REVIEW



MAHMOODA QURESHI, MD Department of General Internal Medicine, Cleveland Clinic MARJAN ATTARAN, MD Department of Gynecology and Obstetrics, Cleveland Clinic

Review of newer contraceptive agents

BSTRACT

Advances in contraceptive technology have made birth control more effective, convenient, and safe. We review the newer products and some under development, including the latest oral contraceptives, injectable progesterone, subdermal progestin implants, progesterone-releasing IUDs, emergency contraception, and male contraception.

KEY POINTS

Newer oral contraceptives contain much lower doses of estradiol than older preparations and use newer progestins with less androgenic activity. They therefore cause fewer side effects.

Depomedroxyprogesterone acetate injections every 3 months are a good contraceptive option for women in whom compliance may be low.

Progestin implants have a failure rate of 0.8 per 100 woman-years for the first 5 years of use, increasing to 2 per 100 woman-years by the 6th year. The implants should be removed after 5 years.

Progesterone-releasing IUDs reduce the cramping and increased menstrual bleeding that often occur with nonmedicated IUDs.

WO THIRDS of reproductive-age women use some form of contraception, yet 55% of the 6.3 million pregnancies each year in the United States are unplanned, and half of these result in abortion.

Physicians should discuss contraception with all sexually active patients of childbearing age. Safe and effective contraceptive agents are available, but need to be adequately provided to at-risk populations.

In this review we focus on the latest improvements in oral contraceptives, longacting contraceptives such as injectable suspensions and subdermal implants, and medicated intrauterine devices (IUDs). Advances in emergency postcoital contraception are also reviewed.

ORAL CONTRACEPTIVES

Oral contraceptives, the most popular method of reversible contraception in the United States,¹ are highly effective, with a failure rate of 0.3 pregnancies per 100 woman-years of typical use, or 0.1 per 100 woman-years of ideal use.

Mechanism of action

Used singly in high doses, both estrogens and progestins can hinder ovulation but may cause side effects such as breakthrough bleeding, endometrial hyperplasia, acne, and weight gain. Used in combination, they act synergistically and suppress ovulation at much lower doses, with fewer side effects.

These hormones work in several ways. Ethinyl estradiol diminishes follicle-stimulating hormone (FSH) secretion, leading to impaired follicular maturation and insufficient estrogen production via the ovaries. Progestins primarily inhibit luteinizing hormone (LH) secretion and thus ovulation. In addition, progestins thicken the cervical mucus (making it impermeable to sperm) and also cause endometrial atrophy, thereby making implantation improbable.

Improvements in newer preparations

Several refinements have made oral contraceptives much safer and better tolerated than when they were introduced in 1960, while maintaining their high rate of effectiveness:

Lower estrogen doses. Early oral contraceptives contained up to 150 μ g of the estrogen ethinyl estradiol; newer preparations contain only 20 to 50 μ g. The side effects of estrogen increase with the dose and are less common with the lower-dose preparations. On the other hand, breakthrough bleeding is seen more often with the low-dose preparations but may also be due to missing pills or endometritis.

Improved progestins. Progestins are synthetic derivatives of testosterone, manipulated to be highly progestogenic and less androgenic. The progestins used in oral contraceptives have evolved through three generations (TABLE 1) and now have very little androgenic activity.

Phasic preparations. Whereas older oral contraceptives contain the same amount of estrogen and progestin in each tablet and are taken for 21 consecutive days in a 28-day cycle, some newer ones are phasic—ie, they vary the dose of progestin to achieve fewer metabolic side effects. Pills that vary the estrogen dose (Estrostep 21) are also available—in theory, these reduce the occurrence of break-through bleeding. A new formulation (Mircette) containing 10 μ g of ethinyl estradiol during days 22 through 26 of the 28-day pack is now available and should reduce breakthrough ovulation as well as break-through bleeding.

Estrogen-related side effects

Although estrogens can cause nausea, breast tenderness, headache, decreased libido, depression, and cyclic weight gain, their most serious potential side effect is venous thromboembolism. Use of current oral contraceptives is not considered a risk factor for cardiovascular disease or breast cancer.²

Venous thromboembolism. Estrogen alters the synthesis of coagulation factors and

TABLE 1

Progestins used in oral contraceptives

First generation (no longer in use) Ethisterone

Second generation Ethynodiol

Levonorgestrel Norethindrone Norethindrone acetate Norgestrel

Third generation (since 1992) Desogestrel Gestodene (not available in the United States) Norgestimate

fibrinolytic enzymes, making thrombosis more likely. The higher the dose, the greater the risk.³ Use of a second-generation or third-generation oral contraceptive is associated with 3 to 4 thromboembolic events per 10,000 woman-years—more than three times the risk in nonusers, but small in absolute numbers, considerably less than with older agents, and half the risk in pregnant women (TABLE 2).^{4–6}

In view of this risk, oral contraceptives are still contraindicated in patients at higher risk, ie, those with any of the following:

• A history of thromboembolism

• Age greater than 35 and cigarette smoking

• A coagulation disorder. Of these, factor V Leiden mutation is the most common, affecting 3% to 5% of the Caucasian population. In contrast, genetic deficiencies of protein C, protein S, and antithrombin III are rare. Factor V Leiden mutation is associated with an incidence of venous thromboembolism of about 6 events per 10,000 womanyears, but taking an oral contraceptive increases the incidence to 29 events per 10,000 woman-years, suggesting that the two act synergistically to promote coagulation.⁵

Should all patients therefore be screened for factor V Leiden mutation before starting oral contraceptives? This step is *not* deemed necessary, since the absolute risk is still very small. However, patients should be asked about any personal or family history of Screening for factor V Leiden mutation is not deemed necessary for oral contraceptive users

TABLE 2

Incidence of venous thromboembolism with oral contraceptive use

POPULATION AND TYPE OF ORAL CONTRACEPTIVE	INCIDENCE PER 10,000 WOMAN-YEARS
Premenopausal women not taking	0.0
oral contraceptives	0.8
Women taking any combined	
oral contraceptive	3–4
Second-generation progestins	
Monophasic levonorgestrel	2.5
Others (not levonorgestrel)	1.8
Third-generation progestins	
Desogestrel + 30 μ g ethinyl estradiol	4.0
Gestodene	4.4
Pregnant women	6.0
Women with factor V Leiden mutation	
Not taking oral contraceptives	5.7
Taking oral contraceptives	28.5

venous thromboembolism, and those with a positive history should be considered for screening. $^{7}\,$

Taking oral contraceptives has a lower risk of thrombosis than does pregnancy

In 1995, epidemiologic studies reported a higher risk of venous thromboembolism with the newer third-generation oral contraceptives than with the second-generation pills.^{8,9} Subsequent studies showed that the apparent increase can be explained by discrepancies in age and age ranges used for controls, confounding factors, greater use of the older drugs in low-risk pateints (ie, the "healthy user" effect), and bias due to differences in prescription practice and venous thromboembolism diagnosis.6,10,11 The literature suggests that the remaining differences may not be clinically significant.¹⁰ Moreover, we know of no plausible biologic basis for a higher incidence of venous thromboembolism with third-generation oral contraceptives.

Cardiovascular disease. Several newer studies have failed to show an increased risk of acute MI with current low doses.^{12–14} Angiographic studies suggest that users of low-dose oral contraceptives actually have *less* atherosclerosis than nonusers, though no reduction in clinical coronary artery disease has been observed.

Moreover, recent studies have demonstrated that the use of second-generation and third-generation oral contraceptives does not increase the risk of stroke.^{10,15} Smoking and increased age are the main determinants of coronary artery disease and stroke, regardless of oral contraceptive use. Current oral contraceptives do not cause hypertension, which was seen to develop in 5% of women taking high-dose (> 50 μ g ethinyl estradiol) agents.

Breast cancer. In theory, oral contraceptives could induce or help propagate estrogen-sensitive breast cancer, although several large epidemiologic studies have not confirmed this risk.^{16,17} In one of these studies,¹⁶ women who were taking oral contraceptives did seem to have a higher incidence of earlystage breast cancer, but this finding was attributed to better detection and closer health surveillance of women taking oral contraceptives. Investigation in this area continues.

Progestin-related side effects

Progestin side effects such as weight gain, acne, and hirsutism are due to their androgenic properties and are far less common with the new progestins. In fact, some of the newer pills are used to *treat* some of these conditions.

Potential progestin side effects

Dyslipidemia. The third-generation oral contraceptives may have less of an adverse effect on lipid levels than older oral contraceptives. In fact, the newer agents increased HDL cholesterol levels and decreased LDL cholesterol levels and have little impact on total cholesterol levels.¹⁸ The long-term implications of these changes is unknown.

Insulin resistance. Neither the secondgeneration nor third-generation progestins are associated with insulin resistance,¹⁸ which was a concern in the original preparations.

Weight gain. A comparative study of women using oral contraceptives, depomedroxyprogesterone acetate (DMPA, Depo-Provera), and levonorgestrel implants showed no significant weight gain at 1 year, nor a statistically significant difference between the three hormonal methods.¹⁹

Health benefits and noncontraceptive uses

Preventing pregnancy in itself has health benefits—taking oral contraceptives is much safer than pregnancy. In addition, oral contraceptives have certain noncontraceptive benefits and uses, which now account for approximately 20% of prescriptions (TABLES 3 AND 4).

Because the new third-generation progestins have no significant androgenic properties, when used in combination with low-dose estrogen they are useful in treating women with androgen-sensitive conditions such as acne, hirsutism, and polycystic ovarian syndrome. Estrogens stimulate the production of sex hormone-binding globulin, thereby reducing the level of circulating free testosterone. Concurrently, the progestin component decreases luteinizing hormone pulsatility, leading to diminished ovarian stromal production of androgens. The oral contraceptive Ortho-Tri-Cyclen is approved by the Food and Drug Administration (FDA) specifically for the treatment of acne.

Older women who do not smoke can use the low-dose (20 μ g ethinyl estradiol) pills both for contraception and to treat perimenopausal menstrual irregularities.

INJECTABLE PROGESTERONE

DMPA, a synthetic progestin, was approved for contraceptive use in 1992 and is the only injectable progestin available in the United States.²¹

DMPA is very effective, with a failure rate of 0.3 per 100 women per year. Fertility returns once the DMPA clears from the body, which may take several months. By one analysis, 90% of women who stopped DMPA to conceive were pregnant within 24 months.²²

Mechanism of action

DMPA prevents contraception by suppressing the luteinizing hormone surge (thereby inhibiting ovulation), inducing cervical mucus changes, and causing the endometrium to atrophy, making it unreceptive to the blastocyst. Some follicular growth may occur due to the subtotal suppression of follicle-stimulating hormone.

TABLE 3

Noncontraceptive benefits of oral contraceptives

Cancer reduction

Ovarian (risk reduced by 80% at 10 years; benefit continues for > 15 years) Endometrial (risk reduced by 60% at 4 years) Colon (possibly)

Gynecological benefits

Reduced risk of: Ectopic pregnancy Pelvic inflammatory disease Functional ovarian cysts

Menstrual improvements

Reduction in: Anemia Premenstrual syndrome Dysmenorrhea Restoration of regular menses

Other benefits

Reduction in: Benign breast disease Premenopausal bone loss²⁰ Acne Atherosclerosis (possibly)

Indications

DMPA is best suited for women who desire contraception over several months to years and in whom compliance may be poor, such as adolescents and patients with mental illness.

DMPA is particularly useful after abortion. It can also be used during lactation (starting at 6 weeks postpartum) and in situations in which estrogen is contraindicated or its metabolism is altered. It can therefore be used in women with valvular heart disease, diabetes, or hypertension, and in those over age 35 who smoke.

Contraindications to DMPA include liver disease, breast cancer, clotting dyscrasias, and cerebrovascular disease.

Noncontraceptive benefits include a reduction in anemia through reduced menstrual flow, and decreased incidence of pelvic inflammatory disease and endometrial cancer. In addition, women with epilepsy have diminished seizure activity while receiving DMPA.²³ Injectable progesterone is a good option for women with poor compliance

TABLE 4

Newer oral contraceptives: Content and noncontraceptive uses

PREPARATIONS	ESTROGEN DOSE	PROGESTIN DOSE	NONCONTRACEPTIVE USES
Monophasic preparations			
Loestrin 1/20	Ethinyl estradiol 20 µg	Norethindrone 1 mg	Ovarian cysts
Alesse	Ethinyl estradiol 20 µg	Levonorgestrel 0.1 mg	Ovarian cysts, perimenopause, use in teens
Desogen, Ortho-Cept	Ethinyl estradiol 30 µg	Desogestrel 0.15 mg	Acne, hirsutism, ovarian cysts
Triphasic preparations			
Ortho Tri-Cyclen	Ethinyl estradiol 35 μg	Norgestimate 0.180 mg (week 1) 0.125 mg (week 2) 0.250 mg (week 3)	Acne, hirsutism
Estrophasic preparations			
Estrostep 21	Ethinyl estradiol 20 μg (week 1) 30 μg (week 2) 35 μg (week 3)	Norethindrone 1 mg	Irregular menses Early cycle breakthrough bleeding
Mircette	Ethinyl estradiol 20 μg (days 1–21) 10 μg (days 22–26)	Desogestrel 0.15 mg	Midcycle bleeding Breakthrough bleeding Estrogen withdrawal headaches

Side effects

Overall, DMPA is safe, has few side effects (namely, irregular menses, headache, gastrointestinal upset, dizziness, and fatigue), and may also cause lower bone mineral density and dyslipidemia.

Thirty percent of women develop irregular menses and spotting during the first 3 months of DMPA therapy. Persistent bleeding can be treated by giving the next dose early or by adding an estrogen such as 1.25 mg of conjugated estrogen for 7 days. Amenorrhea develops in 50% of women at 1 year and 80% at 3 years of use.

Lower bone mineral density was observed in women who used DMPA for a minimum of 5 years than in other premenopausal women, but levels were higher than in postmenopausal controls.^{24,25} The implication for subsequent fracture rates is unknown. Further studies are required.

Although lipid metabolism appears to be influenced adversely, these effects may be

transient and the long-term impact is unknown.^{21,26} The effects on bone and lipids are probably due to lower circulating estradiol levels in women receiving DMPA.

There is no demonstrated increased risk of cervical, ovarian, or breast cancer with DMPA use, and there is no association with hypertension or myocardial infarction.²¹ Studies show no link between weight gain or mood disorders and use of DMPA.²⁶

Dosage

DMPA is given as a 150-mg intramuscular injection every 3 months, ideally within 5 days of onset of menstruation.

Other preparations

Monthly injections of an estrogen-progestin combination are in use outside the United States.²⁷ Combinations of estrogen and a progestin such as medroxyprogesterone acetate or norethindrone lead to monthly withdrawal bleeding and are very efficacious.

SUBDERMAL PROGESTIN IMPLANTS

Levonorgestrel implants (Norplant) have been available in the United States since 1990, although clinical trials were performed as early as 1972 in Chile.

Levonorgestrel implants are effective: the failure rate is 0.8 per 100 woman-years averaged over 5 years, increasing to 2 per 100 woman-years by the 6th year. Thus, after 5 years the implants should be removed. Progestin levels are negligible after removal and fertility returns immediately.

Mechanism of action

Implantable progestins prevent conception primarily by rendering the cervical mucus impenetrable to sperm. They also inhibit ovulation and impair oocyte maturation by blocking the LH surge. However, LH blockade is inconsistent; hence, cyclic luteal activity may be seen, albeit with subphysiologic progesterone levels. Although ovulation may still occur, impaired oocyte maturation and luteal insufficiency prevent pregnancy.

Indications

Progestin implants are best suited for women desiring long-term reversible contraception.

Side effects

Bleeding patterns are variable due to endometrial atrophy and variable cyclic luteal activity. Amenorrhea develops in 5% to 10% of users. Irregular bleeding occurs most frequently in the first year and is usually due to endometrial atrophy. The bleeding can be treated effectively with conjugated equine estrogens 1.25 mg/day for 1 to 2 weeks.

Other side effects account for 14% of removals and include headache (most common), mood changes, local dermatitis, acne, mastalgia, and hair changes.²⁸ Functional ovarian cysts can occur but are usually asymptomatic and are treated expectantly. Levonorgestrel implants are safe to use during lactation and there has been no evidence of teratogenesis in women who conceived while on this agent. There is no clinically significant effect on glucose or lipid metabolism.²⁹ The long-term effect on cancers of the reproductive tract is not yet known.

Dosage and administration

Six rods, each containing 36 mg of levonorgestrel in a Silastic adhesive, are inserted subcutaneously in the upper arm, ideally within 7 days of menstruation. The procedure is simple and can be performed in the office under local anesthetic. However, removal can be cumbersome due to local fibrosis.

New preparations

Norplant-2, a two-rod version of levonorgestrel in a different elastomer, is easier to insert and remove than Norplant and is now available in the United States.³⁰ A single-rod implant (not yet available) containing the third-generation progestin desogestrel is being studied and appears very effective.³¹ New implant systems under investigation use biodegradable materials such as cholesterol and elastomers such as poly E-caprolactone, which is easier to remove.²⁸

PROGESTERONE-RELEASING IUD

The intrauterine device (IUD) is the most commonly used reversible method of contraception in the world.

Progesterone-releasing IUDs were developed to reduce the cramping and increased menstrual bleeding that often occur with nonmedicated IUDs. However, they also have the benefit of an additional mechanism of action: Whereas nonmedicated IUDs prevent conception by producing a sterile intrauterine inflammatory response that is spermicidal, progesterone-releasing IUDs also induce decidualization of the endometrium, inhibiting implantation.³²

Indications

Progesterone-releasing IUDs are best suited for long-term contraception in parous or older women.

Side effects

Amenorrhea or oligomenorrhea is common, due to suppression and atrophy of the endometrium. For a patient suffering from menorrhagia and anemia this may be a welcome side effect. Nevertheless 72% of all removals for menstrual irregularities were due Levonorgestrel IUDs last up to 7 years and should be available in the US soon

CONTRACEPTIVES QURESHI AND ATTARAN

TABLE 5

Oral regimens for postcoital contraception

Estrogen-progestin (Yuzpe regimen)

Ethinyl estradiol 100 µg plus levonorgestrel 0.5 mg, repeated in 12 hours*

Estrogen-progestin (Preven) Ethinyl estradiol 50 µg plus levonorgestrel 0.25 mg, 2 pills, repeated in 12 hours

High-dose estrogen

Ethinyl estradiol 2.5 mg twice daily for 5 days

Levonorgestrel

0.75 mg repeated in 12 hours

Mifepristone (RU486)

600 mg single dose

*Typically, two tablets of Ovral (50 μg ethinyl estradiol plus 0.5 mg norgestrel), or four of Lo/Ovral (30 μg ethinyl estradiol plus 0.3 mg norgestrel) are used to provide similar doses to the Yuzpe regimen; any other combined oral contraceptive may also be used

For emergency contraception, give two doses of estradiol 100 µg plus levonorgestrel 0.5 mg, 12 hours apart to amenorrhea.³³ When patients are counseled appropriately about what to expect regarding bleeding patterns, the number of IUD removals decreases.

The absolute rates of ectopic pregnancy are very low with progesterone-releasing IUDs. Ectopic pregnancy rates are higher with one type of medicated IUD (Progestasert), but lower with the levonorgestrel-containing IUD,^{2,34} compared with rates in women who don't use contraception.

No changes in HDL-cholesterol or coagulation parameters have been observed with the progesterone-releasing IUD.³⁵

Dosage and administration

Progestasert, the only hormone-releasing IUD available in the United States at present, contains 38 mg of progesterone. It must be reinserted every year.

A levonorgestrel IUD, not yet available in the United States but being evaluated for release soon, releases 20 μ g of levonorgestrel per day, thereby suppressing ovulation in 55% of menstrual cycles.³² The levonorgestrel IUD provides contraception for up to 7 years.²

POSTCOITAL (EMERGENCY) CONTRACEPTION

Postcoital (emergency) contraception refers to any method used to prevent pregnancy after unprotected intercourse.

In a recent survey, only 36% of Americans had any knowledge about emergency contraception and only 1% had ever used it.³⁶ The study also showed that, although aware of its existence, very few gynecologists actually prescribe the regimen. Not until February 1997 did the FDA declare the administration of synthetic estrogen and progesterone in oral contraceptives to be an effective and safe method of emergency contraception. Guidelines from the World Health Organization state that the only absolute contraindication to this method is known pregnancy.³⁷

Postcoital contraception can be achieved via a variety of methods: high doses of estrogen, combinations estrogen and progesterone, progestin alone, mifepristone (RU486), and IUD insertion in the immediate postcoital period (TABLE 5).³⁷

How these agents act is not always clear; some may act at different levels. Estrogen and progestin, either alone or concurrently, act primarily by preventing or delaying ovulation.

Combined oral contraceptives. The most commonly used forms of emergency contraception are combined oral contraceptives. Yuzpe and Lancee³⁸ first reported the use of ethinyl estradiol and norgestrel as a postcoital contraceptive in 1977. Two doses of 100 μ g of ethinyl estradiol and 0.5 mg of levonorgestrel are given 12 hours apart, within 72 hours of intercourse. The main side effects of this regimen are nausea and vomiting. Regular birth control pills that contain the same hormones are generally used, but a product specifically licensed by the FDA for emergency contraception was released in September 1998. This product, the Preven Kit, contains a urine pregnancy test to exclude pregnancy prior to taking the hormones.

Progestin-only compounds have been shown to be as effective as the Yuzpe regimen.³⁹ Two doses of levonorgestrel 0.75 mg can be given 12 hours apart, within 48 to 72 hours of unprotected intercourse. This regimen may have a lower incidence of side effects than the Yuzpe regimen.³⁹ In several countries, four tablets of 0.75 mg levonorgestrel are marketed as a postcoital contraceptive agent. Women who require postcoital contraception should be counseled and started on regular contraception.

The copper IUD can both diminish fertilization through a toxic effect on sperm, and impair implantation by causing changes in the endometrium.²

Mifepristone (RU486) is an antiprogesterone that can inhibit ovulation, but if given in the midluteal phase can cause regression of the corpus luteum in 50% of women.⁴⁰ When given as a single 600-mg dose, mifepristone is a very effective method of postcoital contraception. However, since it may delay endometrial maturation, the resulting delay in onset of menstruation can be anxiety-provoking for the patient, even though it effectively prevents pregnancy. Its postcoital use is separate from its use as an early abortifacient. Although widely available in Europe, mifepristone is investigational in the United States, but is being evaluated for emergency contraceptive use for approval in 2001.

MALE CONTRACEPTION

Condoms are the mainstay of male contraception. Advances include the new polyurethane condom, which is thinner, stronger, and less allergenic than the older latex condoms. While male sterilization or vasectomy is a contraceptive choice for some men, it is considered permanent. Reversible vasectomy is being investigated, whereby a polyurethane elastomer plugs the vas deferens but can be removed within 4 years.⁴¹

A variety of male contraceptives are being developed.41 Hormonal methods for terminating sperm production are under investigation, but will probably not be available for another decade. An injection of an androgen and a progestin given every 3 months is under study by the World Health Organization. The androgen suppresses production of gonadotropinreleasing hormone (GnRH), reducing LH and FSH, and therefore inhibiting spermatogenesis. The progestin allows a lower dose of androgen to be used, thereby reducing side effects. Also under development are GnRH receptor antagonists that would block spermatogenesis. Because testosterone production would also decrease, androgen replacement would be required. Short-term contraceptive vaccines for men and women are also being investigated.

REFERENCES

- Trends in oral contraceptive development and utilization. The Contraceptive Report. 1997; 7(5):4–14.
- Speroff L, Glass RH, Kase NG, editors. Clinical gynecologic endocrinology and infertility. Fifth edition. Baltimore: Williams and Wilkins, 1994.
- Norris LA, Bonnar J. The effect of estrogen dose and progestin type on hemostatic changes in women taking low dose oral contraceptives. Br J Obst Gynaecol 1996; 103:261–267.
- Gerstman BB, Piper JM, Tomita DK, Ferguson WJ, Stadel BV, Lundin FE. Oral contraceptive dose and risk of deep venous thrombosis. Am J Epidemiol 1991; 133:32–37.
- Vandenbroucke JP, Koster T, Briet E, Reitsma PH, Bertina RM, Rosendaal FR. Increased risk of venous thrombosis in oral-contraceptive users who are carriers of factor V Leiden mutation. Lancet 1994; 344:1453–1457.
- Farmer RDT, Lawrenson RA, Thompson CR, Kennedy JG, Hambleton IR. Population-based study of risk of venous thromboembolism associated with various oral contraceptives. Lancet 1997; 349:83–88.
- Vandenbroucke JP, van der Meer FJM, Helmerhorst FM, Rosendaal FR. Factor V Leiden: should we screen oral contraceptive users and pregnant women? BMJ 1996; 313:1127–1130.
- Jick H, Jick SS, Gurewich V, Myers MW, Vasilakis C. Risk of idiopathic cardiovascular disease and nonfatal venous thromboembolism in women using oral contraceptives with differing progestin components. Lancet 1995; 346:1589–1593.

- WHO collaborative study of cardiovascular disease and steroid hormone contraception. Effect of different progestins in low oestrogen oral contraceptives on venous thromboembolic disease. Lancet 1995; 346:1582–1588.
- Spitzer WO. The 1995 pill scare revisited: anatomy of a nonepidemic. Hum Reprod 1997; 12:2347–2357.
- Suissa S, Blais L, Spitzer WO, Cusson J, Lewis M, Heinemann L. Firsttime use of newer oral contraceptives and the risk of venous thromboembolism. Contraception 1997; 56:141–146.
- Stampfer MJ, Willett WC, Colditz GA, Speizer FE, Hennekens CH. A prospective study of past use of oral contraceptive agents and risk of cardiovascular disease. N Engl J Med 1988; 319:1313–1317.
- Lewis M, Spitzer WO, Heineman LAJ, MacRae KD, Bruppacher R, Thorogood M on behalf of the Transnational Research Group on Oral Contraceptives and the Health of Young Women. Third-generation oral contraceptives and risk of myocardial infarction: an international case-control study. BMJ 1996; 312:88–90.
- Rosenberg L, Palmer JR, Sands MI, et al. Modern oral contraceptives and cardiovascular disease. Am J Obstet Gynecol 1997; 177:707–715.
- Petitti DB, Sidney S, Bernstein A, Wolf S, Quesenberry C, Ziel HK. Stroke in users of low-dose oral contraceptives. N Engl J Med 1996; 335:8–15.
- Collaborative Group on Hormonal Factors in Breast Cancer. Breast cancer and hormonal contraceptives: collaborative reanalysis of individual data on 53 287 women with breast cancer and 100 239 women without breast cancer from 54 epidemiologic studies. Lancet 1996; 347:1713–1727.

- Rossling MA, Stanford JL, Weiss NS, Habel LA. Oral contraceptive use and risk of breast cancer in middle-aged women. Am J Epidimiol 1996; 144:161–164.
- Godsland IF, Crook D, Simpson R, et al. The effects of different formulations of oral contraceptive agents on lipids and carbohydrate metabolism. N Engl J Med 1990; 323:1375–1381.
- Moore LL, Valuck R, McDougall C, Fink W. A comparative study of one-year weight gain among users of medroxyprogesterone acetate, levonorgestrel implants, and oral contraceptives. Contraception 1995; 52:215–219.
- Recker RR, Davies KM, Hinders SM, Heaney RP, Stegman MR, Kimmel DB. Bone gain in young adult women. JAMA 1992; 268:2403–2408.
- Kaunitz AM, Rosenfield A. Injectable contraception with depot medroxyprogesterone acetate: current status. Drugs 1993; 45:857–865.
- 22. Pardthaisong T. Return of fertility after use of the injectable contraceptive Depo-Provera: updated analysis. J Biosoc Sci 1984; 16:23–34.
- Mattson RH, Cramer JA, Caldwell BVD, Sinconolfi BC. Treatment of seizures with medroxyprogesterone acetate: preliminary report. Neurology 1984; 34:1255–1258.
- Cundy T, Reid OR, Roberts H. Bone density in women receiving depot medroxyprogesterone acetate for contraception. BMJ 1991; 303:13–16.
- 25. Edwards CP, Hertweck SP, Perlman SE, Goldsmith LJ, Sanfilippo JS. A prospective study evaluating the effects of Depo Provera on bone mineral density in adolescent females: a preliminary report [abstract]. Presented at the 12th annual clinical meeting of the North American Society for Pediatric and Adolescent Gynecology; May 28 1998, Palm Beach, FL.
- Westhoff C. Depot medroxyprogesterone acetate contraception: Metabolic parameters and mood changes. J Reprod Med 1996; 41:401–406.
- 27. **Guo-wei S.** Pharmacodynamics effects of once a month combined injectable contraceptives. Contraception 1994; 49:361–385.
- Darney PD. Hormonal implants: contraception for a new century. Am J Obstet Gynecol 1994; 170:1536–1543.
- Kaunitz AM, Illions EH, Jones JL, Sang LA. Contraception. Med Clin North Am 1995; 79(6):1377–1408.

ANNOUNCING

The Alpha One Foundation Research Registry for individuals diagnosed with Alpha,-Antitrypsin Deficiency

Enroll your patients in the Alpha One Research Registry, the only one of its type in the United States.

Patients and their physician will have an opportunity to participate in clinical trials and research protocols to develop new therapies.



All data is confidential and managed by the University of Miami School of Medicine Department of Epidemiology and Public Health

> For Patient Enrollment Call 1-888-825-7421 Ext.6

Visit The Alpha One Foundation on the Web at www.alphaone.org or E-mail us at registry @ alphaone.org

- Gao J, Wang SL, Wu S, Sun B, Hannu A, Tapani L. Comparison of clinical performance, contraceptive efficacy and acceptability of levonorgestrel-releasing IUD and Norplant-2. Contraception 1990; 41:485–494.
- 31. Davies GC, Feng LX, Newton JR. Release characteristics, ovarian activity and menstrual bleeding pattern with a single contraceptive implant releasing 3-ketodesogestrel. Contraception 1993; 47:251–261.
- Chi IC. The progestin-only pills and the levonorgestrel-releasing IUD: Two progestin-only contraceptives. Clin Obstet Gynecol 1995; 38:872–889.
- Sivin I, Stern J. Health during prolonged use of levonorgestrel 20 μg/day and the Copper TCu 380 Ag intrauterine contraceptive devices: a multicenter study. Fertil Steril 1994; 61:70–77.
- Sivin I. Dose- and age-dependent ectopic pregnancy risks with intrauterine contraception. Obstet Gynecol 1991; 78:291–298.
- Nilsson CG. Two-year experience with two levonorgestrel IUDs and one copper releasing IUD: a randomized comparative performance study. Fertil Steril 1983; 39:187–192.
- Delbanco S, Mauldon J, Smith MD. Little knowledge and limited practice: Emergency contraceptive pills, the public and the obstetrician gynecologist. Obstet Gynecol 1997; 89:1006–1011.
- Glasier A. Emergency postcoital contraception. N Engl J Med 1997; 337:1058–1064.
- Yuzpe AA, Lancee AA. Ethinylestradiol and di-norgestrel as a postcoital contraceptive. Fertil Steril 1977; 28:932–936.
- Ho PC, Kwan MSW. A prospective randomized comparison of levonorgestrel with Yuzpe regimen in postcoital contraception. Hum Reprod 1993; 8:389–392.
- Swahn ML. The effect of RU486 administered during the proliferative and secretory phase of the bleeding pattern, hormonal parameters, and the endometrium. Hum Reprod 1988; 3:915–921.
- 41. Alexander NJ. Future contraceptives. Sci Am 1995; 273(3)Sept:136-141.

ADDRESS: Mahmooda Qureshi, MD, Department of General Internal Medicine, E13, The Cleveland Clinic Foundation, 9500 Euclid Avenue, Cleveland, OH 44195.

We Welcome Your Letters

WE ENCOURAGE YOU TO WRITE, either to respond to an article published in the *Journal* or to address a clinical issue of importance to you. You may submit letters by mail, fax, or e-mail.

MAILING ADDRESS

Letters to the Editor Cleveland Clinic Journal of Medicine 9500 Euclid Ave., NA32 Cleveland, OH 44195

FAX

216.444.9385

E-MAIL ccjm@ccf.org

Please be sure to include your full address, phone number, fax number, and e-mail address. Please write concisely, as space is limited. Letters may be edited for style and length. We cannot return materials sent. Submission of a letter constitutes permission for the *Cleveland Clinic Journal of Medicine* to publish it in various editions and forms.