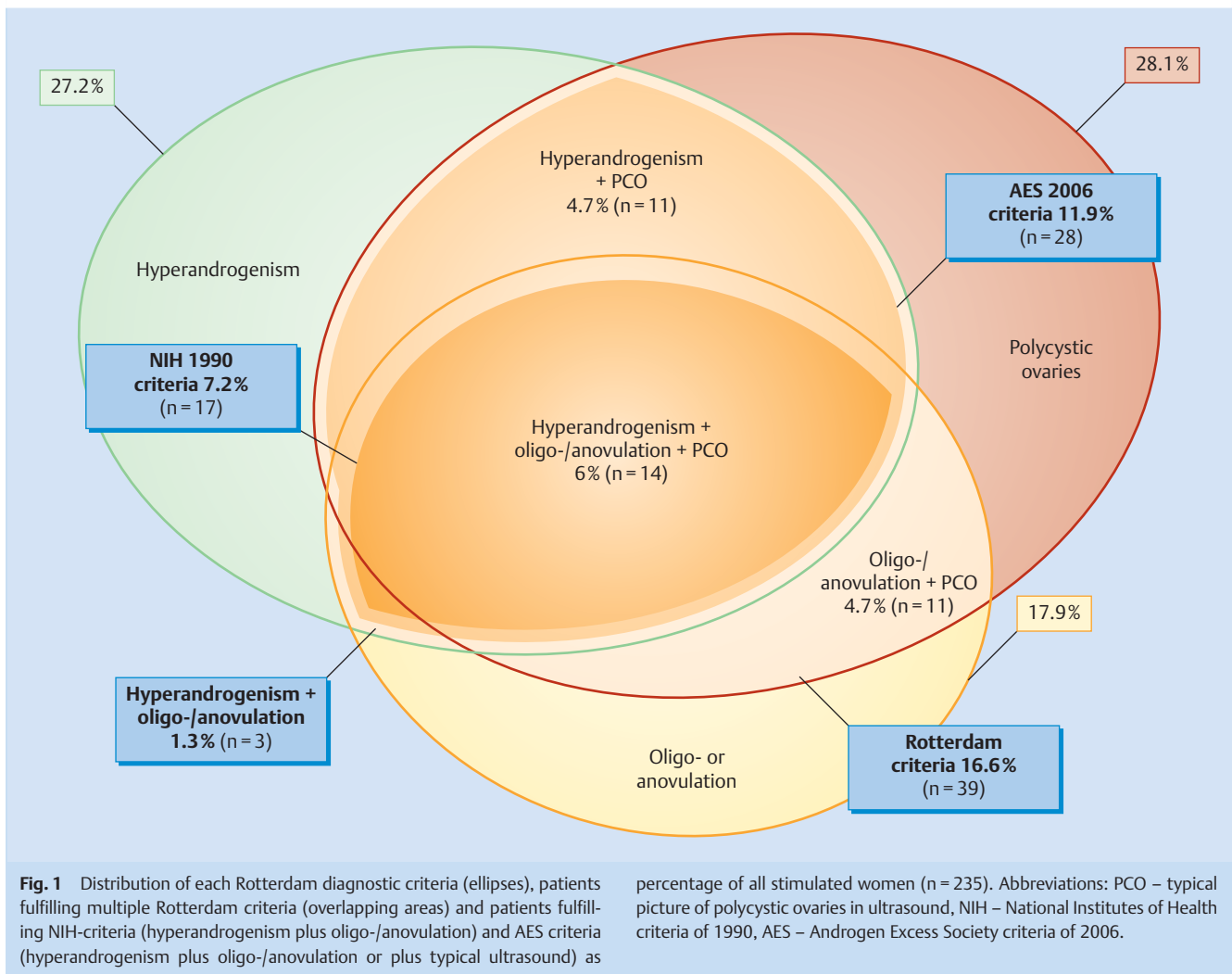
symptoms, abdominal pain or distension and enlarged ovaries but no visible fluids in the pouch of Douglas on ultrasound scan. Moderate OHSS appears with sonographic evidence of ascites accompanying further symptoms but without shift of haemostaseological or biochemical serum parameters. In case of disarrangement of hemostasis, elevated liver enzymes, haematoconcentration, elevated creatinin, dyspnea, oliguria, massive ascites or pleural effusion, severe hyperstimulation is present.

Intervention

For GnRH-agonist protocol, Nafarelin (Synarela[®], Pharmacia, Copenhagen, Denmark), Triptorelin (Decaptyl[®], Ipsen Pharma, Barcelona, Spain, or as Depot: Ferring Arzneimittel GmbH, Kiel, Germany) or Leuprorelin (Enantone[®], Takeda Pharmaceutical Company Limited, Osaka, Japan) were used for the purposes of downregulation. In some cases, an oral contraceptive pill was giv-

en for one cycle overlapping with the agonist downregulation or before the start of stimulation in antagonist cycles, predominantly 30 µg ethinylestradiol and 125 µg levonorgestrel (Minisiston[®]). Cetrotide[®] (Merck Serono, MSD, the Netherlands) or Ganirelix (Orgalutran[®], Organon, Skovlunde, Denmark) were injected in the multidose flexible antagonist-protocol.

We used recombinant FSH Gonal-F[®] (Serono Pharmaceuticals Ltd., Feltham, UK), Puregon[®] (Organon Laboratories Ltd., UK), or urinary hCG Menogon[®] (Ferring Pharmaceuticals, Istanbul, Turkey) for gonadotropin stimulation. The starting dose was calculated depending on age, BMI and basal antimullerian hormone (AMH) of the patient, as well as on ovarian response in previous cycles, diagnosis and types of diagnostic criteria of PCOS and sonographic appearance of the ovarian function (AFC). Decision was made according to the Dose-Finding-Chart (see below, **Fig. 3**). The final oocyte maturation was induced with



10000 IU hCG (Predalon®, Brevactid®) or 250 IE reHCG (Ovitrelle®, Merck Serono, MSD). Ultrasound guided oocyte retrieval was performed 36 hours later under general anesthesia.

Biochemical analyses and outcome

The free androgen index (FAI) was calculated from total testosterone (nmol/l) \times 100/SHBG (nmol/l). For the measurement of LH, FSH, estradiol, progesterone, hCG and prolactin sandwich-immunoassay-Kits from ADVIA Centaur®, Siemens Healthcare Diagnostics, Inc. (Tarrytown, U.S.A.) were used. Radioimmunoassays for free testosterone (Active® Free testosterone RIA), sex hormone binding globulin (SHBG IRMA KIT), dehydroepiandrosterone sulfat (DHEAS-S-7 RIA) and AMH (AMH Gen II ELISA-Kit) were obtained from Beckman Coulter (Galway, Ireland). Asbach Medical Products GmbH (Obrigheim, Germany) provided radioimmunoassays RIA CT for 17-OH-progesterone und androstendione. Beginning with the second half of 2013, we used RIA testosterone direct for total testosterone, Active® Free testosterone, Active® androstendion and SHBG IRMA KIT from Beckman Coulter. DHEAS was quantified by radioimmunoassay Immulite® DHEA-SO4 from Siemens Healthcare Diagnostics, Inc. (Tarrytown, U.S.A.) and 17-OH-progesterone by ELISA from IBL INTERNATIONAL GmbH (Hamburg, Germany). A biochemical pregnancy was confirmed with a serum β -hCG above 10 E/l 12 to 16 days after ET. A gestational sac four to five

weeks after embryo transfer visible on transvaginal ultrasound scan was evaluated as a clinical pregnancy.

Statistical analyses

Following the analysis of the study population, the PCOS- and control group were described and compared using SPSS Statistics (Version 22.0.0.0). Nominal and ordinal data were characterized by frequency, and compared using a χ^2 test or Fisher's exact test for smaller sample sizes. Metrical data were analyzed by descriptive statistics. Using the Levene's test confirmed similar variance of the compared groups and allowed the use of Students t-test, otherwise further significance was calculated by Wilcoxon test (Mann-Whitney-U-Test). $P < 0.05$ was considered as statistically significant.

Results

Characterization of the study population and collectives

The study included 370 stimulation cycles of 235 women treated at the infertility unit of the University Hospital in Dresden within three years. In vitro fertilization (IVF) was performed in 43.8% of the cycles, intracytoplasmic sperm injection (ICSI) in 54.9% of cases, and in the remaining percentage IVF/ICSI splitting was performed.

Table 2 Patient characteristics of 235 patients treated with controlled stimulation for ART (mean \pm SD) and cycle characteristics of 370 controlled stimulation cycles and outcome of 329 ET (mean \pm SD). Bold font: $p < 0,05$, considered as statistically significant.

	Control group n = 196	Patients with PCOS n = 39	p-value
Age (years)	34.13 \pm 4,13	32.739 \pm 4.11	$p = 0.055$
BMI (kg/m ²)	23.30 \pm 4,35	25.86 \pm 5.70	$p = 0,007$
AMH (ng/ml)	2.71 \pm 2,33	7.50 \pm 4.72	$p < 0,001$
Duration of infertility (years)	4.22 \pm 2,83	4.33 \pm 2.92	$p = 0.825$
Primary infertility (%)	66.8	53.8	$p = 0.121$
Number of cycles (n)	305	65	
Number of embryo transfers (n)	270	59	
Agonist protocol (%)	74.1	72.3	$p = 0.476$
FSH starting dose (IU)	172.17 \pm 67.228	131.92 \pm 66.366	$p < 0,001$
Total FSH dose (IU)	1929.16 \pm 895., 893	1721.94 \pm 812.872	$p = 0.086$
Duration of stimulation (days)	10.40 \pm 2.053	11.72 \pm 2.886	$p < 0,001$
Step up (%)	35.7	53.8	$p = 0,025$
Dose adjustment (%)	44.6	60.0	$p = 0,025$
Max. serum E2 (pg/ml)	1641.35 \pm 936.13	1672.77 \pm 1100.22	$p = 0.812$
No. of follicles \geq 16 mm (n)	7.96 \pm 4.112	9.81 \pm 4.253	$p = 0,001$
No. of oocytes (n)	8.41 \pm 4.758	9.95 \pm 5.311	$p = 0,022$
Fertilization rate (%)	0.559 \pm 0.298	0.608 \pm 0.275	$p = 0.356$
Ratio embryo transfer/stimulation cycle	270/305 88.5%	59/65 90.8%	
Biochemical pregnancy rate/ET (%)	42.2	42.4	$p = 0.547$
Clinical pregnancy rate/ET (%)	34.4	32.2	$p = 0.434$
Moderate or severe OHSS (%)	15.7	16.9	$p = 0.852$
Early pregnancy loss (%)	10.8	21.1	$p = 0.191$

PCOS was diagnosed according to Rotterdam criteria in 16.6% of patients. If NIH criteria were applied, 7.2% of the study population were considered having PCOS and with AES criteria 11.9%. The distribution of Rotterdam phenotypes and the classification of FAS of the study group is visualized in **Fig. 1**. For our study we explicitly differentiate patients in group FAS III with hyperandrogenism, metabolic syndrome, polyfollicular ovaries from FAS IV, because women with FAS III are at higher risk for hyperstimulation. FAS IV group women also show metabolic syndrome and hyperandrogenism but ovaries with normal or even low ovarian reserve. In terms of fertility treatment, those patients often perform as low responders in controlled ovarian hyperstimulation. Therefore we decided to use in this study the Rotterdam criteria to select the study group.

Characteristics of the study groups are shown in **Table 2**, as well as differences in stimulation regime, ovarian response and outcome. The mean age in both groups was not statistically different. Mean BMI and AMH-levels were significantly higher in the PCOS-group.

One cycle had to be canceled for impending hyperstimulation and there was no oocyte retrieval performed after hormonal stimulation in one case because of the absence of sperms after testicular sperm extraction.

Stimulation modality depending on AMH-level

Classifying our cycles into AMH-responder groups a positive trend of the portion of cycles with PCOS diagnosis, polycystic ovarian morphology and oligo- or amenorrhea between the AMH-groups becomes visible (χ^2 test, all $p < 0.001$). This does not apply using the criteria of hyperandrogenaemia as a predictor of response ($p = 0.066$).

The mean FSH-starting and maximum dose, as well as the total dose of FSH differed within the AMH-responder groups, declining by rising AMH-levels (ANOVA $p < 0.001$) (**Fig. 2**).

Splitting the basal AMH-value of excessive responders into additional subgroups, the occurrence of each manifestation of the OHSS is not significant (p -value for mild $p = 0.765$; moderate $p = 0.144$ and severe $p = 0.097$). But on pooling moderate or severe manifestation a significant increase with higher AMH-values becomes visible ($p = 0.005$) (**Fig. 2**). Significantly more follicles developed, higher estradiol levels were reached and oocytes could be retrieved within more sensitive groups (ANOVA all p -values < 0.001).

Outcome

In our study the biochemical pregnancy rate of all patients per ET was 42.2% and the clinical pregnancy rate per ET 34.0%. The biochemical pregnancy rate of 42.4% and the clinical pregnancy rate of 32.2% for PCOS-patients did not differ statistically from those of the control group. 32.1% of all clinical pregnancies were twins, there were no higher multiple pregnancies. Two cases of ectopic pregnancy occurred and 12.5% of clinical pregnancies aborted. Miscarriage rate was similar in both groups (**Table 2**).

A mild OHSS occurred in 3.5%, moderate in 8.4% and severe in 7.6% of all stimulation cycles, which resulted in the need for hospitalization for 1.9% of treatments. All seven patients hospitalized for OHSS had no prior diagnosis of PCOS, and hospitalization does not differ between study groups ($p = 0.611$). Neither a mild, nor a moderate or severe manifestation of OHSS occurred significantly more often in patients diagnosed with PCOS. Albeit, in the PCOS collective, one stimulation cycle (all PN-oocytes) was cryopreserved without fresh-embryo transfer (freeze all) and one cycle was coasted because of impending OHSS.

Analysis of the Rotterdam phenotypes

Highest basal AMH-levels can be found for the phenotype with expression of all three features of the Rotterdam criteria. With

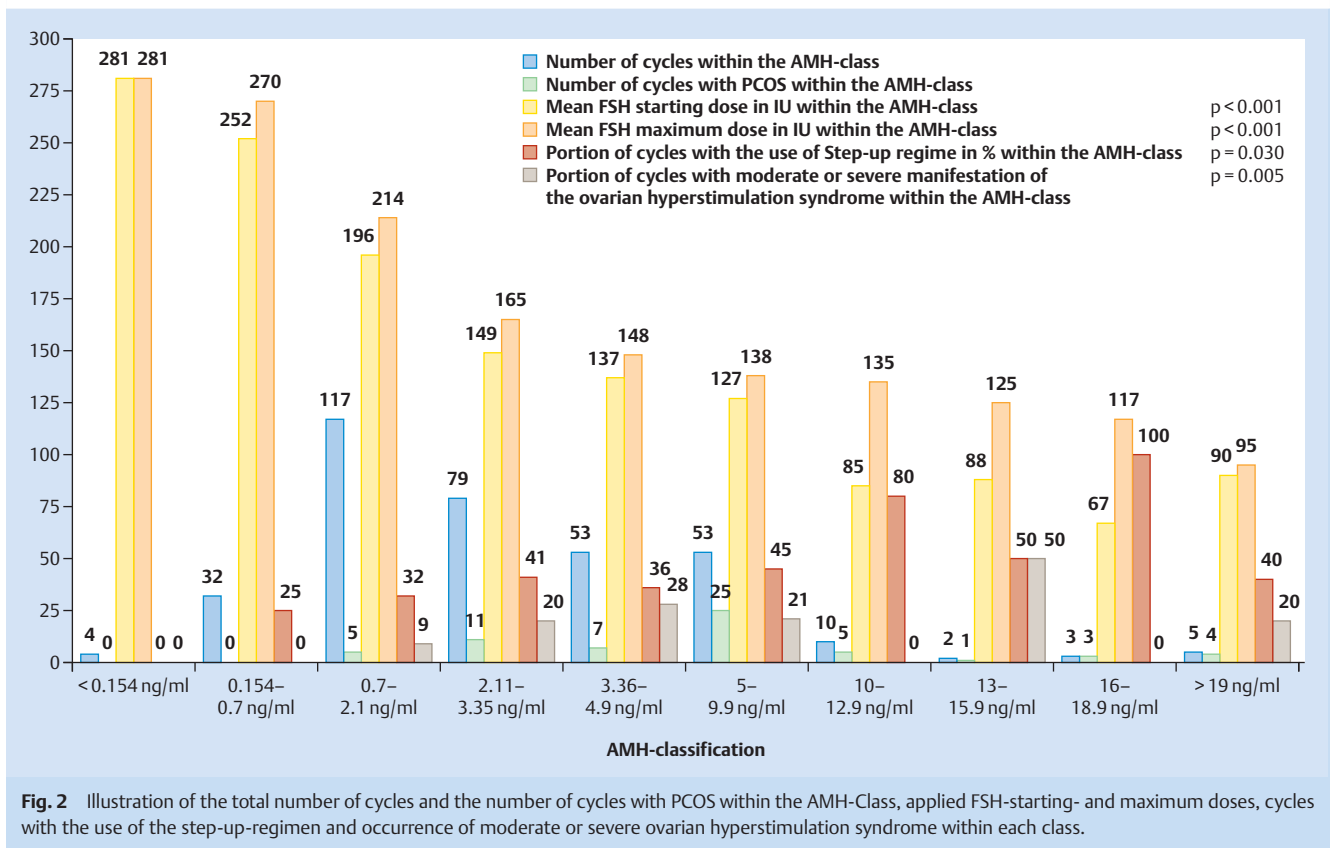


Fig. 2 Illustration of the total number of cycles and the number of cycles with PCOS within the AMH-Class, applied FSH-starting- and maximum doses, cycles with the use of the step-up-regimen and occurrence of moderate or severe ovarian hyperstimulation syndrome within each class.

45%, this group accounted for the largest proportion of all cycles of PCOS patients.

For PCOS-patients with oligo- or anovulation and polycystic ovarian morphology, we applied the lowest mean FSH dose (98 IU). Starting dose of the type with hyperandrogenism and oligo- or anovulation is biased by one patient with three stimulation cycles, accounting for one third of all cycles within this group. Her AMH-level was 1.35 ng/ml, aged 41–44 years and her BMI 32 kg/m². In this case, we decided to start with 200–300 IU. For the Rotterdam phenotype with hyperandrogenism and polycystic ovarian morphology as well as for the phenotype with expression of all criteria, moderate and severe OHSS occurred in more than 10% of cycles. Nevertheless, the distribution of several manifestations of the OHSS shows no significant deviation of the Gaussian distribution (Table 3).

Discussion

Clinical pregnancy rates after ART of women with PCOS differ between 22 and 42% in literature all over the world. This wide spectrum of pregnancy rates is due to different patients' characteristics and treatment methods. Furthermore there are worldwide differing strategies for artificial reproduction with variations of number of transferred embryos, using blastocyst culture and elective single embryo-transfer or pre-implantation genetic diagnosis.

Nevertheless compared to the literature, our calculated stimulation regime represents a feasible option for both PCOS and Non-PCOS patients, with a satisfying result of 34.0% clinical pregnancy rate per embryo transfer.

By investigating patients with and without PCOS diagnosed by Rotterdam criteria, we could not find a difference in the clinical pregnancy rate, confirming results of other authors [10–12]. Results were comparable for PCOS patients, despite significantly higher BMI values compared to the control group, which essentially impairs fertility treatment outcomes [13,14]. Our PCOS study population had a mean BMI of 25,9 kg/m², which is considered overweight according to WHO and confirmed this observation.

The proposed stimulation regime with calculated low-dose stimulation for PCOS did not show elevated frequency of OHSS. Moderate or severe OHSS occurred in 8.4% and in 7.6% of all stimulation cycles and ranges within the rates in literature (5 to 16.6% [10,15,16]). However, lean body weight and low BMI in PCOS increases the OHSS risk [17,18]. Interestingly in our study the need for hospitalization exclusively affected patients without PCOS. Although the absolute number of women with OHSS in all groups was low, the higher awareness for OHSS in women with PCOS and anticipated risk may have resulted in this reduction of risks.

To prevent OHSS, in one case we decided to reduce drastically the gonadotropin dose ("coasting") and in another case we froze all fertilized oocytes without fresh ET. Both treatments were applied to the same patient in subsequent stimulation cycles. The women showed extreme sensitivity to gonadotropins in two following cycles, thus representing a potential candidate for in vitro maturation.

Concerning spontaneous miscarriage rates of patients with PCOS conceiving with ART, most studies did not show a significantly elevated miscarriage rate for women with this diagnosis [19,20]. Corresponding to these results, patients treated in our clinic did

Table 3 Analysis of the Rotterdam phenotypes according to characterization, stimulation modalities and outcome. Bold font: $p < 0,05$, considered as statistically significant.

	No. PCOS	Hyperandrogenism + oligo-/anovulation	Hyperandrogenism + PCO	Oligo- or anovulation+ PCO	Hyperandrogenism + oligo-/anovulation + PCO	p-value
Number of patients (n)	196	3	11	11	14	
Number of cycles (n)	305	10	14	12	29	
Portion of cycles of the phenotype (%)	82.4	2.7	3.8	3.2	7.8	
Portion of cycles with PCOS (%)		15.4	21.5	18.5	44.6	
Mean age (y)	34.14 ± 4.13	35.55 ± 5.20	32.53 ± 4.01	32.10 ± 4.75	32.80 ± 3.65	0.654
Mean BMI (kg/m ²)	23.30 ± 4.35	29.90 ± 1.83	25.49 ± 4.87	26.18 ± 6.56	25.01 ± 6.15	0.613
Mean basal AMH-level (ng/ml)	2.71 ± 2.33	3.71 ± 3.68	5.61 ± 1.83	7.44 ± 4.76	9.84 ± 5.55	0.061
Portion of cycles with step-up regimen (%)	109 (35.7%)	3 (30%)	6 (42.9%)	7 (58.3%)	19 (65.5%)	0.132
Portion without dose adjustment (%)	169 (55.4%)	6 (60%)	7 (50%)	4 (33.3%)	9 (31.0%)	0.132
Portion of cycles with use of oral contraceptives (%)	28 (9.2%)	2 (20%)	0%	1 (8.3%)	14 (48.3%)	< 0.001
Portion of cycles with agonist protocol (%)	226 (74.1%)	6 (60%)	12 (85.7%)	10 (83.3%)	19 (65.5%)	0.419
Mean FSH-Starting dose (IU)	172.17 ± 67.23	245.00 ± 48.31	110.71 ± 27.24	97.92 ± 29.11	117.24 ± 55.94	< 0.001
Mean total FSH dose (IU)	1929.16 ± 895.89	2937.50 ± 976.69	1341.07 ± 342.59	1370.83 ± 596.76	1631.93 ± 589.78	< 0.001
Mean endstimulatory estradiol level (pg/ml)	1641.35 ± 936.127	1140.80 ± 856.31	1957.86 ± 1092.07	1573.42 ± 853.66	1759.69 ± 1239.89	0.323
Portion of cycles with mild OHSS (%)	12 (3.9%)	0	0	0	1 (3.4%)	0.832
Moderate OHSS (%)	25 (8.2%)	0	3 (21.4%)	0	3 (10.3%)	0.260
Severe OHSS (%)	23 (7.5%)	0	1 (7.1%)	1 (8.3%)	3 (10.3%)	0.886

not show a difference. Nevertheless higher BMI and PCOS may contribute additionally to the elevated miscarriage rate [21, 22]. For PCOS, studies report higher numbers of developing follicles and aspirated oocytes compared to controls [10, 23–26]. Similarly, our PCOS group showed significantly higher average numbers of follicles and retrieved oocytes although we used lower FSH-doses as usually recommended in our controlled stimulation protocol. Primarily, outcome parameters are rising with the extent of the reaction to gonadotropin stimulation, coming to a plateau and decreasing with stronger response to stimulation. Decreasing pregnancy [27] and live birth rates [16] are reported when more than 15 oocytes were aspirated. Lower doses of gonadotropins lead to smaller reactions, which would be in favour of more sensitive patients like those with PCOS.

The FAS-classification proposed by Geithövel [3] differentiates the subgroup of women with hyperandrogenemia and response to COS into high and low response. Our data show, that patients in group FAS III, which show hyperandrogenism, metabolic syndrome and polyfollicular/polycystic ovaries are at risk for ovarian hyperstimulation. On the contrary women with FAS IV – even though hyperandrogenism is present – are at risk for low response. Gonadotropins should be adapted to the parameter of ovarian reserve, AMH and AFC.

Analysing applied doses in our regime, adjustments and experiences in literature, we come to the following Dose-Finding-Chart for the FSH-starting dose (● Fig. 3).

On average after calculating the individual dose, we started stimulation with 132 IU FSH in PCOS-patients. Though individually adjusted, it can be necessary to lower this dose down to 75 IU.

These doses range below the doses of the “mild stimulation” established by the group around Fauser et al. [28]. Despite our low-dose strategy, our results are not inferior to those in literature in terms of outcome and risks.

Nelson et al. [6] and Yates et al. [29] recommended an AMH-stratified approach for the decision of the stimulation protocol and FSH dose. For extreme responders (according to AMH above 2 ng/ml and Yates above 3.9 ng/ml) they suggest choosing the antagonist protocol, because cycle cancellation rate is lower and risk of OHSS is already reduced with this type of protocol. From this cut-off, both authors started with a dose of 150 IU FSH/d. Also for extremely low responders, below 0.14 resp. 0.3–2.1 ng/ml, both recommend the antagonist protocol.

In the study of Gera, for cycles with estradiol levels above 2500 pg/ml or more than 30 growing follicles, coasting and if necessary elective oocyte or embryo cryopreservation and subsequent transfer in a hormonal prepared cryo-cycle were initiated. In this publication, the incidence of OHSS is lowered without impairing pregnancy- and live birth rate [30].

Steward emphasizes the need of less aggressive stimulation strategies for patients at risk and refers to the method of the GnRH-antagonist-protocol with the agonist-trigger [16].

Besides intensive support with administration of progesterone and E2 and the separation of the stimulation cycle and oocyte pick-up from the ET, the investigation group of Humaidan, Engmann and Benadiva refined the strategies against the impairment of endometrial quality by luteolysis. They suggest additional support of the luteal phase with low-dose hCG-injections and fresh ET and reached comparable results for clinical and ongoing

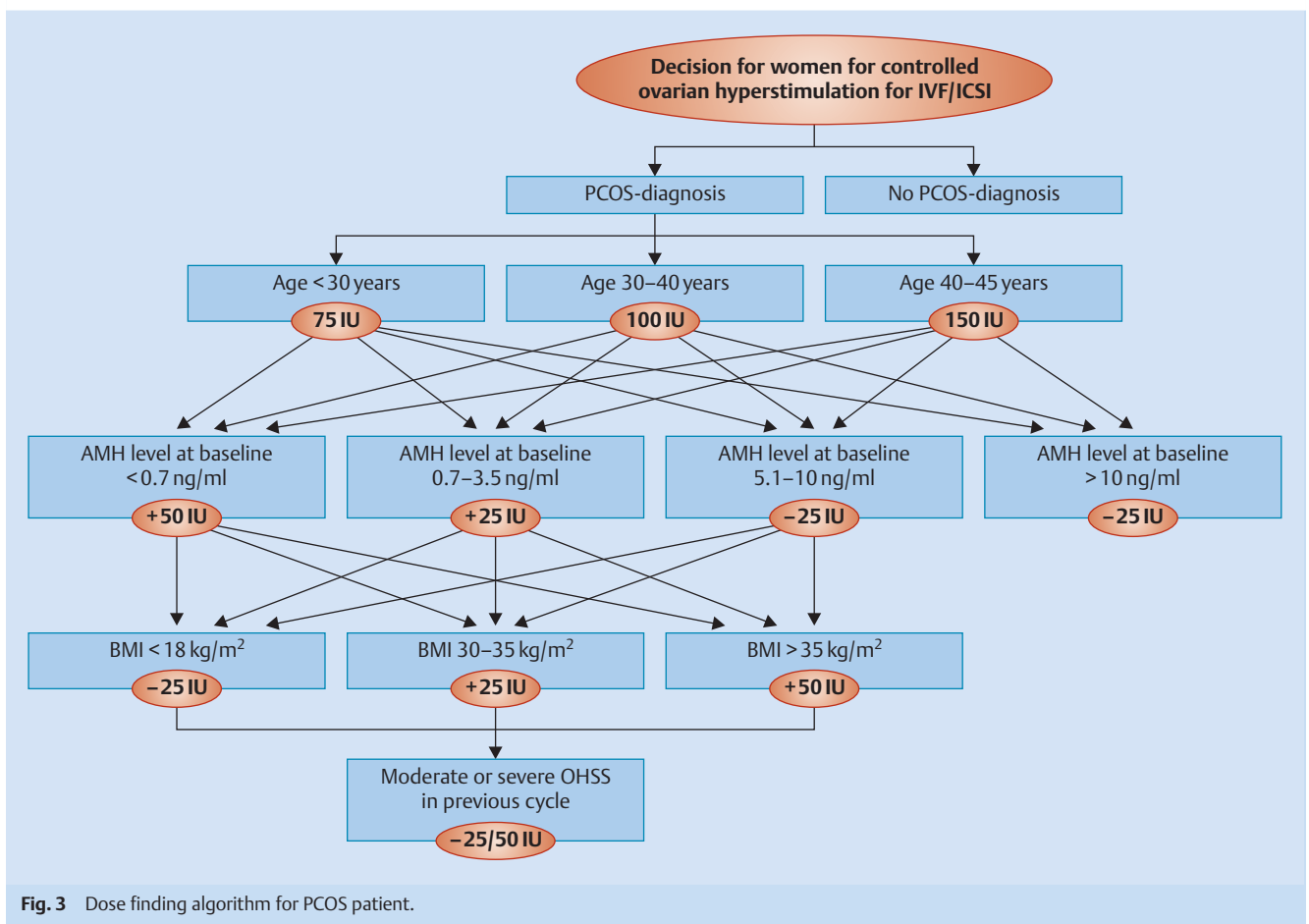


Fig. 3 Dose finding algorithm for PCOS patient.

pregnancy rates as well as OHSS rates in a randomized controlled trial compared to hCG-trigger. For very high-risk patients responding with more than 25 follicles, they still chose to freeze all embryos and transfer in a subsequent cycle [31, 32]. A combination of this strategy and a clear recommendation for an individually adjusted low-dose stimulation is conceivable. Nevertheless, despite the use of GnRH-agonist-trigger and low-dose hCG luteal support, for patients under elevated risk, up to 26% severe OHSS were seen when fresh ET was performed [33].

Abstaining from exogenous hCG luteal support and choosing elective cryopreservation instead of a fresh transfer improves safety of controlled ovarian stimulation for patients at high risk, at the price of lower pregnancy rates [34]. However, even with those preventive measures, six cases of severe OHSS are reported [35–37]. It seems that up to date there is no complete and effective prevention of OHSS. In the case of Ling, the patient showed basal serum AMH level of 64.5 ng/ml and developed more than 40 follicles [37]. Also in the case reports of Gurbuz, one instance of 27 and another one of 52 follicles with 45 retrieved oocytes and serum estradiol levels above 5985 and 10904 pg/ml were observed [36]. All cases show that even without endogenous hCG rising in early pregnancy after fresh ET, early-onset OHSS can occur with any kind of gonadotropin administration. If IVM cycles are primed with 125 IU FSH and ovulation is triggered by exogenous hCG as previously described [34, 38], theoretically the risk of OHSS is even existing with “safe” low-dose IVM-strategy.

There is no strategy to prevent completely the risk of OHSS when gonadotropins are used. On the other hand, very sensitive patients with extremely exaggerated ovarian responses to gonadotropins are extremely rare. The challenge is to identify those patients who are at risk with the well-established approaches and offer the chance to use selectively the safe alternative of IVM. Most fertility centers in the world do not offer the reserve method of IVM. Establishment of IVM techniques and laboratory expertise is for the majority of IVF-centers inaccessible and not cost-effective. Studies show that the technique is feasible, although a longer time of training is required [38, 39]. A solution could be the improvement of existing techniques and experience with exact recommendations from large RCTs and health institutes. Few, individual centers, offering IVM should get the chance to refine their IVM-techniques and knowledge. The majority of fertility centers should filter those patients for IVM and refer them to centers with special expertise in IVH.

The major drawbacks of our study on individually calculated low dose stimulation for PCOS is the retrospective character and the low number of participants. For a following randomised trial, clear guidance for decision-making should be developed out of these first results. The different phenotypes of Rotterdam diagnosed PCOS should be considered in detail.

Conclusion for Clinical Practice

Patients with PCOS represent a challenge for reproductive medicine. We propose a calculated low-dose stimulation strategy with step-up according to ovarian response in long agonist and antagonist protocols.

The choice of the starting dose of FSH has to be calculated by patients' basal AMH level, AFC, age, BMI and PCOS diagnosis. Response to previous stimulation cycles should be integrated as another important clinical parameter, according to our dose finding algorithm.

Conflict of Interest

M. Goeckenjan declares reimbursement of traveling expenses by MSD, Merck Serono and Ferring.

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