PRODUCT MONOGRAPH

Pr KYLEENA®

Levonorgestrel-releasing intrauterine system (19.5 mg)

Progestogen

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## Table of Contents

### PART I: HEALTH PROFESSIONAL INFORMATION
- SUMMARY PRODUCT INFORMATION ..........................................................3
- INDICATIONS AND CLINICAL USE .................................................................3
- CONTRAINDICATIONS ......................................................................................3
- WARNINGS AND PRECAUTIONS ....................................................................4
- ADVERSE REACTIONS ......................................................................................13
- DRUG INTERACTIONS ......................................................................................15
- DOSAGE AND ADMINISTRATION ....................................................................16
- OVERDOSAGE ..................................................................................................26
- ACTION AND CLINICAL PHARMACOLOGY .....................................................27
- STORAGE AND STABILITY ..............................................................................31
- SPECIAL HANDLING INSTRUCTIONS ............................................................31
- DOSAGE FORMS, COMPOSITION AND PACKAGING .....................................31

### PART II: SCIENTIFIC INFORMATION
- PHARMACEUTICAL INFORMATION .................................................................33
- CLINICAL TRIALS ..........................................................................................33
- DETAILED PHARMACOLOGY .........................................................................35
- TOXICOLOGY ..................................................................................................36
- REFERENCES ..................................................................................................38

### PATIENT MEDICATION INFORMATION .....................................................40
KYLEENA®
Levonorgestrel-releasing intrauterine system (19.5 mg)

PART I: HEALTH PROFESSIONAL INFORMATION

SUMMARY PRODUCT INFORMATION

Table 1: Product Information Summary

<table>
<thead>
<tr>
<th>Route of Administration</th>
<th>Dosage Form, Strength</th>
<th>Clinically Relevant Nonmedicinal Ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intrauterine</td>
<td>Intrauterine system / 19.5 mg levonorgestrel (LNG)</td>
<td>Barium sulphate, copper phthalocyanine, polydimethylsiloxane elastomer, polyethylene, polypropylene, silica colloidal anhydrous, silver.</td>
</tr>
</tbody>
</table>

For a complete listing see DOSAGE FORMS, COMPOSITION AND PACKAGING section.

INDICATIONS AND CLINICAL USE

KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) is indicated for conception control for up to 5 years.

Geriatrics

KYLEENA is not indicated for use in postmenopausal women.

Pediatrics (< 18 years of age)

Safety and efficacy have been studied in women aged 18 and over. KYLEENA is not indicated for use before menarche. For additional information on the use of KYLEENA in postmenarcheal adolescents under the age of 18, please see WARNINGS AND PRECAUTIONS-Special Populations.

CONTRAINDICATIONS

KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) is contraindicated in patients with the following conditions:

- known or suspected pregnancy
- current or recurrent pelvic inflammatory disease or conditions associated with increased risk for pelvic infections
- postpartum endometritis or septic abortion during the previous three months
- abnormal uterine bleeding of unknown etiology
• congenital or acquired uterine anomaly, including fibroids, that distort the uterine cavity
• uterine or cervical malignancy
• known or suspected progestogen-dependent neoplasia, including breast cancer
• cervicitis or vaginitis, including bacterial vaginosis or other lower genital tract infections until infection is controlled
• cervical dysplasia
• active liver disease or dysfunction
• actual benign or malignant liver tumours
• hypersensitivity to levonorgestrel or any of the other ingredients in the formulation or components of KYLEENA. For a complete listing, see the DOSAGE FORMS, COMPOSITION AND PACKAGING section of the Product Monograph
• a previously inserted intrauterine contraceptive (IUC) that has not been removed
• recent trophoblastic disease while hCG levels are elevated
• bacterial endocarditis

WARNINGS AND PRECAUTIONS

Serious Warnings and Precautions

- Hormonal contraceptives DO NOT PROTECT against Sexually Transmitted Infections (STIs) including HIV/AIDS. For protection against STIs, it is advisable to use latex or polyurethane condoms IN COMBINATION WITH KYLEENA.
- Cigarette smoking increases the risk of serious adverse effects on the heart and blood vessels. Women should be counseled not to smoke (see Cardiovascular section below).
- Uterine Perforation may occur with the use of intrauterine contraceptives including KYLEENA (see Uterine Perforation section).

Carcinogenesis and Mutagenesis

Breast Cancer

Increasing age, inherited mutations, and a strong family history are the most significant risk factors for the development of breast cancer. Other established risk factors include nulliparity, first full-term pregnancy after the age of 30, menarche before the age of 12, never breastfed a child, and daily alcohol consumption. In some women, the use of hormonal contraceptives may accelerate the growth of an existing but undiagnosed breast cancer. More thorough studies are needed to determine the definitive link between hormonal contraceptive use and the potential risk of breast cancer.
Breast self-examination should be discussed with women receiving hormonal contraceptives. Women should be instructed to notify their healthcare professionals whenever any masses are detected.

Spontaneous reports of breast cancer have been received during postmarketing experience with another levonorgestrel-releasing intrauterine system (LNG-releasing IUS). Two observational studies did not support a causal relationship between breast cancer and the other LNG-releasing IUS, however an elevated breast cancer risk cannot be totally excluded since these studies did not control for confounding factors such as use of oral hormonal contraception by control subjects, genetics and lifestyle and environmental factors such as smoking and alcohol.

There is currently no conclusive evidence of an association between KYLEENA use and development of breast cancer or progression of subclinical breast cancer.

**Cardiovascular**

An individual benefit-risk assessment in relation to continued use of KYLEENA should be carried out in the event of arterial thrombosis. In particular, removal of KYLEENA should be considered if severe arterial disease such as stroke or myocardial infarction occurs. In addition, KYLEENA should be used with caution in patients with a previous history of severe arterial disease such as stroke or myocardial infarction. Women with a history of thromboembolic disorders should be made aware of the possibility of a recurrence. There have been postmarket reports of cardiovascular events, including myocardial infarction and stroke in women using another LNG IUS, although a causal relationship could not be clearly established in these cases.

**Predisposing Factors for Coronary Artery Disease**

Cigarette smoking increases the risk of serious cardiovascular side effects and mortality. Hormonal contraceptives increase this risk, especially with increasing age.

Other women who are independently at high risk for cardiovascular disease include those with diabetes, hypertension, abnormal lipid profile, or a family history of these risk factors.

**Hypertension**

If a significant elevation of blood pressure in previously normotensive or hypertensive subjects occurs at any time during KYLEENA use, KYLEENA removal should be considered.

**Congenital or Valvular Heart Disease**

KYLEENA should be used with caution in women with congenital or valvular heart disease who are at risk of infective endocarditis.

**Endocrine and Metabolism**

**Glucose Tolerance**

Combination and progestogen-only oral contraceptives, including those containing levonorgestrel, may affect glucose tolerance in some users. Diabetic patients, and those with a family history of diabetes, should be observed closely to detect any alterations in carbohydrate...
metabolism. Young diabetic patients whose disease is of recent origin, well controlled and not associated with hypertension or other signs of vascular disease such as ocular fundal changes, should be closely observed. One published clinical study indicated no change in mean daily insulin requirements in women with Type 1 diabetes using another levonorgestrel (LNG)-releasing intrauterine system (IUS) over a 12-month period (1).

Genitourinary

Bleeding Irregularities

Because irregular menstrual bleeding or spotting is common during the first few months of use, endometrial pathology should be excluded prior to insertion of KYLEENA. Irregular bleeding patterns in users of KYLEENA could mask the signs and symptoms of cervical or endometrial cancer. If bleeding irregularities develop after prolonged use, appropriate diagnostic measures should be undertaken.

Patients should be appropriately counseled on the likelihood of changes in menstrual patterns. During the first 3-6 months of KYLEENA use, the number of bleeding and spotting days may be higher and bleeding patterns may be irregular. With continued use, the frequency of amenorrhea and infrequent bleeding increases while the frequency of prolonged, irregular and frequent bleeding decreases. In clinical trials, amenorrhea developed by the end of the first year of use in approximately 12% of KYLEENA users. Over time, infrequent bleeding (1 or 2 bleeding or spotting episodes per 90-day reference period) and/or amenorrhea developed gradually in 26.4% and 22.6% of users, respectively (see ACTION AND CLINICAL PHARMACOLOGY – Pharmacodynamics). In clinical trials, a total of 81 subjects out of 1,697 (4.8%) discontinued due to uterine bleeding complaints. Reduced bleeding increases the level of blood hemoglobin. The possibility of pregnancy should be considered if menstruation does not occur after six weeks or more of amenorrhea, following a pattern of regular menses. A repeat pregnancy test is not necessary in amenorrheic women unless indicated by other symptoms.

Hematologic

An individual benefit-risk assessment in relation to continued use of KYLEENA should be carried out in the event of thrombosis. In particular, removal of KYLEENA should be considered if venous thromboembolic disease such as deep vein thrombosis or pulmonary embolism occurs. Women with a history of thromboembolic disorders should be made aware of the possibility of a recurrence. There have been postmarketing reports of arterial and venous thromboembolism (ATE, VTE) in women using another LNG-releasing IUS, although a causal relationship could not be clearly established in such cases. Epidemiological studies have indicated that women using progestogen-only oral contraceptives may have a slightly increased risk of venous thromboembolism; however, the results are not statistically significant (2-4).

Appropriate diagnostic and therapeutic measures should be undertaken immediately if there are symptoms or signs of thrombosis in users of KYLEENA. Symptoms of thromboembolism include: unilateral leg pain and/or swelling, sudden severe pain in the chest whether or not it radiates to the left arm, sudden breathlessness, sudden onset of coughing, any unusual severe prolonged headache, sudden partial or complete loss of vision, diplopia, slurred speech or aphasia, vertigo, collapse with or without focal seizure, weakness or very marked numbness.
suddenly affecting one side or part of the body, motor disturbances and acute abdomen. Symptoms or signs of retinal thrombosis are: unexplained partial or complete loss of vision, onset of proptosis or diplopia, papilledema, or retinal vascular lesions.

**Other Risk Factors for Venous Thromboembolism**
Other generalized risk factors for venous thromboembolism include but are not limited to a personal history, a family history (the occurrence of VTE in a direct relative at a relatively early age may indicate genetic predisposition), severe obesity (body mass index ≥30 kg/m²) and systemic lupus erythematosus. The risk of VTE also increases with age and smoking. The risk of VTE may be temporarily increased with prolonged immobilization, major surgery, or trauma. Also patients with varicose veins and leg cast should be closely monitored.

**Hepatic/Biliary/Pancreatic**
Steroid hormones may be poorly metabolized in patients with impaired liver function. Acute or chronic disturbances of liver function may necessitate the discontinuation of hormonal contraceptive use until markers of liver function return to normal (see **CONTRAINDICATIONS**).

To date, no studies have examined whether the avoidance of the first-pass effect through the liver, as with non-oral hormonal contraceptives, lessens concerns in women with liver conditions (5).

**Jaundice**
Patients who have had jaundice should be given hormonal contraceptives only with great care and under close observation. If jaundice develops in a patient using KYLEENA, consideration should be given to removing the system. Hormonal contraceptive-related cholestasis has been described in women with a history of pregnancy-related cholestasis. This condition may recur with subsequent hormonal contraceptive use (see **CONTRAINDICATIONS**).

**Neurologic**

**Headache**
KYLEENA should be used with caution in women with a history of severe headache or migraine headache, including migraine with focal neurological symptoms (ie, asymmetrical visual loss or other symptoms indicating transient cerebral ischemia). The onset or exacerbation of migraine or the development of headaches with a new pattern that is recurrent, persistent or severe requires evaluation of the cause and consideration to remove KYLEENA (see **Cardiovascular** section).

**Ophthalmologic**

**Contact Lenses**
Any eye problems or discomfort occurring during use of a hormonal contraceptive, including those relating to the use of contact lenses, should be assessed. If this occurs, an ophthalmologist should be consulted. Temporary or permanent cessation of wear may be advised.
Peri-operative Considerations

Thromboembolic Complications – Post surgery

Women using KYLEENA who require surgery associated with prolonged immobilization should be followed closely for signs and symptoms of thromboembolism.

Psychiatric

Patients with a history of emotional disturbances, especially the depressive type, may be more prone to have a recurrence of depression while using KYLEENA. In cases of a serious recurrence, consideration should be given to removing KYLEENA, since the depression may be drug-related.

Sexual Function/Reproduction

Ectopic Pregnancy

Carefully consider the possibility of an ectopic pregnancy in women who become pregnant while having KYLEENA in place. If a woman becomes pregnant with KYLEENA in place, the relative likelihood of ectopic pregnancy is increased. Approximately half of the pregnancies that occur with KYLEENA in place are ectopic. The possibility of ectopic pregnancy should be considered in the case of lower abdominal pain, especially in association with missed periods, or if an amenorrheic woman starts bleeding.

In clinical trials with KYLEENA, the incidence of ectopic pregnancy was approximately 0.2% per year. Women who choose KYLEENA should be told about the risk of ectopic pregnancy, including the possibility of impaired fertility or loss of fertility. Educate women to recognize and report to their healthcare professional any signs and symptoms of ectopic pregnancy.

Women with a history of ectopic pregnancy were excluded from clinical trials with KYLEENA. The risk of ectopic pregnancy in women who have a history of ectopic pregnancy and use KYLEENA is unknown. Women with a previous history of ectopic pregnancy, tubal surgery or pelvic infection carry a higher risk of ectopic pregnancy.

Expulsion

In clinical trials with KYLEENA, the incidence of expulsion was 3.5% (59 of 1690 subjects over 5 years) and in the same range as that reported for other IUDs and IUSs. Symptoms of the partial or complete expulsion of KYLEENA may include bleeding or pain. However, partial or complete expulsion can occur without the woman noticing it, leading to decrease or loss of contraceptive protection. As KYLEENA typically decreases menstrual bleeding over time, an increase of menstrual bleeding may be indicative of an expulsion (see DOSAGE AND ADMINISTRATION – Expulsion). A partially expelled KYLEENA should be removed. A new system can be inserted at that time provided pregnancy has been excluded.

The risk of expulsion may be increased when the uterus is not completely involuted at the time of insertion. Delay KYLEENA insertion a minimum of six weeks or until uterine involution is complete following a delivery or a second trimester abortion.
A woman should be advised how to check the threads of KYLEENA and to contact her healthcare professional if the threads cannot be felt.

**Ovarian Cysts / Enlarged Ovarian Follicles**

Since the contraceptive action of KYLEENA is due mainly to its local effect on the uterus, ovulatory cycles with follicular rupture usually occur in women of fertile age. Sometimes atresia of the follicle is delayed and folliculogenesis may continue. These enlarged follicles cannot be distinguished clinically from ovarian cysts.

Most of these follicles are asymptomatic, although some may be accompanied by pelvic pain or dyspareunia. In most cases, the enlarged follicles resolve spontaneously over two to three months' observation. Should an enlarged follicle fail to resolve spontaneously, continued ultrasound monitoring and other diagnostic or therapeutic measures may be appropriate. Rarely, surgical intervention may be required.

Ovarian cysts (reported as adverse reactions if they were abnormal, non-functional cysts and/or had a diameter >3 cm on ultrasound examination, regardless of presence or absence of associated symptoms) were reported at least once over the course of clinical trials in 22% of women using KYLEENA, and 0.6% of subjects discontinued because of an ovarian cyst (see ADVERSE REACTIONS – Clinical Trial Adverse Drug Reactions).

**Pelvic Infection**

KYLEENA is contraindicated in women with current or recurrent pelvic inflammatory disease or conditions associated with increased risk for pelvic infections (see CONTRAINDICATIONS). Conditions associated with an increased risk of pelvic inflammatory disease (PID) include established immunodeficiency and acute malignancies affecting blood or leukemias.

The inserter provided with KYLEENA helps protect the system from contamination with micro-organisms during insertion, thereby minimizing the risk of pelvic infection. The exposed product should be handled with aseptic precautions (see DOSAGE AND ADMINISTRATION – Insertion Instructions). Known risk factors for PID include multiple sexual partners, sexually transmitted infections, prior history of PID, and young age. Less common causes of pelvic inflammatory disease include pelvic actinomycosis and pelvic tuberculosis, both of which are extremely rare.

There is an increased risk of PID related to the insertion procedure during 20 days following the insertion of IUDs. Thereafter, the risk of PID during the use of IUDs or levonorgestrel-releasing intrauterine systems is small. In clinical trials, PID was observed in 0.5% of women overall. Patients should be advised to report to their healthcare professionals promptly if they experience symptoms suggestive of PID (6-8). PID may be asymptomatic but tubal damage and resulting fertility issues may still occur.

If recurrent endometritis or pelvic infections are experienced, or if an acute infection does not respond to treatment within a few days, KYLEENA must be removed.
**Sepsis**

There have been very rare postmarket reports of Group A streptococcal sepsis (GAS) temporally associated with IUC insertion. Because death from GAS is more likely if treatment is delayed, it is important to be aware of these rare but serious infections. Aseptic technique during insertion of KYLEENA is essential in order to minimize serious infections such as GAS.

**Uterine Perforation**

Partial perforation (uterine embedment) or complete perforation of the uterus wall or cervix may occur during insertion with intrauterine contraceptives, although the perforation may not be detected until later. Pregnancy may result from partial or complete perforation. If partial or complete perforation occurs, KYLEENA must be located and removed; surgery may be required. Partial perforation (uterine embedment) can result in difficult removal. Delayed detection of perforation may result in migration outside the uterine cavity, adhesions, peritonitis, intestinal perforation and obstruction, abscesses and erosion of adjacent viscera. The number of uterine perforations is linked to the experience of the person inserting the system (9). During clinical trials, perforation occurred rarely, at a rate between 0.1 and 1 per 1000 insertions (see **ADVERSE REACTIONS – Clinical Trial Adverse Drug Reactions**). Clinical trials with KYLEENA excluded breast-feeding women.

In a large, prospective, comparative, non-interventional cohort study (1 year follow-up period) in users of another LNG-IUS¹ and copper IUDs (N = 61,448 women), the incidence of perforation was 1.3 (95% CI: 1.1-1.6) per 1000 insertions in the entire study cohort; 1.4 (95% CI: 1.1-1.8) per 1000 insertions in the cohort of another LNG-IUS, and 1.1 (95% CI: 0.7 - 1.6) per 1000 insertions in the copper IUD cohort. Extending the observational period to 5 years in a subgroup of this study (N = 39,009 women using another LNG-IUS or copper IUD), the incidence of perforation detected at any time during the entire 5-year period was 2.0 (95% CI: 1.6 – 2.5) per 1000 insertions; 2.1 (95% CI: 1.6 – 2.8) and 1.6 (95% CI: 0.9 – 2.5) for another LNG-IUS and copper IUD, respectively.

The study showed that both breast-feeding at the time of insertion and insertion up to 36 weeks after giving birth were associated with an increased risk of perforation (see Table 2). These risk factors were confirmed in the subgroup followed up for 5 years. Both risk factors were independent of the type of IUD inserted.

| Table 2: Incidence of perforation per 1000 insertions for the entire study cohort observed over 1 year, stratified by breastfeeding and time since delivery at insertion (parous women) |
|---|---|
| **Insertion ≤ 36 weeks after delivery** | Breastfeeding at time of insertion | 5.6 (95% CI 3.9-7.9; n=6047 insertions) |
| | Not breastfeeding at time of insertion | 1.7 (95% CI 0.8-3.1; n=5927 insertions) |
| **Insertion > 36 weeks after delivery** | 1.6 (95% CI 0.0-9.1; n=608 insertions) | 0.7 (95% CI 0.5-1.1; n=41910 insertions) |

¹ “Another LNG-IUS” refers to an LNG-IUS that is slightly larger than KYLEENA and contains more levonorgestrel.
The risk of perforation may be increased in women with abnormal uterine anatomy or fixed retroverted uterus.

To reduce the possibility of perforation postpartum, KYLEENA insertion should be delayed a minimum of 6 weeks after delivery or until uterine involution is complete. If involution is delayed, consider waiting until 12 weeks postpartum. Inserting KYLEENA immediately after first trimester abortion is not known to increase the risk of perforation, but insertion after second trimester abortion should be delayed until uterine involution is complete.

To reduce the possibility of perforation, it is important to follow the recommended insertion technique (see DOSAGE AND ADMINISTRATION – Insertion Instructions).

Inform patients before the procedure about the risk of uterine perforation and educate them on possible signs of this complication, including, but not limited to, severe low abdominal pain, which may be associated with bleeding after the procedure, loss of threads, or change in thread length.

**Uterine Embedment**

Embedment of KYLEENA in the myometrium may occur. Embedment may decrease contraceptive effectiveness and result in pregnancy. An embedded KYLEENA must be removed. Embedment can result in difficult removal, and may require surgery.

**Special Populations**

**Pregnant Women/Intrauterine Pregnancy**

The use of KYLEENA during an existing or suspected pregnancy is contraindicated (see also CONTRAINDICATIONS). If a woman becomes pregnant with KYLEENA in place, the likelihood of ectopic pregnancy is increased. If pregnancy occurs with KYLEENA in place, KYLEENA should be removed since any intrauterine system left in place may increase the risk of abortion and preterm labour. Removal of KYLEENA or probing of the uterus may result in spontaneous abortion. In the event of an intrauterine pregnancy with KYLEENA in place, consider the following:

a) Risk of septic abortion

b) Continuation of pregnancy

If KYLEENA cannot be removed or the woman chooses not to have it removed, she should be warned that failure to remove KYLEENA increases the risk of miscarriage, sepsis, premature labor, and premature delivery. Ectopic pregnancy should be excluded. The woman should be followed closely and advised to report any abnormal symptoms, such as fever, chill, cramping, abdominal pain, bleeding, vaginal discharge, or leakage of fluid.

c) Long-term effects and congenital anomalies

When pregnancy continues with KYLEENA in place, long-term effects on the offspring are unknown. Congenital anomalies in live births have occurred infrequently with another LNG-releasing IUS. No clear trend towards specific anomalies has been observed. Because of the intrauterine administration of levonorgestrel and local exposure of the fetus to the hormone, the
possibility of teratogenicity following exposure to KYLEENA cannot be completely excluded. Some observational data support a small increased risk of masculinization of the external genitalia of the female fetus following exposure to progestogens at doses greater than those currently used for oral contraception. Whether these data apply to KYLEENA is unknown.

**Nursing Women**

Hormonal contraceptives are not recommended as the contraceptive method of first choice in breast-feeding women. A published study indicated that during lactation in users of another LNG releasing IUS (52mg), 0.1% of the daily maternal dose of LNG could be transferred to the newborn via milk (10). Although LNG may be found in the breast milk of women using LNG releasing IUS, there does not appear to be a detrimental effect on growth or development of breast-fed infants whose mothers started using the product after six weeks postpartum. Progestogen-only contraceptive methods do not appear to affect the quantity and quality of breast milk. However, isolated cases of decreased milk production have been reported with KYLEENA.

**Geriatrics**

KYLEENA is not indicated for use in postmenopausal women.

**Pediatrics (< 18 years of age)**

Safety and efficacy have been studied in women aged 18 and over. Efficacy is expected to be the same for post-menarcheal adolescents under the age of 18 as for users 18 years and older. KYLEENA is not indicated for use before menarche.

**Monitoring and Laboratory Tests**

**Magnetic Resonance Imaging (MRI)**

Nonclinical testing with another LNG-IUS with the same size silver ring and T-body has demonstrated that a patient can be scanned safely after placement of KYLEENA (“MR Conditional”) under the following conditions:

- Static magnetic field of 3-Tesla or less
- Spatial gradient magnetic field of 36,000 Gauss/cm (T/m) or less
- Maximum whole body averaged specific absorption rate (SAR) of 4W/kg in the First Level Controlled mode for 15 minutes of continuous scanning

In non-clinical testing, the aforementioned LNG-IUS produced a temperature rise of 1.8°C or less at a maximum whole body averaged specific absorption rate (SAR) of 2.9 W/kg, for 15 minutes of MR scanning at 3T using a transmit/receive body coil. A small amount of imaging artifact may occur if the area of interest is in the same area or relatively close to the position of KYLEENA.
**Physical Examination and Follow-up**

Before insertion, the woman must be informed of the efficacy, risks, and side effects of KYLEENA.

A thorough history and physical examination should also be performed prior to insertion, including a blood pressure determination. Breasts, liver, extremities, and pelvic organs should be examined. Cervical smear (Papanicolaou smear) should be performed as needed, according to healthcare professional’s evaluation. Pregnancy and sexually transmitted infections should be excluded, and genital infections have to be successfully treated. For timing of insertion to exclude pregnancy, see **DOSAGE AND ADMINISTRATION – Insertion, Removal and Replacement**.

Women should be re-examined 4 to 12 weeks after insertion and at least once a year thereafter, or more frequently if clinically indicated. At each annual visit, examination should include those procedures that were done at the initial visit as outlined above or per recommendations of the Canadian Task Force on Preventive Health Care.

See also **DOSAGE AND ADMINISTRATION – Medical Examination/Consultation**.

**ADVERSE REACTIONS**

**Adverse Drug Reaction Overview**

The majority of women experience changes in menstrual bleeding pattern after insertion of KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]). Over time, the frequency of amenorrhea and infrequent bleeding increases, and the frequency of prolonged, irregular and frequent bleeding decreases (see **WARNINGS AND PRECAUTIONS – Genitourinary, Bleeding Irregularities**; and **ACTION AND CLINICAL PHARMACOLOGY – Pharmacodynamics**).

The serious adverse reactions of ectopic pregnancy, intrauterine pregnancy, sepsis, pelvic inflammatory disease, perforation, expulsion and ovarian cysts are discussed in **WARNINGS AND PRECAUTIONS**.

**Clinical Trial Adverse Drug Reactions**

*Because clinical trials are conducted under very specific conditions the adverse reaction rates observed in the clinical trials may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials of another drug. Adverse drug reaction information from clinical trials is useful for identifying drug-related adverse events and for approximating rates.*

Adverse drug reactions (ADRs) were collected from a total of 1,697 patients in contraception studies (3-year Phase II study and 3-year Phase III study with extension to 5 years), including 1,425 exposed for one year and 550 who completed the five year study. The data cover approximately 68,000 cycles of exposure or more than 5,225 women-years. ADRs were more common during the first year after insertion of KYLEENA, and then gradually decreased over time.
### Table 3 - Adverse drug reactions, phase II and III clinical trials, N= 1697 women (5225.52 women-years)

<table>
<thead>
<tr>
<th>System Organ Class (MedDRA)</th>
<th>Very common (≥ 10%)</th>
<th>Common (≥ 1% to &lt; 10%)</th>
<th>Uncommon (≥ 0.1% to &lt; 1%)</th>
<th>Rare (≥ 0.01% to &lt; 0.1%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psychiatric disorders</td>
<td></td>
<td>Depressed mood/ Depression</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>Headache</td>
<td>Migraine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>Abdominal/ pelvic pain</td>
<td>Nausea</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td>Acne/ Seborrhoea</td>
<td>Alopecia</td>
<td></td>
<td>Hirsutism</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td>Bleeding changes including increased and decreased menstrual bleeding, spotting, oligomenorrhoea and amenorrhoea Ovarian cyst a Vulvovaginitis</td>
<td>Upper genital tract infection Dysmenorrhea Breast pain/ discomfort Device expulsion (complete and partial) Genital discharge</td>
<td></td>
<td>Uterine perforation b</td>
</tr>
</tbody>
</table>

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*Ovarian cysts had to be reported as AEs if they were abnormal, non-functional cysts and/or had a diameter > 3 cm on ultrasound examination, regardless of presence or absence of associated symptoms.*

*This frequency is based on a large, prospective, comparative, non-interventional cohort study. Please see Postmarket Adverse Drug Reactions. In clinical trials with KYLEENA that excluded breastfeeding women, the frequency of perforation was “rare”.*

In the combined Phase II and III studies, 22% discontinued prematurely due to an adverse reaction. The most common adverse reactions (>1%) leading to discontinuation were increased bleeding (4.5%), abdominal pain/pelvic pain (4.2%), device expulsion (3.1%), acne/seborrhea (2.2%), and dysmenorrhea/uterine spasm (1.3%).

In the clinical trials with KYLEENA (N= 1697), serious adverse reactions occurring in more than a single subject included: ectopic pregnancy/ruptured ectopic pregnancy (10 subjects); pelvic inflammatory disease (6 subjects); missed abortion/incomplete spontaneous abortion/spontaneous abortion (4 subjects); ovarian cyst (3 subjects); abdominal pain (4 subjects); depression/affective disorder (4 subjects); and uterine perforation/embedded device (myometrial perforation) (4 subjects/5259 subjects).

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2 Includes all insertions from clinical trials with a similar IUS, inserter dimensions and identical insertion procedures.
Postmarket Adverse Drug Reactions

In a large, prospective, comparative, non-interventional cohort study (1 year follow-up period) in users of another LNG-IUS and copper IUDs (N = 61,448 women), the incidence of perforation was 1.3 (95% CI: 1.1-1.6) per 1000 insertions in the entire study cohort; 1.4 (95% CI: 1.1-1.8) per 1000 insertions in the cohort of another LNG-IUS, and 1.1 (95% CI: 0.7 - 1.6) per 1000 insertions in the copper IUD cohort. This study showed that breastfeeding at the time of insertion and insertion up to 36 weeks after giving birth are independent risk factors for perforation (see WARNINGS AND PRECAUTIONS – Sexual Function/Reproduction, Uterine Perforation).

The following adverse reactions have been identified during post-approval use of LNG-releasing IUSs. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure:

- Arterial thrombotic and venous thromboembolic events, including cases of pulmonary embolism, deep vein thrombosis and stroke
- Device breakage, which can occur with KYLEENA in place or during its removal. The broken pieces should be located and removed; surgery may be required. Check device integrity when removing KYLEENA.
- Hypersensitivity (including rash, pruritis, urticaria, and angioedema)
- Increased blood pressure
- Isolated postmarketing cases of decreased milk production have been reported in women using a LNG-IUS

DRUG INTERACTIONS

Drug-Drug Interactions

No drug-drug interaction studies have been conducted with KYLEENA.

The effect of hormonal contraceptives may be impaired by drugs which induce liver enzymes, specifically cytochrome P450 enzymes. The influence of these drugs on the efficacy of KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) has not been studied, but it is not believed to be of major importance due to the local action of KYLEENA.

Substances increasing the clearance of levonorgestrel

Substances which may increase the clearance of levonorgestrel include phenytoin, barbiturates, primidone, carbamazepine, rifampicin, and possibly also oxcarbazepine, topiramate, and products containing St. John’s wort.

Substances with variable effects on the clearance of levonorgestrel

When co-administered with sex hormones, many HIV/HCV protease inhibitors and non-nucleoside reverse transcriptase inhibitors can increase or decrease plasma concentrations of the progestin.
**Substances decreasing the clearance of levonorgestrel (enzyme inhibitors)**

Strong and moderate CYP3A4 inhibitors such as azole antifungals (eg, fluconazole, itraconazole, ketoconazole, voriconazole), verapamil, macrolides (eg, clarithromycin, erythromycin), diltiazem and grapefruit juice can increase plasma concentrations of the progestin.

**Drug-Food Interactions**

Interactions with food have not been established.

**Drug-Herb Interactions**

Interactions with herbal products have not been established.

**Drug-Laboratory Test Interactions**

Interactions with laboratory tests have not been established.

**Tissue Specimens**

Pathologists should be advised of KYLEENA therapy when specimens obtained from surgical procedures and Pap smears are submitted for examination.

**Drug-Lifestyle Interactions**

The effect of KYLEENA on the ability to drive or to use machines has not been studied. Patients should be advised not to drive or use machines until they know how they react to KYLEENA.

**DOSAGE AND ADMINISTRATION**

**Recommended Dose**

Following insertion into the uterine cavity, KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) is effective for up to 5 years. KYLEENA must be removed by the end of the fifth year and can be replaced at the time of removal with a new KYLEENA if continued contraceptive protection is desired.

The in vivo release curve is characterized by an initial steep decline that slows down progressively resulting in little change after 1 year until the end of the intended 5-year period of use. Estimated in vivo delivery rates for different time points are provided in Table 4.
Table 4: Estimated *in vivo* release rates based on observed *ex vivo* residual content data

<table>
<thead>
<tr>
<th>Time</th>
<th>Estimated <em>in vivo</em> release rate [micrograms/24 hours]</th>
</tr>
</thead>
<tbody>
<tr>
<td>24 days after insertion</td>
<td>17.5</td>
</tr>
<tr>
<td>60 days after insertion</td>
<td>15.3</td>
</tr>
<tr>
<td>1 year after insertion</td>
<td>9.8</td>
</tr>
<tr>
<td>3 years after insertion</td>
<td>7.9</td>
</tr>
<tr>
<td>5 years after insertion</td>
<td>7.4</td>
</tr>
<tr>
<td>Average over 1st year</td>
<td>12.6</td>
</tr>
<tr>
<td>Average over 5 years</td>
<td>9.0</td>
</tr>
</tbody>
</table>

**Administration**

**Medical Examination/Consultation**

Before insertion, the woman must be informed of the efficacy, risks, and side effects of KYLEENA. A thorough history and physical examination should also be performed prior to insertion, including a blood pressure determination. Breasts, liver, extremities, and pelvic organs should be examined. Cervical smear (Papanicolaou smear) should be performed as needed, according to healthcare professional’s evaluation. Pregnancy and sexually transmitted infections should be excluded and any genital infections must be successfully treated. For timing of insertion to exclude pregnancy, see **DOSAGE AND ADMINISTRATION – Insertion, Removal and Replacement**. The position of the uterus and the size of the uterine cavity should be determined. Fundal positioning of KYLEENA is particularly important in order to ensure uniform exposure of the endometrium to the progestogen, prevent expulsion and maximize efficacy. Because irregular bleeding is common during the first months of therapy with all IUSs, including KYLEENA, it is recommended to exclude endometrial pathology before insertion (see **WARNINGS AND PRECAUTIONS – Genitourinary, Bleeding Irregularities**; and **ADVERSE REACTIONS**). It is also important to counsel women about how their bleeding pattern may change after insertion of KYLEENA. The instructions for insertion should be followed carefully. The patient should be re-examined 4 to 12 weeks after insertion and once a year thereafter, or more frequently if clinically indicated.

KYLEENA is not for use as a postcoital contraceptive.

**Insertion, Removal and Replacement**

It is recommended that KYLEENA only be inserted by healthcare professionals who are experienced in IUS insertions and/or have undergone training on the KYLEENA insertion procedure.

KYLEENA can be distinguished from other IUSs by the silver ring which is visible upon ultrasound as well as the blue colour of the removal threads. The T-body of KYLEENA contains barium sulphate, which makes it visible in X-ray examination.

In women of fertile age, KYLEENA should be inserted within 7 days of the onset of menstruation. In this case no back up contraception is needed. KYLEENA can be inserted any time during the cycle if the healthcare professional can be reasonably certain (as defined by the
World Health Organization) that the woman is not pregnant. If insertion is more than 7 days since menstrual bleeding started, a barrier method of contraception should be used or the patient should abstain from vaginal intercourse for the next 7 days to prevent pregnancy. Consider the possibility of ovulation and conception before using this product. KYLEENA may be replaced by a new system at any time during the cycle. The system can also be inserted immediately after first trimester abortion.

Insertions following second trimester abortion should be postponed for a minimum of 6 weeks or until the uterus is fully involuted. If involution is delayed, wait until involution is complete before insertion.

Postpartum insertions should be postponed until the uterus is fully involuted, and not earlier than 6 weeks after delivery. If involution is substantially delayed, consider waiting until 12 weeks postpartum. In case of a difficult insertion and/or exceptional pain or bleeding during or after insertion, the possibility of perforation should be considered and appropriate steps should be taken, such as physical examination and ultrasound.

If bleeding irregularities develop during prolonged treatment, appropriate diagnostic measures should be taken as irregular bleeding may be a symptom of endometrial polyps, hyperplasia or cancer.

The removal threads may be felt by the partner during intercourse.

KYLEENA can be removed by gently pulling on the removal threads with forceps. If the threads are not visible and the system is in the uterine cavity, it may be removed using forceps. This may require dilatation of the cervical canal or other surgical intervention, such as hysteroscopy.

After removal of KYLEENA, verify that the system is intact. The system should be removed after 5 years of use. If the patient wishes to continue using KYLEENA, a new system can be inserted at the time of removal of the old one. If pregnancy is not desired, removal should be carried out within 7 days of the onset of menstruation in women of fertile age provided that there appears to be a menstrual cycle. If the system is removed at some other time or the woman does not experience regular menses and the woman has had intercourse within a week, she is at risk of pregnancy. To ensure continuous contraception, a new system should be immediately inserted or an alternative contraceptive method should have been initiated.

Insertion and removal may be associated with some pain and bleeding. The procedure may cause a fainting spell or precipitate a seizure in an epileptic patient. It is recommended to wait 24 to 48 hours before having sexual intercourse in the event of general discomfort after insertion of KYLEENA.

**Expulsion**

Symptoms of the partial or complete expulsion of KYLEENA may include bleeding or pain; however, a system may be expelled from the uterine cavity without the patient noticing it. Partial expulsion may decrease the effectiveness of KYLEENA. Since KYLEENA decreases menstrual flow, an increase in menstrual flow may indicate an expulsion. A displaced system should be removed. A new system can be inserted at that time and the patient should be advised on how to check for the presence of the system by feeling for the removal threads.
In clinical trials with KYLEENA, the incidence of expulsion was 3.5% (59 of 1690 subjects over 5 years), and in the same range as that reported for other IUDs and IUSs. Overall, more than half of the expulsions occurred during the first 12 months after insertion.

In clinical trials with KYLEENA, more expulsions occurred in parous women (6.1%), than in nulliparous women (2.5%).

**Lost Removal Threads**

If the threads are not visible upon follow-up examination, they may have retracted into the uterus or broken, or KYLEENA may have broken, perforated the uterus, or been expelled. If the length of the threads has changed since the length at the time of insertion, the system may have become displaced (see **DOSAGE AND ADMINISTRATION – Administration, Expulsion**). Pregnancy must be excluded and the location of KYLEENA must be verified by sonography (KYLEENA contains a silver ring to facilitate detection by ultrasound), X-ray (KYLEENA is radiopaque), or by gentle exploration of the uterine cavity with a probe. If KYLEENA is displaced, remove it. A new KYLEENA may be inserted at that time or during the next menses if it is certain that conception has not occurred.

**Insertion Instructions**

Because the insertion technique is different from other intrauterine devices, it is important that healthcare professionals receive training on the correct insertion technique.

Healthcare professionals should become thoroughly familiar with the insertion instructions in their entirety before insertion of KYLEENA.

KYLEENA is supplied in a sterile package which should not be opened until required for insertion. It is sterilized with ethylene oxide. Do not resterilize. For single use only. Do not use if the seal of the sterile package is broken, or if the package is damaged or opened. The exposed product should be handled with aseptic precautions. Insert before the date indicated on the label.

KYLEENA is supplied with a patient reminder card in the outer package. Complete the patient reminder card and give it to the patient, after insertion.

In women of fertile age, KYLEENA is to be inserted into the uterine cavity within 7 days of the onset of menstruation. In this case no back up contraception is needed. KYLEENA can be inserted any time during the cycle if the healthcare professional can be reasonably certain (as defined by the World Health Organization) that the woman is not pregnant. If insertion is more than 7 days since menstrual bleeding started, a barrier method of contraception should be used or the patient should abstain from vaginal intercourse for the next 7 days to prevent pregnancy. Consider the possibility of ovulation and conception before using this product. KYLEENA is not suitable for use as postcoital contraceptive. KYLEENA can be replaced by a new system at any time in the cycle.

**Preparation for Insertion**

1. Visualize the cervix with the aid of a speculum and thoroughly cleanse the cervix and
vagina with a suitable antiseptic solution.

2. Grasp the upper lip of the cervix with a tenaculum or suitable holding forceps to stabilize the uterus. If the uterus is retroverted, it may be more appropriate to grasp the lower lip of the cervix. Gentle traction on the holding forceps can be applied to straighten the cervical canal. The forceps should remain in position and gentle traction on the cervix should be maintained throughout the insertion procedure.

3. Gently advance a uterine sound through the cervical canal to the fundus to determine the depth and confirm the direction of the uterine cavity, and to exclude any evidence of intrauterine abnormalities (eg, uterine septum, synechiae or submucosal fibroids) or a previously inserted intrauterine contraceptive which has not been removed. If difficulty is encountered, consider dilatation of the canal. If cervical dilatation is required, consider using analgesics and/or paracervical block.

**Insertion**

**Step 1–Opening of the sterile package**

- First, open the sterile package completely (Figure 1). Then use aseptic technique and sterile gloves.

![Figure 1: Sterile Package Containing KYLEENA](image)

**Step 2–Load KYLEENA into the insertion tube**

- To load KYLEENA into the insertion tube, push the slider **forward** in the direction of the arrow to the furthest position (Figure 2).
- IMPORTANT! Do not pull the slider downwards as this may prematurely release KYLEENA. **Once released, KYLEENA cannot be re-loaded.**
Step 3—Setting the flange

- Holding the slider in the furthest position, set the upper edge of the flange to correspond to the sound measurement of the uterine depth (Figure 3).
Step 4–Advance the inserter through the cervix

- While holding the slider in the furthest position, gently advance the inserter through the cervical canal and into the uterine cavity until the flange is approximately 1.5 to 2.0 cm from the external cervical os (Figure 4).

- **NOTE: Do not advance flange to the cervix at this step.** Maintaining the flange 1.5 to 2 cm from the cervical os allows sufficient space for the arms to open (when released) within the uterine cavity.

- **IMPORTANT! Do not force the inserter. If necessary, dilate the cervical canal.**
**Step 5–Release the arms**

- While holding the inserter steady, **pull the slider to the mark** to open the horizontal arms of KYLEENA (Figure 5). Wait approximately 10 seconds for the horizontal arms of KYLEENA to open completely.

![Figure 5: Releasing the Arms of KYLEENA](image-url)


**Step 6–Advance to fundal position**

- Advance the inserter gently towards the fundus of the uterus **until the flange touches the cervix** or you feel fundal resistance. KYLEENA should now be in the desired fundal position (Figure 6).

![Image of KYLEENA in fundal position](image)

**Figure 6: KYLEENA in the Fundal Position**

**Step 7–Release KYLEENA and withdraw the inserter**

- While holding the inserter in place, **pull the slider all the way down** to release KYLEENA from the insertion tube (Figure 7). The threads will release automatically from the internal thread lock of the inserter.

- Gently remove the inserter by pulling it out.

- **Cut the threads perpendicular** to the thread length, for example, with sterile curved scissors, leaving about 2-3 cm visible outside of the cervix. **NOTE:** Cutting the threads at an angle may leave sharp ends.
KYLEENA® Product Monograph

KYLEENA insertion is now complete.

In case difficulties arise during insertion, the patient complains of pain, or if there is any doubt that KYLEENA is in the correct position, verify with ultrasound or X-ray. Remove KYLEENA if it is not positioned properly in the intrauterine cavity, and insert a new one. A removed system must never be reinserted.

Patients should be re-examined 4 to 12 weeks after insertion and once a year thereafter, or more frequently if clinically indicated.

Use of Sanitary Pads

The use of sanitary pads is recommended. If tampons are used, they should be changed carefully to avoid inadvertently pulling the KYLEENA removal threads.

Removal/Replacement of KYLEENA

KYLEENA is removed by pulling on the threads with a pair of forceps (Figure 8).

A KYLEENA system should not remain in the uterus longer than 5 years.
Continuation of Contraception after Removal

KYLEENA should be removed after 5 years. If KYLEENA has been used for a longer period of time than 5 years, pregnancy should be ruled out before insertion of a new system.

If pregnancy is not desired and if a woman wishes to continue using KYLEENA, a new system can be inserted immediately after removal any time during the cycle.

If a patient with regular cycles wants to start a different contraceptive method, time removal and initiation of the new method to ensure continuous contraception. Either remove KYLEENA during the first 7 days of the menstrual cycle and start the new method immediately thereafter or start the new method at least 7 days prior to removing KYLEENA if removal is to occur at other times during the cycle.

If a patient with irregular cycles or amenorrhea wants to start a different contraceptive method, start the new method at least 7 days before removal.

OVERDOSAGE

Not applicable. KYLEENA is an intrauterine system.
**ACTION AND CLINICAL PHARMACOLOGY**

**Mechanism of Action**

KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) consists of a small polyethylene T-shaped body with a cylindrical reservoir containing levonorgestrel around the vertical stem of the T body (see Figure 10). In addition, the vertical stem contains a silver ring to aid in detection by sonography located close to the horizontal arms. The white T-body has a loop at one end of the vertical stem and two horizontal arms at the other end. Blue colored removal threads are attached to the loop. The vertical stem of the IUS is loaded in the insertion tube at the tip of the inserter.

![Figure 9: KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg])](image)

KYLEENA is a long acting reversible contraceptive (LARC). After insertion in the uterus, KYLEENA releases levonorgestrel continuously for up to 5 years. Intrauterine administration allows a very low daily dosage of progestin (average over 5 years of 9.0 mcg per 24 hours), as the hormone is released directly to the target organ. KYLEENA contains a total of 19.5 mg of levonorgestrel; the *in vivo* release curve is characterized by an initial steep decline that slows down progressively resulting in little change after 1 year until the end of the intended 5-year period of use. Estimated *in vivo* delivery rates for different time points are provided in Table 4. KYLEENA does not contain any estrogen.
Pharmacodynamics

KYLEENA has mainly local progestogenic effects in the uterine cavity. The high levonorgestrel concentration in the endometrium down-regulates endometrial estrogen and progesterone receptors. The endometrium becomes relatively insensitive to the circulating estradiol and a strong antiproliferative effect is seen. Morphological changes of the endometrium and a weak local foreign body reaction were observed during use. Thickening of the cervical mucus prevents passage of the sperm through the cervical canal. The local milieu of the uterus and of the fallopian tubes inhibits sperm mobility and function, preventing fertilization. The local mechanism by which continuously released LNG contributes to the contraceptive effectiveness of KYLEENA has not been conclusively demonstrated.

The use of KYLEENA does not alter the course of future fertility; upon removal of KYLEENA, women return to their normal fertility. In a 5-year study with KYLEENA, 116 of 163 (71.2 %) women who discontinued because of the wish for pregnancy, and with follow-up information available, had become pregnant during the 12-month follow-up.

The duration and volume of menstrual bleeding and menstrual blood loss gradually decreases after the first several months of use, after an initial increase in the number and irregularity of bleeding and spotting days. With continued use, bleeding patterns vary from regular menstruation in some women, to irregular or prolonged bleeding in other women, and amenorrhea in others. In most women, there is a trend over time towards less frequent and shorter episodes of bleeding.

Menstrual bleeding patterns were recorded by all women throughout their participation in clinical trials with KYLEENA, starting from the day of insertion. Menstrual bleeding patterns were assessed using the World Health Organization 90-day reference period method (11). Data reported for each reference period reflect the number of women actually enrolled in trials during that specific reference period and for whom valid bleeding diaries were returned. The following bleeding patterns were observed:

| Table 5: Bleeding patterns by 90-day reference period |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| KYLEENA                         | First 90 days   | Second 90 days  | End of year 1   | End of year 3   | End of year 5   |
| Amenorrhea                       | < 1 %           | 5 %             | 12 %            | 20 %            | 23 %            |
| Infrequent bleeding              | 10 %            | 20 %            | 26 %            | 26 %            | 26 %            |
| Frequent bleeding                | 25 %            | 10 %            | 4 %             | 2 %             | 2 %             |
| Prolonged bleeding*              | 57 %            | 14 %            | 6 %             | 2 %             | 1 %             |
| Irregular bleeding               | 43 %            | 25 %            | 17 %            | 10 %            | 9 %             |

*Subjects with prolonged bleeding may also be included in one of the other categories (excl. amenorrhea)

Overall, women experienced reduced bleeding with KYLEENA. Over five years, the percentage of women experiencing amenorrhea increased gradually from <1% to 23%. Rates of frequent bleeding decreased to 2% and prolonged bleeding decreased to 1% at five years. Incidence of irregular bleeding decreased, while infrequent bleeding increased during the first year and remained stable thereafter (12).
Amenorrhea is defined as no bleeding or spotting throughout the 90 day reference period. Infrequent bleeding is the occurrence of 1 or 2 bleeding/spotting episodes per 90 day reference period. Frequent bleeding is the occurrence of more than 5 bleeding/spotting episodes per 90 day reference period. Prolonged bleeding is the occurrence of bleeding/spotting episodes lasting more than 14 days per 90 day reference period. Irregular bleeding is the occurrence of 3 to 5 bleeding/spotting episodes and less than 3 bleeding/spotting-free intervals of 14 days or more per 90 day reference period.

In the phase III study, the mean number of bleeding/spotting days in the first 30 days was 17.3 and decreased to 4.6 days after one year (see Figure 10).

![Figure 10: Mean number of bleeding/spotting days by 30-day reference periods in the phase III trial]

The altered menstrual bleeding pattern that occurs with KYLEENA use is a result of the direct action of levonorgestrel on the endometrium and is not due to the suppression of the ovulatory cycle. There is no clear difference in follicle development, ovulation, or estradiol and progesterone production in women with different bleeding patterns. Ovarian function is normal and estradiol levels are maintained even when users of KYLEENA are amenorrheic.
In clinical trials with KYLEENA, ovulation was observed in the majority of women studied. Evidence of ovulation was seen in 23 out of 26 women in the first year, in 19 out of 20 women in the second year and in all 16 women in the third year. In the fourth year, evidence of ovulation was observed in the one woman remaining in the subset and in the fifth year, no women remained in this subset.

Endometrial histology has been investigated in clinical studies examining the intrauterine release of levonorgestrel at rates ranging from approximately 5 to 20 mcg/day after insertion. During treatment, a strong progestogenic effect was observed in the majority of cases. The decrease in number of bleeding days and amount of menstrual bleeding seen during the clinical studies reflects the high degree of endometrial suppression.

In the phase III pivotal study, cervical histology was evaluated at screening and annually. Epithelial cell abnormalities were seen in 1.7% of women at screening, and the maximum was observed at Month 24 with 6.3% of women. At the Month 60 visit at the end of the extension phase, epithelial cell abnormalities were detected in 4.3% of women.

**Pharmacokinetics**

**Absorption**

Following insertion, levonorgestrel (LNG) is immediately released from the IUS into the uterine cavity. More than 90% of the released levonorgestrel is systemically available. Maximum serum concentrations of levonorgestrel are reached within the first two weeks after insertion of KYLEENA. Seven days after insertion, a mean LNG concentration of 162 pg/mL was determined. Thereafter serum concentrations of LNG decline over time to reach mean concentrations of 91.3 pg/mL after 3 years and 83.1 pg/mL after 5 years. The geometric mean serum concentrations were calculated based on a population pharmacokinetic model. With the use of an LNG-IUS, the high local drug exposure in the uterine cavity leads to a strong concentration gradient from the endometrium to the myometrium (gradient: endometrium to myometrium >100-fold), and to low concentrations of LNG in serum (gradient: endometrium to serum>1000-fold).

**Distribution**

LNG is bound non-specifically to serum albumin and specifically to sex hormone-binding globulin (SHBG). Less than 2% of the circulating LNG is present as free steroid. LNG binds with high affinity to SHBG. Accordingly, changes in the concentration of SHBG in serum result in an increase (at higher SHBG concentrations) or in a decrease (at lower SHBG concentrations) of the total LNG concentration in serum. The concentration of SHBG declined on average by approximately 30% during the first 3 months after insertion of KYLEENA and remained relatively stable over the 5 year period of use. The apparent volume of distribution of LNG is reported to be approximately 1.8 L/kg.

**Metabolism**

LNG is extensively metabolized. The most important metabolic pathways are the reduction of the Δ4-3-oxo group and hydroxylations at positions 2α, 1β and 16β, followed by conjugation.
CYP3A4 is the main enzyme involved in the oxidative metabolism of LNG. The available in vitro data suggest that CYP mediated biotransformation reactions may be of minor relevance for LNG compared to reduction and conjugation.

**CYP3A4**

The available in vitro data suggest that CYP mediated biotransformation reactions may be of minor relevance for LNG compared to reduction and conjugation.

**Excretion**

The total clearance of LNG from plasma is approximately 1.0 mL/min/kg. Only trace amounts of LNG are excreted in unchanged form. The metabolites are excreted in feces and urine at an excretion ratio of about 1. The excretion half-life is about 1 day.

**Special Populations and Conditions**

**Geriatrics (> 65 years of age)**

KYLEENA is not indicated for use in postmenopausal women.

**Pediatrics (< 18 years of age)**

Safety and efficacy have been studied in women aged 18 and over. KYLEENA is not indicated for use before menarche.

**Hepatic Insufficiency**

KYLEENA has not been studied in women with hepatic impairment. KYLEENA is contraindicated in women with acute liver disease or liver tumour (see CONTRAINDICATIONS).

**Renal Insufficiency**

KYLEENA has not been studied in women with renal impairment.

**STORAGE AND STABILITY**

Store KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) at controlled room temperature (between 15°C and 30°C).

Keep out of reach of children and pets.

**SPECIAL HANDLING INSTRUCTIONS**

KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) should be handled with aseptic precautions. Used KYLEENA systems should be considered biohazardous waste and disposed of accordingly. Care should be taken to ensure the remaining hormonal ingredients are not introduced into water/sewer systems.

**DOSAGE FORMS, COMPOSITION AND PACKAGING**

KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) contains 19.5 mg of levonorgestrel in a cylindrical-shaped reservoir composed of a whitish or pale yellow matrix of levonorgestrel and polydimethylsiloxane covered with a semi-opaque membrane made of polydimethylsiloxane and silica (see Figure 9). The reservoir is mounted on the vertical stem of a
The vertical stem contains a silver ring to facilitate detection by sonography which is located close to the horizontal arms. The T-body is pigmented with barium sulphate and has a loop at one end of the vertical stem and two horizontal arms at the other end. Blue polypropylene removal threads are attached to the loop of the T-body.

KYLEENA is supplied in a sterile package. The vertical stem of KYLEENA is loaded in the insertion tube at the tip of the EvoInserter (Figure 11), which is used to insert KYLEENA into the uterine cavity. The arms of KYLEENA are pre-aligned in the horizontal position.

The EvoInserter consists of a symmetrical two-sided body and slider that are integrated with flange, 2-sided scale, lock, curved insertion tube and plunger. The narrow insertion tube has an outer diameter of 3.8 mm. The blue removal threads are contained within the insertion tube and the ergonomic handle, enabling single-handed loading. Once KYLEENA has been inserted, the inserter is to be discarded.

Figure 11: KYLEENA EvoInserter
PART II: SCIENTIFIC INFORMATION

PHARMACEUTICAL INFORMATION

Drug Substance

Proper name: Levonorgestrel
Chemical name: 18,19-Dinorpregn-4-en-20-yn-3-one, 13-ethyl-17-hydroxy-, (17α)-(-) (CAS)

Molecular formula: C_{21}H_{28}O_{2}
Molecular weight: 312.45

Structural formula:

Physicochemical properties: Levonorgestrel is a white to off-white crystalline powder, practically insoluble in water and slightly soluble in ethanol and acetone. The melting range is between 234°C and 239°C.

CLINICAL TRIALS

The contraceptive efficacy of KYLEENA (levonorgestrel-releasing intrauterine system [19.5 mg]) has been evaluated in a multi-centre (Europe, North America, and South America), open-label, randomized clinical study for 3 years with an extension to 5 years. The study included 1452 women aged 18-35 including 39.5% (574) nulliparous women of whom 84% (482) were nulligravid using KYLEENA. Approximately 40% of women in this study were 25 years or younger. Approximately 80% of women enrolled in the study were Caucasian, 11% Hispanic, 5% Black and 1% of Asian origin. The mean body mass index was 25.3 kg/m² (range 15.2 – 57.6 kg/m²). Women less than six weeks postpartum, with a history of ectopic pregnancy, with clinically significant ovarian cysts or with HIV or otherwise at high risk for sexually transmitted infections were excluded.

The 1 year Pearl Index was 0.16 and the Pearl Index after 5 years was 0.29. The failure rate was approximately 0.2% at 1 year and the cumulative failure rate was approximately 1.4% at 5 years. The failure rate also includes pregnancies due to undetected expulsions and perforations. KYLEENA provides more than 99% efficacy for each year of use. Contraceptive failure rate
with KYLEENA is <0.5% for each year of use. Kaplan-Meier failure rates associated with KYLEENA are provided in Table 6 below.

Table 6: Kaplan-Meier Failure Rates and Contraceptive Efficacy of KYLEENA

<table>
<thead>
<tr>
<th>Time point</th>
<th>Number of women</th>
<th>Number of pregnancies</th>
<th>Kaplan-Meier failure rate (%)</th>
<th>Failure Rate 95 % CI</th>
<th>Contraceptive efficacy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year 1</td>
<td>1,452</td>
<td>2</td>
<td>0.178</td>
<td>0.044–0.709</td>
<td>99.8%</td>
</tr>
<tr>
<td>Year 2</td>
<td>1,206</td>
<td>4</td>
<td>0.371</td>
<td>0.139–0.988</td>
<td>99.6%</td>
</tr>
<tr>
<td>Year 3</td>
<td>1,010</td>
<td>4</td>
<td>0.423</td>
<td>0.159–1.123</td>
<td>99.6%</td>
</tr>
<tr>
<td>Year 4</td>
<td>773</td>
<td>1</td>
<td>0.147</td>
<td>0.021–1.038</td>
<td>99.8%</td>
</tr>
<tr>
<td>Year 5</td>
<td>636</td>
<td>2</td>
<td>0.333</td>
<td>0.083–1.324</td>
<td>99.7%</td>
</tr>
<tr>
<td>5 years (Cumulative)</td>
<td>1,452</td>
<td>13</td>
<td>1.445</td>
<td>0.823–2.531</td>
<td>98.5%</td>
</tr>
</tbody>
</table>

The Kaplan-Meier failure rate reflects the probability of getting pregnant (%) after each year of use. With the ongoing risk of pregnancy over time, the cumulative failure rate increases as the period of analysis increases. Contraceptive efficacy is often expressed as 1- contraceptive failure rate. For example, the contraceptive efficacy of KYLEENA for the first year of use: 99.8% (1-0.00178 = 0.99822) (13).

Of the 1452 women treated with KYLEENA, 1445 (99.5%) had a successful insertion. The first insertion attempt was successful in 1390/1452 women (95.7%); the second attempt was successful in 55/57 (96.5 %) women. Study investigators assessed the insertion procedure as easy in 1302/1452 women (89.7 %). The insertion procedure was evaluated as easy in 84.3% and 93.2% of nulliparous and parous women, respectively.

Most women experienced either no pain (18.5%, 268/1452) or only mild pain (47.0 %, 683/1452) during insertion of KYLEENA. A total of 400 (27.5 %) women experienced moderate pain and 100 (6.9%) experienced severe pain. There was a trend towards less pain in parous women compared to nulliparous women.

In the pivotal study, 60% (870/1452) of women completed the planned treatment duration of 3 years with KYLEENA (14). Of the 707 women that continued in the 2 year extension phase, 78% (550/707 women) completed the overall treatment duration of 5 years (12). During the first 3 years, the majority of premature discontinuations were due to adverse events (278 subjects, 19.1%). In the 2 year extension phase, only 5.1% of premature discontinuations were due to an adverse event. Over the five years, the frequency of discontinuations due to progestogen-related side effects or any bleeding problems/abnormalities (including amenorrhea) was 3.1% and 5.2%, respectively.
General Information

The following table gives typical pregnancy rates for various forms of birth control, including no birth control. The reported rates represent the number of women out of 100 who would become pregnant within the first year of use (15).

Table 7: Reported Pregnancies per 100 Women Within the First Year of Use

<table>
<thead>
<tr>
<th>Method</th>
<th>Rate (per 100 women)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hormonal Intrauterine system (IUS)</td>
<td>less than 1</td>
</tr>
<tr>
<td>Copper Intrauterine device (IUD)</td>
<td>less than 1</td>
</tr>
<tr>
<td>Progesterone Injection</td>
<td>6</td>
</tr>
<tr>
<td>Combined hormonal contraceptive (pill, patch or ring)</td>
<td>9</td>
</tr>
<tr>
<td>Diaphragm</td>
<td>12</td>
</tr>
<tr>
<td>Male condom</td>
<td>18</td>
</tr>
<tr>
<td>Female condom</td>
<td>21</td>
</tr>
<tr>
<td>Sponge, spermicide</td>
<td>12-28</td>
</tr>
<tr>
<td>Withdrawal method</td>
<td>22</td>
</tr>
<tr>
<td>Natural family planning</td>
<td>24</td>
</tr>
<tr>
<td>No method</td>
<td>85</td>
</tr>
</tbody>
</table>

DETAILED PHARMACOLOGY

Human Pharmacology

See Part I: ACTION AND CLINICAL PHARMACOLOGY – Pharmacokinetics.

Pharmacodynamics

Levonorgestrel (LNG) is a 19-nortestosterone derivative with potent progestogenic effects, but no significant estrogenic activity.

In rabbits, evidence of transformation of the endometrium was observed after subcutaneous administration of 0.01 mg LNG corresponding to 2 mcg/kg/day. Transformative effects are also histologically recognizable in the rabbit endometrium when LNG is administered orally in doses ranging from 0.03 to 0.3 mg per animal corresponding to approximately 6 to 60 mcg/kg/day.

In pregnant rats, ovariectomized within the first 4 days after conception, the subcutaneous administration of 0.002 mg LNG had a blastocyst-maintaining effect. The antiestrogenic or progestogenic activity of levonorgestrel has also been demonstrated in various test models in rats and mice. The potency of LNG is significantly higher than progesterone and about 83 times stronger than chlormadinone acetate.

LNG does not have any significant estrogenic activity and androgenic effects are only detectable after large doses. LNG also influences the gonadotrophic function of the anterior lobe of the pituitary gland in all experimental tests.
Like other progestogens, relative large doses of LNG lead to increases in insulin secretion in rats and dogs.

**TOXICOLOGY**

Levonorgestrel (LNG) is widely used in gynecology in various ways: either in combination as the progestin component in oral contraceptives, and in hormonal replacement therapy or alone for contraception in minipills, subdermal implants and intrauterine systems. The toxicological profile of LNG after systemic and local administration is well known and the available evidence indicates that this hormone is a safe drug in the above-mentioned indications.

**Acute Toxicity**

Single dose toxicity studies with LNG in rats and mice using the oral, intraperitoneal and subcutaneous route of application indicate a low acute toxicity with lethal doses in the g/kg body weight range.

**Long-term Toxicity**

Systemic tolerance studies were performed in mice, rats, dogs and monkeys either with LNG or the racemate d,l-norgestrel. Systemic tolerance findings were as expected and in line with findings that have been described in these species after prolonged treatment with other progestins. Most findings could directly be related to the progestogenic activity of LNG on target organs or on the endocrine regulation. Furthermore, clinical and laboratory findings observed in one or more species, such as increased body weight gains, alterations in serum lipid (cholesterol or triglyceride) levels, increases in plasma fibrinogen, increases in blood sedimentation rate and the decreases in red blood cell parameters are also well known effects of repeated dosages of progestogens, including progesterone in the affected species.

The effect of LNG on the induction of mammary hyperplasia (acinar and/or ductal) in dogs is considered a species-specific effect. No compound-related alterations were observed in tumourigenicity studies with LNG in mice, rats and monkeys. In dogs, long-term oral treatment with the highest daily dose of 0.5 mg/kg LNG led to an increased incidence in benign mammary tumours. This is a well known effect of progestogens in dogs, the mechanism of which is assumed to be due to stimulation of growth hormone secretion that exerts a mammatrophic effect in this species. Endocrine regulation and the sensitivity of the mammary gland to progestogens and growth hormone are known to be unique for the dog, and have no predictive value for human risk assessment. Thus, taking into consideration biocompatibility and the absence of genotoxicity of the LNG-IUS and its components, the absence of proliferative or neoplastic changes in the local and systemic tolerance studies in rats and monkeys, and the extensive clinical experience with LNG and with another LNG-releasing intrauterine system (IUS) (52 mg), KYLEENA is regarded to possess no carcinogenic potential.

In chronic toxicity studies in monkeys over 9 months, intrauterine administered LNG-IUSs have been systemically and locally well tolerated. Exposure to the LNG-IUS induced typical pharmacological effects of LNG such as amenorrhea or reduced intensity and frequency of menses. Histological findings of the endometrium, which were observed both in the LNG-treated and placebo control animals, such as slight superficial hemorrhage, slight to moderate superficial
necrosis, slight inflammatory cell infiltration and slight to moderate focal hyperplasia, were considered to be due to the mechanical effect of the IUS. Further progestational effects, such as atrophy of endometrial glands accompanied by decidualization of endometrial stroma, are typical endocrine effects of LNG and not indicators of local intolerance.

**Reproductive Toxicology**

An early developmental formulation of KYLEENA was studied in rabbits following intrauterine administration. The absence of compound-related maternal toxicity, embryotoxicity, teratogenicity or effects on fetal development in pregnant rabbits suggests that KYLEENA is not a reproductive toxicant.

Reproduction toxicity tests were performed with LNG or norgestrel, for evaluation of effects on fertility and peri-/postnatal toxicity in rats and on embryotoxicity in mice, rats and rabbits. No adverse effects are to be expected at clinically relevant dose levels.

**Mutagenicity**

Mutagenicity tests were performed with LNG to detect gene mutations and chromosomal aberrations. Based on the results obtained from in vitro and in vivo tests, there is no evidence that LNG has a mutagenic potential. Furthermore, LNG has been tested into the potential to induce DNA-repair in rat hepatocytes in vitro (Unscheduled DNA Synthesis-test) and into its DNA-adduct forming potential in human liver slices in vitro. None of these tests gave an indication of a genotoxic potential of LNG.

Neither the LNG-IUS nor its components showed any evidence of a genotoxic potential in vitro or in vivo.

**Biocompatibility studies**

A wide range of biocompatibility tests were performed with the materials and components of KYLEENA. Many of the tests have been conducted during the development of other LNG-releasing IUSs (52 mg and 13.5 mg), which mainly consist of the same materials as KYLEENA. Therefore, studies conducted with other LNG-releasing IUSs (52 mg and 13.5 mg), or modified versions of it, are also relevant for KYLEENA. There was no evidence of any bioincompatibility including absence of a genotoxic potential of the components of KYLEENA. No systemic or local intolerance was detected in chronic studies in rats (intramuscular, subcutaneous) or Cynomolgus monkeys (intrauterine implantation). The endometrial effects of the modified intrauterine systems observed in monkey studies were similar to those seen in clinical studies.

It is unlikely that cytotoxic concentrations of silver ions can be achieved in vivo by the silver ring used in KYLEENA. Also, taking into consideration that there was no evidence of hemolytic potential or mutagenicity in vitro and no local or systemic intolerance in vivo after exposure to the silver ring for up to 39 weeks, there is no cause for concern for the use of the silver ring in KYLEENA for detection and differentiation by ultrasound.

Based on the results of the studies conducted with KYLEENA materials after ethylene oxide sterilization and considering guidance given by the ISO standard 10993, the limit of 2 ppm is considered to be toxicologically justified and not pose a risk to human health.
REFERENCES


KYLEENA®
Levonorgestrel-releasing intrauterine system

Read this carefully before you start taking KYLEENA and each time you get a refill. This leaflet is a summary and will not tell you everything about this drug. Talk to your healthcare professional about your medical condition and treatment and ask if there is any new information about KYLEENA.

**Serious Warnings and Precautions**

- **Hormonal Contraceptives including KYLEENA** DO NOT PROTECT against Sexually Transmitted Infections (STIs) including HIV/AIDS. To protect yourself against STIs you can use condoms with KYLEENA.

- Cigarette smoking increases the risk of serious side effects on the heart and blood vessels. You should talk to your healthcare professional about the risks of smoking.

- **KYLEENA may become attached to (embedded) or go through the wall of the uterus. This is called perforation. If you experience severe abdominal or pelvic pain with or without vaginal bleeding contact your healthcare professional immediately.**

**What is KYLEENA used for?**

KYLEENA is used to prevent pregnancy for up to 5 years.

**How does KYLEENA work?**

KYLEENA is an intrauterine system (IUS). It is placed in your uterus by your healthcare professional. KYLEENA works by slowly releasing a low amount of a hormone called levonorgestrel directly into the uterus every day. Levonorgestrel is similar to a sex hormone produced naturally by the body. It is also used in many birth control pills (the “Pill”).

KYLEENA may work to prevent pregnancy in several ways including:

- reducing the monthly thickening of the lining of the uterus,
- thickening the mucus in the cervix (this makes it harder for sperm to pass through the cervix), and
- impairing sperm movement and function.

Together, these actions prevent the sperm and egg from coming into contact and work together to prevent pregnancy.

KYLEENA does not contain any estrogen.

In clinical trials, there was less than 1 pregnancy per year for every 100 women using KYLEENA.

KYLEENA is a long acting reversible contraceptive (LARC). LARCs are highly effective in preventing pregnancy. They can be used for a long period of time and are easy to use.
Other Ways to Prevent Pregnancy

Other methods of birth control are available to you. When used properly, other methods of birth control work well enough for many women.

The table below shows the typical pregnancy rates for different methods of birth control. It also shows the pregnancy rate when no birth control is used. The rates show the number of women out of 100 who would become pregnant within the first year of use.

<table>
<thead>
<tr>
<th>Reported Pregnancies per 100 Women Within the First Year of Use</th>
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<tr>
<td>Hormonal Intrauterine system (IUS)</td>
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</tr>
<tr>
<td>Natural family planning</td>
</tr>
<tr>
<td>No birth control</td>
</tr>
</tbody>
</table>

The pregnancy rates listed in the table vary widely. This is because of differences in how carefully and regularly people use each method of birth control. Regular users may have lower pregnancy rates. Others may expect pregnancy rates in the middle ranges. This does not apply to IUDs because they are placed in the uterus and do not depend on user compliance.

What are the ingredients in KYLEENA?

Medicinal ingredients: levonorgestrel

Non-medicinal ingredients: barium sulphate, copper phthalocyanine, polydimethylsiloxane elastomer, polyethylene, polypropylene, silica colloidal anhydrous, silver.

KYLEENA comes in the following dosage form:

Intrauterine system (IUS): 19.5 mg levonorgestrel

Do not use KYLEENA if you have or have had any of the following conditions:

- allergies to the hormone levonorgestrel, or to any of the other ingredients of KYLEENA or to the components of the container
- are pregnant or think you may be pregnant
- currently have a pelvic infection [pelvic inflammatory disease (PID)] or have had one multiple times
- an untreated infection of your vagina and/or cervix
- an infection of your uterus after delivering a baby
- bleeding from the vagina that has not been explained
- a condition that changes the space inside your uterus, such as large abnormal growths (fibroids)
- abnormalities of the cells in the cervix (your healthcare professional can tell you if you have this)
• a known or suspected tumour that grows in response to progestogen (a hormone). This includes breast cancer
• liver disease or problems
• liver tumour
• an infection of your uterus after an abortion in the past 3 months
• cancer of the uterus or the cervix
• already have an IUS or IUD in your uterus
• tumours related to pregnancy
• an infection of the heart valves or lining of the heart (bacterial endocarditis)

To help avoid side effects and ensure proper use, talk to your healthcare professional before you take KYLEENA. Talk about any health conditions or problems you may have, including if you:
• are breastfeeding
• have given birth in the last 36 weeks
• have had a stroke, heart attack or any heart problems
• have an abnormality of your heart or if you have any problem with your heart valves
• have a history of blood clots (thrombosis)
• have a history of migraine, dizziness or blurred vision
• have severe headaches
• have or have had liver problems or jaundice (a yellowing of the skin, whites of the eyes and/or nails)
• are diabetic or have a family history of diabetes
• have or someone in your family has high blood pressure or abnormal blood lipid levels (blood fat levels)
• smoke
• have Systemic Lupus Erythematosus
• have a history of emotional problems such as depression
• have a family history of blood clots, heart attacks, or strokes

Other warnings you should know about:

If you see a different healthcare professional, inform him/her that you are using KYLEENA. You should inform your healthcare professional if you are scheduled for Magnetic Resonance Imaging (MRI). KYLEENA can be safely scanned with MRI under most standard conditions.

Tell your healthcare professional if you are scheduled for any lab tests. Certain tests may be affected by hormonal birth control. Also, tell your healthcare professional if you are scheduled for surgery requiring prolonged bed rest.

KYLEENA should be used only under the supervision of your healthcare professional. Visit your healthcare professional 4 to 12 weeks after you have KYLEENA placed. You should have a follow-up visit at least once a year. Your visits may include a blood pressure check, breast exam, pelvic exam (including a Pap smear) and an abdominal exam. Carefully follow all directions given to you. Otherwise, you may become pregnant.

If you and your healthcare professional decide that, for you, the benefits of KYLEENA outweigh the risks, you should be aware of the following:
1. **Diabetes**

If you have diabetes and use KYLEENA, your blood sugar levels should be watched closely.

2. **Infections**

In the first 3 weeks after placement of an IUS/IUD, there is an increased risk of a serious pelvic infection called pelvic inflammatory disease (PID). Other known risk factors for PID are having multiple sexual partners, frequent intercourse, sexually transmitted infections (STIs) and young age.

PID can cause serious problems such as difficulty getting pregnant, development of a pregnancy outside the uterus (ectopic pregnancy), or pelvic pain that does not go away. PID is usually treated with antibiotics. More serious cases of PID may require surgery.

Tell your healthcare professional right away if you have signs of PID. These can include long-lasting or heavy bleeding, unusual vaginal discharge, pain in your lower stomach area, painful sex, chills or fever.

3. **Ectopic Pregnancy**

While using KYLEENA, the risk of pregnancy is low. However, if you get pregnant while using KYLEENA, it is more likely that the pregnancy will be ectopic. This means that the pregnancy is not in the uterus. Ectopic pregnancy is a serious condition. Tell your healthcare professional immediately if you have lower abdominal pain, especially if you have missed a period and/or have unexpected bleeding. These can be signs of an ectopic pregnancy.

4. **Cysts on the Ovary**

Cysts (fluid filled sacs) on the ovary commonly occur in women using KYLEENA. These cysts usually disappear within a few months on their own. However, cysts can sometimes cause pain and may need medical attention.

5. **Uterine Perforations**

KYLEENA may become attached to or go through the wall of the uterus. This is called perforation and it happens most often during placement. Perforation is uncommon. If this happens, KYLEENA must be removed.

The risk of perforation is higher in women who are breastfeeding when KYLEENA is placed. The risk is also higher when placement takes place up to 36 weeks after a delivery. The risk of perforation may be increased in women with a mis-shaped uterus or a uterus that is fixed and leans backward.

6. **Use While Breastfeeding**

You can use KYLEENA when you are breastfeeding. Small amounts of the hormone in KYLEENA have been found in the breast milk of women using a different IUS. This does not appear to affect the health of your baby when you start using KYLEENA 6 weeks after delivery. Levonorgestrel does not appear to affect the amount or the quality of breast milk. However, isolated cases of decreased milk production have been reported among women using KYLEENA.

7. **Use in Pregnancy**

If you become pregnant with KYLEENA in place, you should have it removed as soon as possible. If it is left in place during pregnancy, the chances of having a miscarriage or early delivery increase. It is not known if KYLEENA can cause long-term effects on the baby if it stays in place during pregnancy. Removing KYLEENA may cause a miscarriage. Talk with your healthcare professional about the benefits and risks of continuing the pregnancy.
8. **Use After Pregnancy and Abortion**

After childbirth, KYLEENA should be placed only when the uterus has returned to normal size. KYLEENA should not be placed for at least 6 weeks after delivery.

KYLEENA can be placed right after an abortion which took place in the first three months of pregnancy. If an abortion takes place in the second trimester, placement of KYLEENA should be delayed for 6 weeks or until the uterus has returned to normal size.

9. **Pregnancy After Stopping KYLEENA**

If you wish to become pregnant, ask your healthcare professional to remove KYLEENA. Your usual fertility level should return after removal.

10. **Breast Self-Examination**

Your healthcare professional should show you how to do breast self-examinations when you are using KYLEENA. If you find any lumps or see any other changes talk to your healthcare professional immediately.

11. **Broken KYLEENA**

KYLEENA may break, most often during a difficult removal. Broken pieces must be found and removed. Surgery may be needed to do this.

**Driving or Using Machines**

The effect of KYLEENA on the ability to drive or to use machines has not been studied. Do not drive or use machines until you know how you react to KYLEENA.

**How Will KYLEENA Affect My Periods?**

KYLEENA may affect your menstrual cycle. For the first 3 to 6 months, your period may become irregular and the number of bleeding days may increase. You may also have frequent spotting or light vaginal bleeding. Some women have heavy or prolonged bleeding during this time. Contact your healthcare professional if this persists.

Overall, you are likely to have a gradual reduction in the amount and number of days of bleeding each month. Some women eventually find that periods stop altogether.

When KYLEENA is removed, periods soon return to normal.

**What if I Stop Having Periods?**

Over time, your menstrual period may gradually disappear when using KYLEENA. This is because of the effect of the hormone on the lining of the uterus. The normal monthly thickening of the uterine lining with blood does not happen; therefore, there is little or no bleeding, as happens during a usual menstrual period. It does not necessarily mean you have reached menopause or are pregnant.

If, however, you are having regular menstrual periods and then do not have one for 6 weeks or longer, it is possible that you may be pregnant. You should speak to your healthcare professional.
Tell all healthcare professionals about all the medicines you take. This includes KYLEENA and any other drugs, vitamins, minerals, natural supplements or alternative medicines.

Certain drugs may interact with KYLEENA. They can make it less able to prevent pregnancy. Hormonal birth control may become less reliable if you are taking drugs that affect your liver (such as primidone, barbiturates, phenytoin, carbamazepine, and rifampicin). KYLEENA is not likely to be affected by these drugs. This is because it releases a very small amount of hormone directly inside your uterus. If you are taking other medications, ask your healthcare professional if you need to use an additional method of birth control.

KYLEENA is visible during X-ray exams. KYLEENA also contains a small silver ring that makes it visible by ultrasound.

How to take KYLEENA:

What it looks like:
KYLEENA consists of a small, white, T-shaped body made from soft, flexible plastic. The vertical and horizontal arms of the T-body are about 3 cm long. The vertical arm is surrounded by a narrow, cylindrical shaped drug reservoir that contains levonorgestrel. Two fine blue plastic threads are attached to the tip of the vertical arm. These threads are used for removal of the system. The threads can also be used to check if KYLEENA is in place. In addition, the vertical stem contains a silver ring located close to the horizontal arms, which is visible under ultrasound. KYLEENA is packaged with the EvoInserter (which is used to place KYLEENA).

How is KYLEENA Placed?
Before KYLEENA is placed, you will have a pelvic exam to determine the position and size of your uterus. You may also have a Pap smear and a breast exam. If necessary, you may also have other tests (ie, for infections including STIs).

Your healthcare professional will place the thin flexible plastic tube of the insertion device containing KYLEENA into your uterus. Once KYLEENA is in the correct position, your healthcare professional will withdraw the tube and leave KYLEENA in your uterus. Your healthcare professional will trim the removal threads to a suitable length. The procedure takes only a few minutes.
Most women experience either no pain or only mild pain during placement of KYLEENA. Some women may experience some discomfort. You may wish to discuss the need for a painkiller or local anesthetic with your healthcare professional.

Some women may feel dizzy after KYLEENA is placed. This feeling goes away after a short rest. The placement may cause a seizure in patients who have epilepsy. You may also experience some bleeding during or just after placement. After placement you may feel some cramp-like menstrual pain; however, this usually stops within a few days.

It is not likely, but KYLEENA may go through the wall of the uterus during placement and come to rest outside the uterus. If this happens KYLEENA must be removed.

**When Should KYLEENA be Placed?**

KYLEENA should be placed within 7 days of the start of your period. In this case, no back up birth control is needed. If it is certain that you are not pregnant, KYLEENA may also be placed at any other time during your cycle. Tell your healthcare professional if you have had unprotected sex since your last period. If KYLEENA is placed more than 7 days since your period started, use a condom or diaphragm, or do not have sex for the next 7 days. KYLEENA cannot be used as emergency birth control.

When replacing an existing KYLEENA for a new one, it is not necessary to wait for your period. A new KYLEENA can be placed at any time of the cycle.

**How Quickly Does KYLEENA Start to Work?**

KYLEENA starts to work right away if it is placed within 7 days of the start of your period. It is best to wait 24-48 hours before having intercourse in case of discomfort. If KYLEENA is placed more than 7 days after the start of your period, use a condom or diaphragm for the next 7 days. Alternatively, do not have sex for the next 7 days.

**How Often Should I Have KYLEENA Checked?**

You should have KYLEENA checked about 4 to 12 weeks after it is placed, again at 12 months and then once a year until it is removed. KYLEENA can stay in place for up to 5 years before it must be removed. You should receive a Patient Reminder Card from your healthcare professional after the placement of KYLEENA. Keep this card until KYLEENA is removed and bring it with you to every medical appointment.

**How Can I Check if KYLEENA is in Place?**

After each menstrual period or about once a month, you should check by feeling if the threads are still in place. Your healthcare professional will show you how to do this. Do not pull on the threads as you may accidentally pull KYLEENA out.

If you cannot feel the threads, this may mean that KYLEENA has fallen out (expulsion) or gone through the uterus (perforation). See your healthcare professional and in the meantime, use another method of birth control that does not include hormones. You should also see your healthcare professional if you can feel the lower end of KYLEENA itself.
Will KYLEENA be felt during Sexual Intercourse?
You and your partner should not be able to feel KYLEENA during intercourse. If you can feel KYLEENA or if you feel any pain, avoid having sex until your healthcare professional has checked if KYLEENA is still in the correct position.

The removal threads may be felt by your partner during intercourse.

Can Tampons be Used?
Use of sanitary pads is recommended. If tampons are used, you should change them carefully so as not to pull the threads of KYLEENA.

Can KYLEENA Fall Out?
Although unlikely, KYLEENA may fall out on its own. This is called expulsion. If this happens, you are not protected against pregnancy. An unusual increase in the amount of bleeding during your period might be a sign that this has happened. If you think KYLEENA has fallen out, use a non-hormonal method of birth control and talk to your healthcare professional.

Removal of KYLEENA
KYLEENA should not be left in place for more than 5 years. You should see your healthcare professional when you want to have KYLEENA taken out. Removal of KYLEENA is usually very easy.

You could become pregnant as soon as KYLEENA is removed, so you should use another method of birth control if you do not want to become pregnant. Talk to your healthcare professional about the best birth control methods for you, because your new method may need to be started 7 days before KYLEENA is removed to prevent pregnancy.

If you wish to continue using KYLEENA after 5 years, your healthcare professional can place a new system immediately after removing the old system. If the same KYLEENA system has been left in place for longer than 5 years, you may become pregnant. Pregnancy should be ruled out before placement of a new system.

What are possible side effects from using KYLEENA?
These are not all the possible side effects you may feel when taking KYLEENA. If you experience any side effects not listed here, contact your healthcare professional. Please also see the box called “Serious Warnings and Precautions.”

Side effects with KYLEENA are more common during the first months of use; they gradually decrease over time. Differences in your menstrual bleeding are the most common side effects of KYLEENA during the first months after the system is placed, but these effects should decrease over time.

Very common side effects (may affect more than 1 in 10 women):
- acne/oily skin

Common side effects (may affect up to 1 in 10 women):
- nausea
- hair loss
- infection of the ovaries, fallopian tubes or uterus
- painful menstruation
- breast pain/discomfort
- discharge from the vagina

Uncommon side effects (may affect up to 1 in 100 women):
- excess hair on the face, chest, stomach or legs

Women using a different IUS have reported cases of the following events. Talk with your healthcare professional if any of these occur:
- blood clots in the lungs (symptoms may include sharp pain in the chest, shortness of breath, coughing up blood), blood clot in the leg (symptoms may include pain, warmth, redness and swelling in the leg) or blood clot in the brain (stroke) (symptoms may include sudden severe headache, vomiting, dizziness, fainting, problems with vision or speech, weakness or numbness in the face, arm or leg)
- breakage of the IUS
- increased blood pressure

Few women using KYLEENA after delivery have reported less milk production.

### SERIOUS SIDE EFFECTS AND WHAT TO DO ABOUT THEM

<table>
<thead>
<tr>
<th>Symptom/ effect</th>
<th>Talk with your Healthcare Professional</th>
<th>Get immediate medical help</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Only if severe</td>
<td>In all cases</td>
</tr>
<tr>
<td><strong>VERY COMMON</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Vaginal bleeding changes:</strong> increased or decreased menstrual bleeding, spotting, infrequent periods or absence of bleeding</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>Vaginal Infection</strong> (inflammation of the vagina or surrounding area): itching, or unusual or increased discharge from the vagina</td>
<td></td>
<td>✓</td>
</tr>
<tr>
<td><strong>Headache</strong></td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>Abdominal pain/ pelvic pain</strong></td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>Cysts on the ovaries:</strong> Pelvic pain, painful intercourse, abdominal bloating or swelling, pain during bowel movements. (Cysts usually disappear on their own within a few months and may not show symptoms. Serious cysts are uncommon.)</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>COMMON</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Expulsion (KYLEENA has fallen out):</strong> An unusual increase in the amount of bleeding during your period</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>Migraine</strong></td>
<td>✓</td>
<td></td>
</tr>
</tbody>
</table>
### SERIOUS SIDE EFFECTS AND WHAT TO DO ABOUT THEM

<table>
<thead>
<tr>
<th>Symptom/ effect</th>
<th>Talk with your Healthcare Professional</th>
<th>Get immediate medical help</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Only if severe</td>
<td>In all cases</td>
</tr>
<tr>
<td><strong>Depressed Mood/Depression:</strong> Sad mood that won’t go away. If you have a history of depression, your depression may become worse</td>
<td></td>
<td>✓</td>
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<tr>
<td><strong>UNCOMMON</strong></td>
<td></td>
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<tr>
<td><strong>Pelvic Infection:</strong> Constant pain in the lower stomach area, together with fever or unusual discharge from the vagina</td>
<td>✓</td>
<td></td>
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<tr>
<td><strong>Intrauterine pregnancy, miscarriage, or ectopic pregnancy:</strong> Constant pain in the lower stomach area, together with nausea or breast tenderness and/or vaginal bleeding</td>
<td>✓</td>
<td></td>
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<tr>
<td><strong>Perforation of the uterus</strong> [attached to (embedded) or gone through the wall of the uterus]: Severe lower abdominal pain which may be together with bleeding</td>
<td>✓</td>
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<tr>
<td><strong>VERY RARE</strong></td>
<td></td>
<td></td>
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<tr>
<td><strong>Allergic reaction</strong> including itchiness, rash, swelling of face and lips, cheeks, tongue and/or throat</td>
<td></td>
<td>✓</td>
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</table>

If you have a troublesome symptom or side effect that is not listed here or becomes bad enough to interfere with your daily activities, talk to your healthcare professional.
Reporting Side Effects
You can help improve the safe use of health products for Canadians by reporting serious and unexpected side effects to Health Canada. Your report may help to identify new side effects and change the product safety information.

3 ways to report:
- Online at MedEffect (www.healthcanada.gc.ca/medeffect);
- By calling 1-866-234-2345 (toll-free);
- By completing a Consumer Side Effect Reporting Form and sending it by:
  - Fax to 1-866-678-6789 (toll-free), or
  - Mail to: Canada Vigilance Program
            Health Canada, Postal Locator 1908C
            Ottawa, ON
            K1A 0K9

Postage paid labels and the Consumer Side Effect Reporting Form are available at MedEffect.

NOTE: Contact your healthcare professional if you need information about how to manage your side effects. The Canada Vigilance Program does not provide medical advice.

Storage
Store at controlled room temperature (between 15°C and 30°C).
Keep out of reach and sight of children.

If you want more information about KYLEENA:
- Talk to your healthcare professional
- Find the full product monograph that is prepared for healthcare professional and includes this Patient Medication Information by visiting the Health Canada website (www.healthcanada.gc.ca); the manufacturer’s website http://www.bayer.ca or by contacting Bayer at 1-800-265-7382.

This leaflet was prepared by:

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