The levonorgestrel-releasing intrauterine system: Safety, efficacy, and patient acceptability

Megan N Beatty
Paul D Blumenthal
Department of Obstetrics and Gynecology, Stanford University, Stanford, CA, USA

Abstract: The levonorgestrel-releasing intrauterine system (LNG-IUS) is a safe, effective and acceptable form of contraception used by over 150 million women worldwide. It also has a variety of noncontraceptive benefits including treatment for menorrhagia, endometriosis, and endometrial hyperplasia. The LNG-IUS has also been used in combination with estrogen for hormone replacement therapy and as an alternative to hysterectomy. Overall, the system is very well tolerated and patient satisfaction is quite high when proper education regarding possible side effects is provided. However, despite all of the obvious benefits of the LNG-IUS, utilization rates remain quite low in the developed countries, especially in the United States. This is thought to be largely secondary to the persistent negative impressions from the Dalkon Shield intrauterine experience in the 1970s. This history continues to negatively influence the opinions of both patients and health care providers with regards to intrauterine devices. Providers should resolve to educate themselves and their patients on the current indications and uses for this device, as it, and intrauterine contraception in general, remains a largely underutilized approach to a variety of women’s health issues.

Keywords: Mirena®, levonorgestrel-releasing, intrauterine system, intrauterine contraceptive device

Introduction

Worldwide, intrauterine devices (IUDs) are the most widely used and effective reversible contraceptive method and are safe, extremely “low maintenance” contraceptives. They are second only to female sterilization as the most prevalent method of family planning worldwide (13.6% vs 20.5%). Over 150 million women worldwide use the IUD for contraception. However, for a variety of reasons, including the continued concern many women have about the morbidity and mortality associated with IUDs resulting (whether they know it or not) from the Dalkon shield experience of the 1970s, there is a large disparity between the numbers of users in developing versus developed countries. There is an especially low utilization rate within the United States. Nearly 15% of women who use contraception in less developed countries choose the IUD, compared with only 8% of women in developed countries and only 2% in the United States. Two thirds of worldwide IUD use (over 100 million women) is represented by China alone.

There is currently a wide variety of IUDs available worldwide; some are inert, some are copper containing, and some are medicated with levonorgestrel or indomethacin. They also come in a wide variety of sizes and shapes including T-shaped and “frameless” devices. This article will focus specifically on the LNG-IUS, exploring...
mechanism of action, safety, efficacy, variety of uses, and patient acceptability.

**Mechanism of action of the levonorgestrel-releasing intrauterine system**

The levonorgestrel-releasing intrauterine system (LNG-IUS) has been available in Europe since 1990 and the United States since 2000. It is marketed under the name Mirena® (Bayer Schering Pharma, Berlin, Germany). The LNG-IUS provides highly effective contraception for up to five years, with potential for approval for up to seven years in the near future. The mechanisms of action of the LNG-IUS are similar to that of levonorgestrel implants or levonorgestrel-containing mini-pills, although it accomplishes these effects with much lower peak serum levels than other progestin-containing contraceptives (0.1–0.4 ng/ml vs 1.7–15.2 ng/ml with combined and progestin-only oral contraceptives, respectively, and 5.4 ng/ml for combined vaginal preparations).

The LNG-IUS is a T-shaped device composed of a cylinder containing 52 mg of LNG covered by a rate-controlling membrane which serves to regulate the rate of hormonal release (Figure 1). Initially, 20 µg of levonorgestrel is released every 24 hours from this polymer cylinder. This decreases to 11 µg every 24 hours by the end of five years, with an average release rate of 14 µg per day over the life of the IUS. Levonorgestrel, a highly potent second generation progestin, thickens cervical mucus and suppresses endometrial proliferation (preventing decidualization of the stroma). This creates a hostile environment for sperm survival, inhibiting motility and capacitation with the net effect combining to prevent fertilization.

The LNG-IUS also produces endometrial thinning with fragile superficial vessels which, in the unlikely event of fertilization, may prevent implantation. The low serum levels of absorbed progestin are below the threshold for inhibition of ovulation, so that most women with the LNG-IUS continue to ovulate regularly.

As a result of these various contraceptive actions, the efficacy rate of the LNG-IUS is high, with only 0.1% of women experiencing an unintended pregnancy within the first year of typical use. In four clinical studies, representing more than 10,000 woman-years of use, the average Pearl Index (a statistical estimation of the number of unintended pregnancies in 100 woman-years of exposure to a contraceptive method) was 0.1.

**Candidates and contraindications for intrauterine contraception and the LNG-IUS**

**Candidates and contraindications for intrauterine devices in general**

Intrauterine contraceptive devices are appropriate contraceptive options for women who desire a very convenient, long-term contraceptive or who are considering sterilization. While there are contraindications to utilization of an IUD, they are relatively few compared to some other methods. Some contraindications, for example pregnancy, are absolute and undisputed. However, other contraindications, such as patients at increased risk of sexually transmitted infections or nulliparous women, are more controversial. The recommendations regarding these issues vary based on which specific organization is consulted (Table 1).

One of the most contentious contraindications to IUD use relates to patients who are perceived to be at increased risk for sexually transmitted infections (STIs). This concern likely dates back to the experience with the Dalkon Shield in the 1970s. The Dalkon Shield IUD was associated with an unacceptably high risk of pelvic inflammatory disease, tubal infertility and septic abortion, to a significant extent secondary to its braided polyfilament tail. It was subsequently removed from the market, and is thought by many to be responsible for the initial significant decline and continued slow increase in numbers of patients interested in using IUDs for contraception. After the
Table 1 Contraindications to the levonorgestrel-releasing intrauterine system according to organizational recommendations

<table>
<thead>
<tr>
<th>Condition</th>
<th>ACOG</th>
<th>WHO (for the definition of risk categories see Appendix B)</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uterine anomaly (including fibroids causing distortion of the uterine cavity)</td>
<td>Contraindicated</td>
<td>Risk category 4</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>History of pelvic inflammatory disease (past three months only)</td>
<td>Contraindicated</td>
<td>No contraindication</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Post-partum endometritis or septic abortion in the last three months</td>
<td>Contraindicated</td>
<td>Immediate insertion only, risk category not defined</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Active cervicitis/vaginitis</td>
<td>Contraindicated</td>
<td>Initiation of LNG-IUS is risk category 4, continuation is risk category 2 for cervicitis; both are risk category 2 for vaginitis</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Immunosuppression (eg, leukemia, AIDS, intravenous drug abuse)</td>
<td>No recommendation</td>
<td>Initiation is risk category 3, continuation category 2. If AIDS patient is clinically well on antiretroviral therapy, initiation is also category 2. Not contraindicated in HIV.</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Known or suspected cervical dysplasia/genital bleeding of unknown etiology</td>
<td>Contraindicated</td>
<td>Initiation is risk category 4, continuation is risk category 2</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Known or suspected breast carcinoma</td>
<td>Contraindicated</td>
<td>Risk category 4 for current, category 2 for past with no evidence of disease for the last five years</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>Contraindicated</td>
<td>Risk category 4</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>Post-partum &lt;48 hours (including insertion immediately after delivery of the placenta)</td>
<td>Contraindicated</td>
<td>Risk category 4 if not breast feeding, category 3 if breast feeding</td>
<td>No recommendation</td>
</tr>
<tr>
<td>Post-partum ≥48 hours to &lt;4 weeks</td>
<td>No recommendation</td>
<td>Risk category 3</td>
<td>No recommendation</td>
</tr>
<tr>
<td>Active viral hepatitis, liver tumor (benign or malignant), cirrhosis</td>
<td>Not recommended for “current liver disease”</td>
<td>Risk category 3 for malignant hepatoma, benign hepatocellular adenoma, or decompensated cirrhosis, category 2 for focal nodular hyperplasia</td>
<td>Contraindicated</td>
</tr>
<tr>
<td>History of, or condition that predisposes to, ectopic pregnancy</td>
<td>No contraindication</td>
<td>No contraindication</td>
<td>Recommends caution</td>
</tr>
<tr>
<td>Current DVT/PE</td>
<td>No recommendation</td>
<td>Risk category 3, category 2 if established on anticoagulant therapy</td>
<td>No recommendation</td>
</tr>
</tbody>
</table>

Abbreviations: ACOG, American College of Obstetricians and Gynecologists; AIDS, acquired immunodeficiency syndrome; DVT, deep vein thrombosis; HIV, human immunodeficiency virus; PE, pulmonary embolism; WHO, World Health Organization.

dismal experience with the Dalkon Shield physicians understandably were, and despite current reassuring data often continue to be, hesitant to prescribe this for unmarried women or patients with multiple sexual partners because of the persistent concern over serious pelvic infections and implications for future fertility. However, numerous studies on the currently available versions of the IUD have shown that, overall, IUD users are no more
likely to suffer from PID than control women. One model estimated that the risk of clinical PID secondary to IUD use is only 0.15%.6 Another recent study evaluating a high-risk population in West Africa showed the risk of PID related to IUD insertion to be only 0.075%.7 One randomized trial (with data collection in the early 1980s) showed a decreased rate of PID among LNG-IUS users compared to users of a copper-containing IUD.8 The IUD may, therefore, be a reasonable option for women who may be at risk for sexually transmitted infections. This is especially the case where prevalence in the community as a whole may be high but individual risk may be low (ie, no current or recent history of an STI or the patient uses condoms regularly if she has more than one partner). Morrison and colleagues developed a checklist to help with risk assessment for STIs in candidates for IUD insertion which may be helpful with management decisions (Appendix A).9 Of note, all women must still be counseled on safe sex practices, and condom use must be encouraged.

The appropriateness of IUD use in nulliparous women has also been controversial because of a variety of concerns specific to this group. First, because of the smaller size of the uterus it was thought that there might be an increased expulsion rate. In addition, because of the more stenotic nulliparous os, the device may cause increased pain with insertion. There has also been concern over a possible increased rate of infertility in these women, again secondary to the concern over increased rates of PID with the IUD and the thought that nulliparous women are less likely to be in stable, monogamous relationships. However, these concerns are not supported by recent studies. A retrospective cohort study of 129 nulliparous and 332 parous women showed that rates of expulsion for the levonorgestrel-releasing IUDs were 0% to 0.2% per year, and that nulliparous women did not show more complications than parous women.10 A comparative cross-sectional study of 227 nulliparous women and 2,080 parous women showed similar reasons for removal between the two groups: 49.1% vs 48.2% because of device expiration, 21.7% vs 15.4% desired planned pregnancy, 4.8% vs 6.3% were secondary to accidental pregnancy, 1.6% vs 1.1% because of PID. There were actually decreased rates of expulsion in the nulliparous group (1.6% vs 5.1%), and slightly increased rates of pain/bleeding (14.5% vs 10.2%).11 According to the medical eligibility criteria for contraceptive use developed by the World Health Organization (WHO), nulliparity is a condition for which the advantages of using intrauterine contraception generally outweigh the theoretical or proven risks (risk category 2. For the full definition of WHO risk categories please see Appendix B).12 The intrauterine device is, therefore, generally considered to be a safe and effective contraceptive which should be offered to nulliparous women as part of their family planning counseling.

There are no WHO restrictions for using IUDs among women who have uncomplicated valvular disease. When valvular heart disease is complicated by pulmonary hypertension, atrial fibrillation, or subacute bacterial endocarditis, prophylactic antibiotics should be administered to women with these conditions before an IUD is inserted to prevent endocarditis.

Candidates and contraindications for the LNG-IUS

The WHO provides highly useful guidelines regarding the appropriateness of the LNG-IUS for women with a variety of medical conditions (see Appendix B for the definition of WHO risk categories for eligibility for use of an intrauterine device). According to the WHO, women who have any degree of hypertension can use a LNG-IUS if no other risk factors for cardiovascular disease are present. When multiple risk factors do exist, the risk of cardiovascular disease may increase substantially. For this reason, because progestins have been shown in clinical trials to influence lipid metabolism,13 there has been concern, albeit theoretical, about the effect of levonorgestrel-releasing IUDs on lipid levels in women with a history of heart disease or stroke.14 However, these concerns about the effect of the LNG-IUS on lipid profiles have not been borne out. A recent study of 48 patients who had the LNG-IUS inserted for menorrhagia showed no significant change from baseline in mean total cholesterol, triglycerides, low-density lipoprotein (LDL) or very low density lipoprotein (VLDL) at the end of one year.15 Another study of 92 women with the LNG-IUS inserted for menorrhagia showed a marginally significant reduction in total cholesterol from baseline, a reduction in HDL at 6 months which reverted to baseline by one year, while triglycerides, LDL, apolipoprotein A1 and apolipoprotein B remained stable over the course of 18 months.16 Therefore, though the WHO recommendations warn of a theoretical effect of the LNG-IUS on lipid levels and caution its use in women with multiple cardiac risk factors, studies do not show this to be a significant problem associated with the LNG-IUS.

Progestins are thought to increase the risk for deep vein thromboses (DVTs) and pulmonary emboli, although this increase is substantially less with a levonorgestrel-releasing IUS than with combined oral contraceptives. Since risks associated with the LNG-IUS are even greater for women
who have a current DVT, the WHO states that the risks of using the LNG-IUS in such a situation usually outweigh the benefits (risk category 3) until they are established on anticoagulant therapy (risk category 2).\(^{12,16}\)

Among women who are infected with the human immunodeficiency virus (HIV) or are at risk of becoming infected with this pathogen, there is no known interaction between antiretroviral therapy and LNG-IUS use. The WHO, however, classifies insertion of a LNG-IUS in a woman with acquired immunodeficiency syndrome (AIDS) as a category 3 risk and continuation of the IUD as a category 2 risk unless the patient is clinically well and on antiretroviral therapy. In this case, insertion of an IUS is then considered to provide benefits that generally outweigh any theoretical or proven risk. Although IUD use is not associated with an increased risk of HIV transmission to sexual partners, these women should be counseled about safe sex practices and condom use must be encouraged.\(^{12}\)

**Insertion of the LNG-IUS**

Although the intrauterine contraceptive device should ideally be inserted within the first seven days following a woman’s menstrual cycle, it can be inserted at any time during the cycle if it is reasonably certain that she is not pregnant and has not been at risk of pregnancy during that cycle. As discussed in more detail later (please see “Efficacy of the LNG-IUS”), an IUD may be safely inserted immediately after uncomplicated spontaneous or induced abortions, including those that are performed during the second trimester. Expulsion, pregnancy, and removal rates after first-trimester termination are comparable to those found in studies in which the device was inserted during or immediately after menses. The shorter the period of gestation at the time of spontaneous or induced abortion, the lower the event rates following IUD insertion. The LNG-IUS can also be safely inserted immediately after childbirth, but there are associated risks with insertion during the immediate postpartum period. If an IUD is inserted within the first 48 hours after childbirth, the risk of expulsion is greater than interval insertion, although the risk of expulsion is somewhat lower if the IUD is inserted within 10 minutes of the delivery of the placenta. If an IUD is inserted 48 hours and up to four weeks after childbirth, the risk of perforation is greater. According to the WHO medical eligibility criteria for contraceptive use, there are no restrictions for inserting an IUD four or more weeks after childbirth; however, no distinction is made between a vaginal or caesarean delivery. The type of delivery is important, because a cesarean section can distort the uterine anatomy. If the distortion is found to be significant during a bimanual examination, then inserting the IUD may be aided by uterine ultrasonography.\(^{17}\)

Adhering to the proper insertion method and using a sterile or “no-touch” technique are sufficient to decrease the risk of infection, uterine perforation, and expulsion. Data indicate that antibiotic prophylaxis does not prevent the rare occurrence of infections acquired at the time of insertion. A Cochrane review performed to assess the effectiveness of prophylactic antibiotic administration before IUD insertion in reducing IUD-related complications and discontinuations within three months of insertion showed little benefit from prophylactic antibiotics on early removal or occurrence of PID following insertion.\(^{18}\) Prophylactic antibiotics are therefore not recommended prior to insertion of the IUD.

Pain may occur during the insertion procedure that may result from a vasovagal reaction, the need for cervical dilation, difficulty with insertion, and uterine perforation. While some clinicians administer ibuprofen or another nonsteroidal anti-inflammatory drug (NSAID) to control such pain, a recent study of 2,019 first-time IUD users found that ibuprofen had no significant impact on reducing pain.\(^{19}\) In a comparative study conducted in the United Kingdom, 102 women presenting for IUD insertion were randomly assigned to 1 of 3 groups: no pretreatment, pretreatment with 2% lignocaine gel (Instillagel) applied to the cervical canal, and pretreatment with an inert gel. Pain scores for women in the nontreatment groups were significantly higher (p < 0.025) than those in the lignocaine group. These results suggest that a local lignocaine gel has the potential for reducing the pain and discomfort experienced during IUD insertion.\(^{20}\) In an attempt to reduce cervical resistance, a randomized, controlled trial was undertaken among 80 nulliparous women randomly assigned to receive sublingually 400 µg misoprostol and 100 mg diclofenac or 100 mg diclofenac alone one hour prior to IUD insertion. Following treatment with misoprostol, insertion was significantly easier with fewer difficult and failed attempts at insertions than in the control group. However, pain scores, estimated using a visual analogue scale (VAS; 1–10), were not different between the groups.\(^{21}\) Therefore, there may be some benefit to use of misoprostol for cervical ripening prior to LNG-IUS insertion as well as local anesthetic use for pain control, while NSAIDs appear to be of little benefit.

**Efficacy of the levonorgestrel-releasing IUS**

The IUD, in general, is one of the most effective forms of contraception available today, with a global cumulative
The LNG-IUS is also an effective option for women to choose immediately post abortion. Multiple studies have shown that copper-releasing IUDs are safe and effective during this timeframe. Fewer studies have been done looking specifically at the LNG-IUS under this circumstance; however, they have shown it to be at least as effective as the copper IUD. One study followed 305 women who received the Mirena® IUS immediately post-abortion for up to five years. There were a total of two pregnancies for a pregnancy rate of 0.8% at five years. There was a discontinuation rate secondary to expulsion of 7.1% at one year and 10.5% at five years which is obviously higher than the interval expulsion rate of 2%–3% per year, but not significantly different than that of the NovaT (8.6% at one year and 15.4% at five years). A LNG-IUS may, therefore, be safely inserted immediately after either uncomplicated spontaneous or induced abortions.

The levonorgestrel-releasing intrauterine system can also be safely inserted immediately after childbirth, but there is also an increased risk of expulsion compared with interval insertion. A study of 19 women who received a post placental LNG-IUS, defined as placement within 10 minutes of placental delivery, showed an expulsion rate of 10.5% and no infections. There are no studies that specifically examine the timing of LNG-IUS insertion post partum, but one done with a copper-releasing IUD showed post-placental IUD insertion to have significantly lower expulsion rate (14.3% complete and 22.6% partial) than early post partum insertion, defined as taking place 10 min to 72 hours after delivery of the placenta, at the end of one year (18.6% complete and 51.2% partial). Both rates were much higher than interval insertion (3.8% complete and 3.1% partial). There were no perforations in either of the post partum groups, and 2.3% perforation rate in the interval groups. Insertion of the LNG-IUS is therefore a reasonable, almost effortless, option for post placental insertion if the woman is properly counseled about the increased risk of expulsion versus interval insertion and receives proper counseling on checking the IUS strings and receives appropriate clinical follow-up. For many low-resource settings, since the cervix (or uterus in the case of insertions at Cesarean delivery) is open at delivery and the patient is commonly in lithotomy position, post-placental insertion may reduce the barriers to interval insertions such as lack of speculae, tenacula, sounds, exam tables, and already crowded clinics.

Noncontraceptive benefits of LNG-IUS

Like oral contraceptives, the levonorgestrel-releasing intrauterine system confers important noncontraceptive health benefits. The Mirena® LNG-IUS can be effective in treating a variety of gynecological disorders including menorrhagia, dysmenorrhea, pain associated with endometriosis, anemia, endometrial hyperplasia and can be used as an alternative to hysterectomy for women with bleeding problems as well as an adjunct to estrogen replacement therapy.

The LNG-IUS as treatment for menorrhagia

Menorrhagia, defined as >80 mL of menstrual blood loss, is a common health problem affecting up to 2.5 million women in the United States annually. It accounts for a loss of 8% of employee wages in the USA, and is a major reason for women to present to their health care provider for invasive procedures, including hysterectomy. The prevalence of menorrhagia increases with increasing age, peaking in the perimenopausal period. Nearly half a million women annually in the United States alone undergo hysterectomy because of bleeding disorders, with the associated health care costs, time from work, and morbidity that accompany this operative treatment. There are a variety of medical and surgical treatments available for menorrhagia, including prostaglandin synthetase inhibitors, antifibrinolytic agents, oral contraceptive pills, and endometrial ablation. However, most of these treatments have only been shown to improve menstrual bleeding by 20%–50%. In addition, the associated side effects and repetitive nature of these treatments significantly impair patient compliance. In contrast, the LNG-IUS has been used in numerous studies as a treatment of menorrhagia, with a reduction in menstrual blood flow of 86%–97%. One study comparing the LNG-IUS and norethisterone showed that the LNG-IUS reduced menstrual blood flow by 94%, with norethisterone by a close 87%. However, after three cycles of treatment, 76% of the
LNG-IUS group wished to continue the treatment, compared with only 22% of the medical therapy group.\textsuperscript{31}

In addition, the LNG-IUS has been used in numerous studies to compare outcomes with hysterectomy for treatment of menorrhagia. A randomized controlled trial of 236 women assigned either to LNG-IUS or hysterectomy showed that after one year the two treatments were associated with equal improvements in health status, quality of life, and psychosocial well-being, but the IUS was more cost-effective. Of note, 20% of the women in the LNG-IUS group did go on to receive hysterectomies because of continued bleeding.\textsuperscript{32} Another study which looked specifically at the cost-effectiveness of oral contraceptives versus LNG-IUS versus surgical management (including both ablation and hysterectomy) for dysfunctional uterine bleeding (DUB) showed the LNG-IUS to be the most cost-effective of all three treatments.\textsuperscript{33} A Cochrane review of the subject concluded that use of the LNG-IUS results in a significant decrease from baseline in the amount of menstrual bleeding, and that it is more cost-effective as a treatment for menorrhagia than hysterectomy both at one and five years.\textsuperscript{34} The LNG-IUS is, therefore, a satisfactory, effective, and economical alternative to medical and surgical treatment of menorrhagia.

The LNG-IUS as treatment for endometriosis

Endometriosis is a significant problem affecting 5%–10% of reproductive age women in the United States.\textsuperscript{35} It is associated with chronic pelvic pain, dyspareunia and infertility, and is often a significant detriment to a patient’s quality of life. Treatment has historically consisted of some combination of nonsteroidal anti-inflammatory medications (NSAIDs), progestational medications such as depot medroxyprogesterone acetate (DMPA) that function as anti-estrogens, ovulation suppression with oral contraceptive pills, androgenic medications such as danazol, gonadotropin-releasing hormone (GnRH) analogues to induce temporary pseudo-menopause, and surgical ablation. However, the hypoestrogenic side effects associated with many of these medical treatments and the invasive nature of the surgical treatment are a limiting factor to both the clinical ability to continue the treatment long term and patient compliance. The LNG-IUS has recently been studied as an alternative for treating the pain and symptoms associated with endometriosis, as it addresses both the patient compliance and the long-term use issues. A randomized-controlled clinical trial of 82 women with chronic pelvic pain from endometriosis compared 39 women treated with the LNG-IUS with 43 women treated with a GnRH analogue. After six months, both groups showed significant improvement in pain scores throughout the six months of treatment, with those women with stage III or IV endometriosis receiving the fastest improvement. There was no difference in quality of life reported, and the LNG-IUS group had the benefits of fewer hypoestrogenic side effects, and only requiring one intervention every five years.\textsuperscript{36} After 36 months, 59% (n = 23) of the women in the LNG-IUS group were still using the device. The VAS pain score in this group was 0–3 (indicating excellent pain control) in 82.6%, and ≥7 in only 8.6%. In the former users of the GnRH analogue, 28% were using some form of hormonal contraceptives to control pain, and 47.5% continued to have excellent pain control, while only 2.5% had a pain score ≥7. Therefore, while both treatments are effective in pain control for up to 3 years, the LNG-IUS may be the treatment of choice for women who do not wish to become pregnant because it can provide both effective pain control and contraceptive protection.\textsuperscript{37}

Another study of 34 women with surgically staged minimal to moderate endometriosis who were treated with the LNG-IUS for up to 36 months showed a decrease in the visual analog pain scale (VAS) from an initial score of 7.7/10 to 2.7 at 36 months. There was also a decrease in the verbal rating scale (VRS) of both dysmenorrhea and noncyclic pelvic pain from an initial 25/96 to 8.4 after 36 months as well as a decrease in total days of pain per 28-day period from 15.0 to 6.0 after 12 months.\textsuperscript{38}

A randomized-controlled clinical trial of 40 women who underwent conservative surgical treatment for moderate to severe endometriosis compared 20 women who received a postoperative LNG-IUS with 20 women who were managed expectantly postoperatively. The study showed that 10% of the women who received the LNG-IUS had a recurrence of moderate to severe endometriosis after 12 months, compared with 45% of the expectantly managed group. There was also an absolute risk reduction in recurrence of dysmenorrhea in the LNG-IUS group of 35%, as well as a decrease in recurrence rates of dyspareunia.\textsuperscript{39} The LNG-IUS appears to be an effective treatment for the pain symptoms associated with endometriosis, and is associated with fewer intervention requirements, fewer hypoestrogenic side effects, and increased patient satisfaction rates than many of the other alternative treatments clinically available.

The LNG-IUS as treatment for adenomyosis

Adenomyosis is a relatively common disorder affecting women, usually in their 40s and 50s, characterized by the
presence of heterotopic endometrial glands and stroma in the myometrium with hyperplasia of the adjacent smooth muscle. The symptoms include menorrhagia (40%–50%), dysmenorrhea (15%–30%), and metrorrhagia (10%–12%). Historically, diagnosis and definitive treatment for adenomyosis has been hysterectomy. However, diagnosis has recently become possible using a combination of transvaginal ultrasound and magnetic resonance imaging. As a result, there has been a concerted effort on the part of gynecologists to find an effective, but less invasive treatment for the disorder. The options for this condition include endometrial ablation, danazol and GnRH agonists. However, endometrial ablation is not effective for this condition as it does not penetrate deeply enough into the myometrium. In addition, as discussed previously with regards to treatment for endometriosis, in light of the side effects of the other forms of medical treatment listed, the LNG-IUS is becoming an increasingly attractive option for treatment of adenomyosis. It is hypothesized to work on adenomyosis in two ways. First, as discussed earlier, it causes decidualization and atrophy of the endometrium, therefore decreasing the amount of menstrual flow. Secondly, the levonorgestrel down regulates estrogen receptors in glandular and stromal endometrial tissues. This likely prevents further estrogen stimulation of the adenomyosis foci within the myometrium, causing them to atrophy and shrink. This may lead to decreased menstrual flow by allowing the myometrium to better contract, limiting the blood loss during menses, and also accounts for the decreased size of the uterus noted.

Several studies have been performed to evaluate the effectiveness of the LNG-IUS as a treatment for adenomyosis. In one small study of 25 women with menorrhagia associated with adenomyosis, the subjects were treated with the LNG-IUS and followed for up to 12 months. The average pictorial blood loss decreased by approximately 75% (211 ± 61 to 44 ± 18 mL). In addition, both uterine volume and endometrial thickness decreased significantly, while there was a significant increase in hemoglobin, serum iron and ferritin levels.

Another study was performed on 47 women with adenomyosis. Each received a LNG-IUS and were followed for up to 36 months. At the end of 36 months 32 women were still enrolled in the study. In these women, serum hemoglobin and ferritin levels increased significantly and CA-125 values significantly decreased. In addition, uterine volume decreased from 156.85 ± 49.79 to 118.64 ± 41.35 mL at 12 months. However, uterine volume increased from 24 to 36 months, to 139.87 ± 29.93 mL. At 36 months, this difference was not significantly different from the initial pretreatment volume. Pain scores decreased significantly from 8.55 ± 1.02 to 1.93 ± 0.95 by 6 months and remained there at 24 months. Menstrual blood flow was also markedly reduced, with a reduction of >90%, and remained similar at 24 months. However, as with uterine size, the pain scores and blood flow increased again after 36 months, though they were still significantly less than the initial values. Though the exact cause for this phenomenon is unknown, it is hypothesized that it could be attributed to the decreased amount of hormone released by the LNG-IUS daily over time. Regardless, the LNG-IUS does appear to be an effective treatment for adenomyosis with improvement in pain and bleeding scores over two years. Further studies will need to address whether replacing the LNG-IUS after 2–3 years will keep the pain and bleeding scores from increasing again over time.

The LNG-IUS as treatment for endometrial hyperplasia

Endometrial hyperplasia is classified according to increasingly abnormal architectural and cytologic criteria as simple, complex, and atypical hyperplasia. Cytologic atypia is the most important prognostic factor with regard to progression to endometrial cancer. For nonatypical hyperplasia, there is a 1%–3% chance of progression to cancer, with a 72% chance of regression after expectant management. In contrast, for atypical hyperplasia there is an 8%–30% chance of progression to endometrial carcinoma, with only a 54% chance of spontaneous regression with expectant management. In addition, endometrial cancer can coexist with atypical hyperplasia in up to 25% of cases. Because of the high potential for progression of atypical endometrial hyperplasia to carcinoma hysterectomy is generally considered to be the standard of care unless the patient desires to retain potential fertility or there are medical contraindications to surgery. In contrast, because nonatypical hyperplasia is generally considered to be low risk for progression to cancer, many patients and providers consider hysterectomy too invasive a treatment. Though there is no consensus on the best way to treat these women, they have often been treated with oral progestins. However, because of the systemic nature of the treatment there can be significant side effects that limit compliance with treatment, and when the treatment is discontinued the hyperplasia can recur. As a result, numerous studies have been done to evaluate the LNG-IUS as an alternative therapy to hysterectomy for those with endometrial hyperplasia without atypia. One study of 12 women diagnosed with hyperplasia without atypia and eight women with atypia was performed to evaluate the
long-term (three-year) remission rate after treatment with a LNG-IUS. Nineteen of the twenty women in this study developed a normal endometrium, with the other woman continuing to have focal atypical hyperplasia with a normal endometrial stripe at the end of 36 months. In addition, at the beginning of the study all women with atypical hyperplasia showed progesterone receptor expression in the epithelial cells. This expression declined significantly over the course of the study, marking the strong antiproliferative effect of the LNG-IUS, with its inhibition of estrogen bioactivity and suppression of the endometrium.45

A larger study was done involving long term (up to 106 months) follow up of 258 women with endometrial hyperplasia. Of these 258 women, 85 women were treated with 10 mg oral medroxyprogesterone acetate (MPA) for 10 days per cycle for three to six months, 66 women had a LNG-IUS placed for 3–108 months, and 107 women underwent control biopsy alone with observation. After six months 100% of the LNG-IUS group responded, compared with 54% of the oral MPA group, while 50% of the observation group spontaneously regressed. After 58–106 months, 44 of the women still retained their LNG-IUS, and 100% of these women responded, compared with 63% of the women who had their LNG-IUS removed, 60% of the oral MPA group and 49% of the observation only group. There was no significant difference between these last three groups.46 Though larger studies with long term follow up need to be done for more definitive evidence, the studies so far do seem to indicate that the LNG-IUS is a superior treatment modality for endometrial hyperplasia than low dose oral progestins or observation alone.

The LNG-IUS as adjunct to estrogen replacement therapy

It is clear that in peri- and postmenopausal females who retain their uteri and desire estrogen replacement therapy secondary to climacteric side effects it is essential to provide progestin medication in order to counteract the proliferative effect of estrogen on the endometrium. The standard treatment has been to use a cyclic oral progestin preparation at the end of each cycle in combination with the estrogen. However, the oral regimen is associated with cyclic vaginal bleeding which is burdensome to some, and can also have a variety of androgenic side effects. In addition, multiple studies have shown possible links with systemic progesterone and increased cardiovascular risks in addition to increased breast cancer risks.47,48 The LNG-IUS has been studied as an alternative method of providing locally acting progestin which can accomplish the endometrial protective effects with a minimal amount of adverse side effects secondary to its significantly decreased serum concentration. One study of 40 perimenopausal women complaining of climacteric symptoms evaluated two treatment regimens: three week cyclic treatments with 2 mg oral estradiol (E₂) valerate combined with 250 µg of oral levonorgestrel for the last 10 days of each cycle versus 2 mg of E₂ valerate continuously with the LNG-IUS. After one year the subjective symptoms in both groups were improved, and none of the women had endometrial proliferation. However, in the LNG-IUS group 15/18 women were amenorrheic, versus all women in the oral LNG group who continued with cyclic bleeding.49

Another study reviewed the available literature (19 studies with a total of 826 subjects) regarding progestin intrauterine devices in combination with estrogen, with a duration of six months to five years, and determined that none of the subjects developed endometrial hyperplasia throughout the course of the study.50 Hampton and colleagues evaluated 82 perimenopausal women treated with oral estrogen and the LNG-IUS for long term protection of the endometrium. Nonproliferative endometrium was present in 98.6% of the participants at the end of 12 months, 98.6% at 24 months, 95.5% at 36 months, 96.8% at 48 months, and 95.2% at 60 months. There were no cases of hyperplasia throughout the entire 60 month period. Amenorrhea was present in 54.4% of the women after 12 months, and 92.7% after 60 months. Overall the treatment was well tolerated, with 79.8% of the women still continuing with the estrogen and LNG-IUS after the full 60 months of the study and 92.4% of the participants rating the treatment as “good” or “very good”.51

The LNG-IUS thus appears to be a highly effective and acceptable method of providing endometrial protection for peri- and postmenopausal women who are taking estrogen replacement therapy.

Patient acceptability of the LNG-IUS

The LNG-IUS is a safe and effective device which has been shown to be useful not only as a contraceptive, but as a tool which can be used for a variety of gynecologic conditions or disorders. However, the IUS is not without its side effects. The most common side effect associated with the LNG-IUS is menstrual irregularity. In fact, nearly 25% of women discontinue using the LNG-IUS because of amenorrhea.1 Approximately 20% of LNG-IUS users will be amenorrheic by the end of 12 months,2 and 70% of users will be oligomenorrheic or amenorrheic by 24 months.52 In addition, though the systemic absorption is much smaller than with
other progestin-containing contraceptives, there is some absorption and it can occasionally be associated with a variety of undesirable side effects such as ovarian cysts, acne, weight gain, depression and decreased libido. However, the device seems to be well tolerated overall. A three-year study which specifically set out to evaluate the long-term acceptability of the LNG-IUS followed 165 women over 36 months. The study showed a three-year continuation rate of 90.3%. In total, 97% of participants reported alterations in their menstrual pattern, with 34% reporting a decrease in the amount of bleeding, 17% had persistent spotting or intermenstrual bleeding, and 56% reported at least a temporary period of amenorrhea. 81% of the women with amenorrhea viewed it as a positive change for them. There was also a significant improvement in the amount of pain associated with menstruation, with a decrease from 60% to 29% in women reporting dysmenorrhea after 36 months. The number of women who expressed that they were very satisfied with the LNG-IUS increased steadily with the duration of the treatment, with 29% after two weeks, 56% after two months, 69% after six months and 77% after 36 months.53

Another study of 78 women who had the LNG-IUS inserted for menstrual disorders also evaluated patient satisfaction through questionnaires. 12% of the women had the IUS prematurely removed, with the major reason being pain and heavy bleeding. However, the majority of participants were satisfied with their results. 78% of the women reported improvement in their periods, and 84% reported improvement in menstrual discomfort. 72% of the women reported they would use the LNG-IUS again, 73% would recommend it to their peers, and the overall satisfaction rate was 76%.54

The amount of satisfaction with the LNG-IUS, not surprisingly, appears to correlate with how well informed patients are about the possible side effects associated with it. A study that involved the evaluation of 17,914 questionnaires of current LNG-IUS users showed that 74% were very or fairly satisfied with it. User satisfaction correlated with the amount of information provided regarding different symptoms (menstrual irregularities, greasy hair/skin, pregnancy and PID) regardless of whether or not the patient actually experienced that specific symptom. In particular, the women who were warned of the possibility of amenorrhea were more satisfied than the women who were not.55

Conclusion
The levonorgestrel-releasing intrauterine system is a safe, effective and acceptable form of contraception being used by over 150 million women worldwide. It also has a multitude of noncontraceptive benefits including improvement in menorrhagia, decreased pelvic pain associated with endometriosis and adenomyosis, as well as a treatment for endometrial hyperplasia, an alternative to hysterectomy and an adjunct to estrogen replacement therapy. Overall the device is very well tolerated and patient satisfaction is quite high when the proper education regarding possible side effects has been provided. Despite all of the obvious benefits of the LNG-IUS, utilization rates remain quite low in the developed countries, and especially the United States. This is largely attributed to the negative history of the Dalkon Shield intrauterine device in the 1970s, which continues to negatively influence the opinions of both patients and health care providers. In addition, cost remains a barrier in less developed countries where demand might be high but the device is currently unaffordable in the public sector. Recently, with a concerted effort to train residents and providers in IUD and IUS use, a reversal of trends has been noted and the IUS is rapidly gaining market share in the US. This safe, acceptable and highly effective method will likely contribute significantly to the contraceptive method mix in the coming years.

Disclosure
The authors report no conflicts of interest in this work.

References


## Appendix A  Checklist for sexually transmitted infection (STI) risk assessment of intrauterine device (IUD) candidates

### Checklist for STI Risk Assessment of IUD Candidates

<table>
<thead>
<tr>
<th>Name:</th>
<th>Date:</th>
</tr>
</thead>
</table>

Please circle the appropriate answers below:

1. **Is the client less than 25 years old?**
   - Yes: 1
   - No: 0

2. **Is she currently living apart from her husband or partner?**
   - Yes: 1
   - No: 0

3. **During the last year, has she had bleeding between periods or bleeding or spotting within 24 hours after sex?**
   - Yes: 1
   - No: 0

4. **Is her school education less than secondary level?**
   - Yes: 1
   - No: 0

5. **How many different sexual partners has she had during the last 3 months? Check A if none, B if one, C if more than one.**
   - A. None: 0
   - B. One:
     - No condom use: 0
     - Some condom use: 1
     - Always used condoms: 0
   - C. More than one:
     - No condom use: 1
     - Some condom use: 1
     - Always used condoms: 0

Add up the circled scores. **Total Score:** [ ]

### Scoring

<table>
<thead>
<tr>
<th>Recommended Action</th>
<th>Low Cervical Infection Population (&lt;10%)</th>
<th>High Cervical Infection Population (≥10%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Counsel/refer for an IUD insertion without any reservations</td>
<td>If score is 0–2</td>
<td>If score is 0</td>
</tr>
<tr>
<td>Consider presumptive treatment for chlamydia/gonorrhea (if available) or counsel/refer to use another contraceptive method</td>
<td>If score is 3+</td>
<td>If score is 1+</td>
</tr>
</tbody>
</table>

*The level may change from less than secondary level to a higher educational level if almost all women (>90%) have at least some secondary education.*
### Appendix B WHO risk categories for eligibility for use of an intrauterine device\(^2\)

<table>
<thead>
<tr>
<th>Risk category</th>
<th>Description of category</th>
<th>With clinical judgment</th>
<th>Without clinical judgment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A condition for which there is no restriction for the use of the contraceptive method</td>
<td>Use the method in any circumstances</td>
<td>Use the method</td>
</tr>
<tr>
<td>2</td>
<td>A condition where the advantages of using the method generally outweigh the theoretical or proven risks</td>
<td>Generally use the method</td>
<td>Use the method</td>
</tr>
<tr>
<td>3</td>
<td>A condition where the theoretical or proven risks usually outweigh the advantages of using the method</td>
<td>Use of the method not usually recommended unless other more appropriate methods are not available or not acceptable</td>
<td>Do not use the method</td>
</tr>
<tr>
<td>4</td>
<td>A condition which represents an unacceptable health risk if the contraceptive method is used</td>
<td>Method not to be used</td>
<td>Do not use the method</td>
</tr>
</tbody>
</table>

**Notes:** “Clinical Judgement” means that a trained health care provider or the clinical resources to perform clinical tests and/or examinations are present.