Affecting 8 to 13% of women of reproductive age and 21% in high risk groups, polycystic ovary syndrome (PCOS) is the most prevalent reproductive disorder causing significant health consequences for women impairing quality of life and increasing morbidity. The Rotterdam diagnostic criteria for PCOS are now internationally endorsed and are based on two of three features: oligo- or anovulation, hyperandrogenism (clinical or biochemical), and polycystic ovaries. Insulin resistance and hyperinsulinemia play an important role in the pathophysiology and metabolic manifestations of PCOS, independent of but exacerbated by obesity. Early diagnosis is key in addressing symptoms, improving quality of life, and identifying fertility problems, as well as for long-term considerations including metabolic, cardiovascular, and psychological factors.
psychosocial features. There is increasing evidence of the effectiveness of lifestyle interventions on nonreproductive as well as reproductive outcomes. Medical therapy is effective as are infertility treatments in PCOS. Here, we outline a summary of the most recent insights in diagnostic criteria, clinical features, and management of women with PCOS including a brief summary of the recommendations from the latest international evidence-based guidelines. The full guideline and freely available translation resources can be found at https://www.monash.edu/medicine/sphpm/mchri/pcos and the full recommendations summary is published elsewhere.

**Diagnostic Criteria**

The Rotterdam diagnostic criteria are commonly used globally to diagnose PCOS and were the criteria endorsed by the recent international evidence-based guidelines on the diagnosis and management of PCOS. The Rotterdam diagnosis for PCOS requires two of the following features: oligo- or anovulation, clinical and/or biochemical hyperandrogenism, and polycystic ovaries. In adults if oligo- or anovulation and clinical and/or biochemical hyperandrogenism are present, ultrasound may not be required. Additionally, other endocrine etiologies such as thyroid disease, nonclassic congenital adrenal hyperplasia, and hyperprolactinemia should be excluded with further evaluation guided by clinical judgement. The guideline refines Rotterdam criteria to highlight that in adolescents or those within 8 years of menarche, ultrasound is not a reliable discriminator for PCOS. Hence in this group, both oligo- or anovulation and clinical and/or biochemical signs of hyperandrogenism are required and ultrasound is not recommended for the purpose of diagnosis.

**Ovulatory Dysfunction and Irregular Menstrual Cycles**

When irregular or absent menstrual cycles occur, a diagnosis of PCOS should be considered at approximately 85 to 90% of women with oligomenorrhea and 30 to 40% of women with amenorrhea have PCOS. Ovulatory dysfunction is assessed after 1 year post-menarche; before this time period, irregular cycles are part of the normal pubertal transition. Ovulatory dysfunction is clinically reflected by irregular menstrual cycles, which are defined as shorter than 21 days or longer than 45 days in women between 1 and 3 years post-menarche, and less than 21 or more than 35 days in women over 3 years postmenarche to perimenopause. Additionally, when women present with primary amenorrhea by the age of 15 or more than 3 years post-thelarche, ovulatory dysfunction is probable. It is, however, possible even with regular cycles to have ovulatory dysfunction. Luteal phase biochemical serum progesterone levels can be used to assess ovulation.

**Hyperandrogenism**

In women with PCOS, hyperandrogenism is highly prevalent with a recent study reporting hyperandrogenism in 78% of women with PCOS and even higher prevalence in overweight women. When evaluating symptoms of hyperandrogenism, hirsutism should be assessed using the modified Ferriman-Gallwey score (mFG), where a level ≥4 to 6 indicates hirsutism. Alopecia can be assessed using the Ludwig visual score. For acne, no universally accepted visual assessment is available. When clinical signs of hyperandrogenism are unclear or absent, measurement of calculated free testosterone, free androgen index, or calculated bioavailable testosterone should be undertaken. High-quality assays such as liquid chromatography-mass spectrometry/mass spectrometry and extraction/chromatography immunoassays are most accurate. Evaluation should be undertaken when women are not on hormonal contraception. If total or free testosterone is not elevated, consideration can be given to assessing androstenedione and dehydroepiandrosterone sulfate (DHEAS). In a small prospective cross-sectional study of 82 patients with PCOS, DHEAS showed higher levels in PCOS patients with PCO morphology compared with controls. Furthermore, a negative correlation between DHEAS levels and mean and maximum ovarian volume was reported. However, these findings remain unexplored in a clinical setting and there is currently no clear evidence that DHEAS is associated with ovarian morphology in PCOS.

**Polycystic Ovarian Morphology**

Where oligo- or anovulation and clinical and/or biochemical hyperandrogenism are present in adults, ultrasound is not strictly necessary in diagnosing PCOS; however, it will help identify the complete phenotype and may be useful for other indications in PCOS. The guideline recommends using a follicle number per ovary of more than 20 follicles (2–9 mm) and/or an ovarian volume ≥10 mL using transducer frequency ≥8 MHz. For older ultrasound equipment, the criterion is an ovarian volume ≥10 mL. Due to the high incidence of polycystic ovaries and nonspecificity in those with a gynecological age of less than 8 years postmenarche, ultrasound is not recommended at this life stage for the purposes of diagnosis. Women with polycystic ovarian morphology (PCOM) exhibit higher levels of anti-Mullerian hormone (AMH) and higher levels of AMH are associated with anovulation and hyperandrogenism. Current research is evaluating whether serum AMH can replace ultrasound in the diagnosis of PCOM; however, the guideline does not recommend AMH for the detection of PCOM or the diagnosis of PCOS. Current barriers to applicability of AMH include poorly defined patient and control populations, variable quality assays, and no international standards. Additionally, there is a lack of data on clustering of AMH levels with other PCOS features and relationships to long-term outcomes and these are areas of future research.

**Prevalence and Phenotypes**

The prevalence of PCOS varies depending on the diagnostic criteria, phenotypes, and populations studied. Bozdag et al reviewed a total of 55 prevalence studies. According to the diagnostic criteria of National Institutes of Health, Rotterdam, and AE-PCOS Society, the rates of PCOS prevalence were 6, 10, and 10%, respectively. The range for prevalence on Rotterdam criteria was 8 to 13%. The Rotterdam criteria have four phenotypes (Table 1). The classic phenotype women
present with hyperandrogenism and oligomenorrhea with (A) or without (B) PCO on ultrasound. In the “ovulatory phenotype,” women have hyperandrogenism and PCO (C). In the “non-hyperandrogenic phenotype,” there is oligomenorrhea and PCO, without overt hyperandrogenism (D).14 Prevalence of phenotypes is variable, as this depends greatly on how the population was identified. In an Indian population, among all PCOS women, 56% presented with phenotype A, 15% with phenotype B, 11% with phenotype C, and 18% with phenotype D. Phenotypes A and B were seen more in obese women, with more hyperandrogenemia, insulin resistance, and worse cardiometabolic profile. Metabolic syndrome prevalence was lowest in phenotype D.15 However in other studies, these differences were not as clear. The guideline has emphasized the need for defining phenotypes in research, but the clinical relevance of this remains somewhat unclear at present.

Clinical Features

Reproductive

PCOS is reported as the cause of anovulatory infertility in 70% of women, making it the most common cause of ovulatory dysfunction.16 In a large community-based cohort study, infertility was reported in 72% of women with PCOS compared with 16% in women without PCOS. However, encouragingly, this study also reported that women with PCOS had the same number of children as those without.17

Women with PCOS may also have an increased risk of miscarriage18 and pregnancy complications. A recent meta-analysis found a significantly higher risk of developing gestational diabetes, pregnancy-induced hypertension, preeclampsia, preterm birth, and caesarean section. Metaregression failed to provide evidence of a significant effect between outcome and body mass index (BMI).19 Maternal complications appear to be frequent in women with hyperandrogenic PCOS compared with women with normoandrogenic PCOS.20 Women with PCOS should be informed of the additional risks in pregnancy5 and health practitioners need to be aware of these increased risks.21

PCOS is also associated with an increased risk of endometrial cancer, although it remains unclear whether this risk is independent of other risk factors common in women with PCOS: obesity, diabetes, metabolic syndrome, anovulation.22 A large Danish cohort study of 12,070 women with PCOS reported a fourfold increased risk for endometrial cancer and two- to fourfold increased risks for colon, kidney, and brain tumors. They found no association between PCOS and either breast or ovarian cancer.23 Although the absolute risk of endometrial cancer is still low, health professionals should have a low threshold for excluding endometrial cancer in women with PCOS.5 When persistent thickened endometrium and/or risk factors including prolonged amenorrhea, abnormal vaginal bleeding, or excess weight are present, investigation is indicated using transvaginal ultrasound and/or endometrial biopsy.5

Metabolic Syndrome

PCOS is underpinned by insulin resistance and hyperinsulinemia.3 A recent meta-analysis of clamp assessments of insulin action in PCOS found a decrease in insulin sensitivity of 27% in women with PCOS compared with controls, independent of BMI, age, or diagnostic criteria. BMI exacerbated insulin resistance by 15% in women with PCOS and had a greater impact on insulin resistance in PCOS than in controls.24 PCOS is associated with impaired glucose tolerance (IGT) in up to 30% and type 2 diabetes in up to 10% of women with PCOS.25 When followed up over 10 years, the age-standardized prevalence of diabetes was 39.3% in women with PCOS compared with 5.0% of controls of similar age.26 When investigating differences between phenotypes, Panidis et al reported that in phenotype A, insulin resistance is more prevalent. In women with phenotype C, insulin resistance is not different compared with BMI-matched controls.27 Furthermore, an association between the number of ovarian follicles and insulin resistance was recently described. Ovarian follicle numbers were a significant predictor of insulin resistance in women with PCOS.28 Glycemic status should be assessed (using an oral glucose tolerance test [OGTT], fasting plasma glucose, or HbA1c) at baseline in all women with PCOS and should be repeated every 1 to 3 years depending on other individual risk factors for diabetes present.5 A 75-g OGTT is recommended for women with additional risk factors and preconception and during pregnancy.

Women with PCOS have an adverse cardiovascular risk profile including dyslipidemia, hypertension, and obstructive sleep apnea. A case–control study of 1,550 women with PCOS reported significantly lower levels of HDL and higher levels of total cholesterol, low density lipoprotein, triglycerides, and both systolic and diastolic blood pressure compared with controls, independent of BMI.29 A recent study reported that insulin and androgens may have opposing effects on lipid profiles in women with PCOS, with changes in the production of phospholipids, free fatty acids, and bioactive lipids.30 Studies have also found evidence for endothelial dysfunction in women with PCOS, matched for BMI.31,32 Furthermore, coronary artery calcification is associated with PCOS with odds ratios ranging from 2.3 to 2.4.33,34 The Coronary Artery Risk Development in Young Adults (CARDIA) study found that women with both hyperandrogenemia and oligomenorrhea (phenotype A) had an increased prevalence of coronary artery calcifications and increased carotid intima-media thickness, but women with

Table 1 Polycystic ovary syndrome phenotypes

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<th>Phenotype</th>
<th>Androgen excess</th>
<th>Ovulatory dysfunction</th>
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<tr>
<td>A</td>
<td>Androgen excess</td>
<td>Ovulatory dysfunction</td>
<td>Polycystic ovarian morphology</td>
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<td>B</td>
<td>Androgen excess</td>
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isolated oligomenorrhea or hyperandrogenemia did not. Clinicians should also be aware that it is probable that PCOS precedes and contributes to the development of obstructive sleep apnea. However, it is possible that OSA contributes to the presentation and worsens the clinical manifestations of PCOS. Both conditions are associated with comorbidities including depression, fatigue, hypertension, dyslipidemia, insulin resistance, and IGT. Women with PCOS have increased cardiovascular risk factors but evidence around long-term outcomes on cardiovascular events is limited. A recent sub-group analysis of 106 women with PCOS and 171 control women within a larger prospective cohort study of women through post-menopause (mean time (SD) since menopause 19.85 years (9.94)) reported no increased risk for stroke or ischaemic heart disease compared to the control women. (Meun et al, JCEM, 2018). However, given limited large data sets on long-term data on the morbidity and mortality for cardiovascular disease in PCOS on the background of elevated risk factors it is recommended to offer all women with PCOS a metabolic and cardiovascular assessment measuring weight, height, waist circumference, BMI, blood pressure, fasting lipid profile and monitoring of regular weight changes. If screening identifies CVD risk factors such as obesity, cigarette smoking, dyslipidemia, hypertension, IGT, and lack of physical activity, women should be considered to be at risk of CVD and screening planned as appropriate.

**Psychosocial Health**

There is increasing evidence that women with PCOS are more likely to suffer from mood disorders. The underlying mechanisms are still unclear; however, factors such as obesity, insulin resistance, elevated androgens, and clinical features including distress related to hirsutism, weight gain, acne, and infertility are also drivers.

A recent systematic review focused on moderate and severe symptoms of anxiety and depression. It reported a 4.18 increased odds of moderate or severe symptoms of depression and a 5.62 increased odds of symptoms of anxiety in women with PCOS compared with controls. The increased risk of depression in women with PCOS persisted after adjustment for BMI. Women with PCOS and symptoms of depression had higher mean values of insulin resistance, BMI, hirsutism, and infertility. Mood disturbances, together with other factors including self-esteem, body image, and sexual function, negatively influence quality of life in women with PCOS. The magnitude of psychological consequences of PCOS highlights the need for routine screening for anxiety and depressive symptoms. If the screen is positive, further assessment and/or referral for assessment and treatment should be completed by suitably qualified health professionals.

**Treatment of Nonreproductive Outcomes**

Combined oral contraceptive pills (COCPs) are recommended as first-line medical treatment for the management of hyperandrogenism and regulation of menstrual cycles in women with PCOS. The effects of COCPs are based on estrogen increasing sex hormone binding globulin (SHBG) levels which in turn decrease free testosterone and progesterone, inhibiting luteinizing hormone (LH) secretion resulting in a decline in androgen production and suppression of 5α-reductase activity which reduces conversion of T to DHT. Prolonged use of COCPs is effective in decreasing mFG score, total testosterone, free testosterone, androstenedione, and DHEAS and increasing SHBG. Rates of depression and anxiety have not shown a significant change after treatment with COCPs in PCO; however, studies in general populations have noted an impact of the COCP on libido and mood.

Previous studies have raised concerns of COCPs affecting insulin resistance and triglycerides; however, a recent overview of systematic reviews found no evidence of increased cardiovascular events in prolonged use of COCPs. There is limited information on which COCP preparation type or duration of therapy is optimal for women with PCOS, with all agents reducing SHBG and improving clinical outcomes. Therefore, to minimize complications and based on general population data, recommendations are to use the lowest effective estrogen dose and natural estrogen preparations, balancing efficacy, metabolic risk profile, side effects, costs,
and availability. In combination with COCPs, metformin could be used to avoid weight gain and further suppress hyperandrogenism.

Because of the metabolic features of PCOS such as insulin resistance and hyperinsulinemia, insulin-sensitizing agents, especially metformin, have been used as a treatment option for PCOS for several decades. Metformin reduces insulin levels leading to a reduction of ovarian and adrenal secretions of androgens, pituitary secretion of LH, and increase of SHBG levels. This mechanism may lead to the favorable effect metformin has on BMI, menstrual frequency, systolic blood pressure, fasting glucose levels, fasting insulin levels, testosterone, triglyceride levels, and LH levels. Results, however, do show high levels of heterogeneity. An effect has not been reported on mFG scores, cholesterol levels, or free androgen index. A systematic review and meta-analysis demonstrated that the combination of lifestyle modification and metformin use is associated with lower BMI, subcutaneous adipose tissue, and improved menstruation compared with the combination of lifestyle and placebo. Hence, metformin may be used as an adjuvant to lifestyle modification but not as a substitute for it. When combined with lifestyle intervention, metformin may have an additive effect in improving cardio metabolic risk, especially in high metabolic risk groups. Hence, metformin use is recommended in PCOS for weight and metabolic effects. Commencement at a lower dose is recommended with subsequent titration to minimize mild and self-limiting side effects.

The role of antiandrogens in the treatment of hirsutism in PCOS is controversial; however, there is some low-quality evidence that flutamide and spironolactone reduce hirsutism. A recent study showed a higher reduction in hirsutism when COCPs were used in combination with the antiandrogen bicalutamide compared with COCP alone. In combination with COCPs, antiandrogens should only be considered to treat hirsutism after a minimum of 6 months of COCPs after cosmetic therapy has failed. However, most of these adjuvant medications are used off label and are not registered for the treatment of hyperandrogenism. Bariatric surgery can improve menstrual irregularity, hirsutism, and infertility as reported in a recent systematic review and meta-analysis. The incidence of PCOS decreased by nearly 40% at study endpoint, with nearly 50% improvement of menstrual irregularity and 30% improvement in hirsutism. However, registry studies demonstrate concerns around pregnancy outcomes. For this reason, the guidelines recommend that bariatric surgery can be considered in PCOS after lifestyle therapy fails; however, it should not be recommended for the purpose of improving fertility outcomes pending further research.

### Treatment of Reproductive Outcomes

Interventions in fertility management should be preceded by the optimization of lifestyle and psychological health to improve reproductive and obstetric outcomes. Pharmacological options include ovulation induction with letrozole, clomiphene citrate (CC) metformin, and gonadotrophins. Letrozole is considered first-line treatment for women with PCOS and anovulatory infertility. Letrozole followed by timed intercourse improves live birth rate and clinical pregnancy rates compared with CC with no difference in rates of ovarian hyperstimulation syndrome (OHSS) and reduced multiple pregnancy. Regulations, however, vary internationally on use of letrozole for treatment of infertility and treatment is usually off-label requiring explanation and consent.

CC is recommended for infertility treatment in PCOS both alone and with metformin. Pregnancy and live birth rates are increased, but is associated with increased multiple pregnancy rates (5–8%) and OHSS (<1%). There is also evidence of a benefit from metformin on clinical pregnancy rates and ovulation rates compared with placebo and when combined with CC compared with CC alone. Literature also suggests that metformin is effective in nonobese women. CC could therefore be combined with metformin, rather than persisting with CC alone, especially in women with PCOS who are CC resistant.

Stimulating follicle development and growth with exogenous gonadotrophins for ovulation induction in women with CC-resistant PCOS is well established, and could be used as second-line pharmacological treatment when first-line therapy has failed. Addition of metformin may increase live birth rate among women undergoing ovulation induction with gonadotrophins.

For anovulatory infertility in women who are clomiphene resistant, laparoscopic ovarian drilling is a successful second-line treatment for ovulation induction. It offers several advantages including avoiding OHSS and multiple pregnancies, negating the need for complex monitoring, the favorable effect on physiological ovulatory cycles, and the possible long-term reproductive and endocrine benefits.

In vitro fertilization (IVF) is indicated for women with anovulatory PCOS after failure to respond to ovulation induction or if there are other indications such as tubal damage or male subfertility. A gonadotrophin-releasing hormone antagonist protocol is favored over an agonist protocol because of the reduced OHSS risk. When an agonist protocol is used, metformin could improve clinical pregnancy rate and reduce OHSS risk.

### Summary

Polycystic ovary syndrome is a complex reproductive, metabolic, and psychological condition that affects women's health and quality of life across the life-course. Physicians should be aware of the clinical features and risks for women with PCOS and screen and manage accordingly. Clinicians should focus on lifestyle adjustments as the first-line management to improve reproductive, metabolic, cardiovascular, and psychosocial outcomes focusing on weight management and physical exercise. In addition, pharmacological therapy in the form of COCPs and metformin may be useful. For the management of anovulatory infertility with no other factors, lifestyle intervention is recommended as the first-line management. If this is unsuccessful, ovulation induction using letrozole is the first-line medical management, and after this clomiphene citrate and metformin may have an additive effect. Gonadotrophins and laparoscopic surgery are second line and IVF therapy is the third-line therapy.
where other treatments have failed. The latest international evidence-based guidelines detail these treatments along with providing translation resources for health professional or women with PCOS. Together these are designed to provide a valuable resource aiding clinicians in optimal assessment and management of PCOS.

Funding
H.J.T. and J.A.B. are NHMRC-funded research fellows. 

Conflicts of Interest
The authors have no conflicts of interest to disclose.

References
4 Moran LJ, Hutchison SK, Norman RJ, Teede HJ. Lifestyle changes in women with polycystic ovary syndrome. Cochrane Database Syst Rev 2011;16(2)
11 Priouka A, Farmakiotis D, Katsikis I, Macut D, Gerou S, Panidis D. Anti-Mullerian hormone levels reflect severity of PCOS but are negatively influenced by obesity: relationship with increased luteinizing hormone levels. Am J Physiol Endocrinol Metab 2009;296(02):E238–E243
13 Pankhurst MW, Shorakae S, Rodgers RJ, Teede HJ, Moran LJ. Efficacy of predictive models for polycystic ovary syndrome using serum levels of two antimullerian hormone isoforms (proAMH and AMH_{	ext{NC}}). Fertil Steril 2017;108(05):851–857.e2
28 Hong SH, Sung YA, Hong YS, Jeong K, Chung H, Lee H. Polycystic ovary morphology is associated with insulin resistance in women with polycystic ovary syndrome. Clin Endocrinol (Oxf) 2017;87(04):375–380


Farrell K, Antoni MH. Insulin resistance, obesity, inflammation, and depression in polycystic ovary syndrome: biobehavioral mechanisms and interventions. Fertil Steril 2010;94(05):1565–1574


Glinborg D, Altinok ML, Mumm H, Hermann AP, Ravn P, Andersen M. Body composition is improved during 12 months’ treatment with metformin alone or combined with oral contraceptives compared with treatment with oral contraceptives in polycystic ovary syndrome. J Clin Endocrinol Metab 2014;99(07):2584–2591


Velazquez EM, Mendoza S, Hamer T, Sosa F, Gueck CJ. Metformin therapy in polycystic ovary syndrome reduces hyperinsulinemia, insulin resistance, hyperandrogenemia, and systolic blood pressure, while facilitating normal menses and pregnancy. Metabolism 1994;43(05):647–654


66 Franik S, Eltrop SM, Kremer JAM, Kiesel L, Farquhar C. Aromatase inhibitors (letrozole) for subfertile women with polycystic ovary syndrome. Cochrane Database Syst Rev 2018;5:CD010287