PRODUCT INFORMATION

Implanon® Implant

NAME OF THE MEDICINE

Etonogestrel sub-dermal implants.

Etonogestrel is a white to nearly white crystalline powder also known as 3-ketodesogestrel or (17α)-13-ethyl-17-hydroxy-11-methylene-18,19 dinorpregn-4-en-20-yn-3-one.

Molecular formula $C_{22}H_{28}O_2$ Molecular mass 324.44 CAS No. 54048-10-1

DESCRIPTION

Implanon is a sub-dermal contraceptive implant, consisting of a co-axial rod, placed inside an applicator. The rod consists of a core containing a mixture of the drug substance, etonogestrel, and ethylene vinylacetate copolymer and a skin consisting of ethylene vinylacetate.

Each rod has a length of 4.0 cm and a diameter of 2.0 mm and is located inside the stainless steel part of the cannula of the applicator.

**Implant**
- Core: Ethylene vinylacetate copolymer (28% vinyl acetate) 46 mg - proprietary ingredient
- Skin: Ethylene vinylacetate copolymer (14% vinyl acetate) 15 mg - proprietary ingredient

**Applicator**
Acrylonitrile-butadiene-styrene body with a stainless steel needle and a polypropylene needle shield.

**Qualitative and Quantitative Composition**
One implant contains 68 mg etonogestrel; the release rate is 60-70 $\mu$g/day during week 5-6, and decreases to approximately 35-45 $\mu$g/day at the end of the first year, to approximately 30-40 $\mu$g/day at the end of the second year, and to approximately 25-30 $\mu$g/day at the end of the third year.

PHARMACOLOGY

**Pharmacodynamic properties**
Implanon is a non-biodegradable, etonogestrel (ENG) containing implant for subdermal use. ENG is the biologically active metabolite of desogestrel, a progestagen widely used in oral contraceptives (OCs). It is structurally derived from 19-nortestosterone and binds with high affinity to progesterone receptors in the target organs. The contraceptive effect of Implanon is achieved primarily by inhibition of ovulation. Ovulations were not observed in the first two years of use and only rarely in the third year. Besides inhibition of ovulation Implanon also causes changes in the cervical mucus, hindering the passage of spermatozoa. The three-year Pearl Index is 0.00 (95% Confidence Interval 0.00 – 0.18). This high degree of protection against pregnancy is obtained among other reasons because, in
contrast to OCs, the contraceptive action of Implanon is not dependent on the regular intake of pills. The contraceptive action of Implanon is reversible, which is apparent by a rapid return of a normal menstrual cycle after removal of the implant.

Although Implanon inhibits ovulation, ovarian activity is not completely suppressed. Mean oestradiol concentrations remain above the level seen in the early-follicular phase. In a two-year study, in which the bone mineral density in 44 Implanon users has been compared with that in a control group of 29 IUD-users no adverse effects on bone mass have been observed. During the use of Implanon no clinically relevant effects on lipid metabolism have been observed. The use of progestagen-containing contraceptives may have an effect on insulin resistance and glucose tolerance.

In addition, in clinical trials it was shown that Implanon users often have a less painful menstrual bleeding.

Pharmacokinetic properties

Absorption
After insertion of Implanon, ENG is rapidly absorbed into the blood stream. Ovulation-inhibiting concentrations are reached within 1 day. Maximum serum concentrations (between 472 and 1270 pg/mL) are reached within 1 to 13 days. The release rate of the implant decreases with time. As a result serum concentrations decline rapidly over the first few months. By the end of the first year a mean concentration of approximately 200 pg/mL (range 150-261 pg/mL) is measured, which slowly decreases to 156 pg/mL (range 111-202 pg/mL) by the end of the third year. The variations observed in serum concentrations can be partly attributed to differences in body weight.

Distribution
ENG is 95.5-99% bound to serum proteins, predominantly to albumin and to a lesser extent to SHBG. The central and total volume of distribution are approximately 27 l and 220 l respectively and hardly change during the use of Implanon.

Metabolism
ENG is hydroxylated, reduced and conjugated to sulfates and glucuronides. *In vitro* data provide evidence that metabolism of etonogestrel, similar to that of other contraceptive steroids, is catalysed by CYP3A4 (See 'Interaction with other medicines').

Elimination
After IV administration of ENG, the mean elimination half-life is approximately 25 hours and the serum clearance is approximately 7.5 l/hour. Both clearance and elimination half-life remain constant during treatment period. The excretion of ENG and its metabolites, either as free steroids or as conjugates, is in urine and faeces (ratio 1.5:1). After insertion of Implanon in lactating women, etonogestrel is excreted in breast milk with a milk/serum ratio of 0.44-0.50 during the first four months. In lactating women using Implanon, the mean transfer of etonogestrel to the infant is approximately 2.2% of the maternal etonogestrel daily dose (values normalized per kg body weight). Concentrations show a gradual and statistically significant decrease from about 20 to 15 and 10 ng/kg/day at month 1, 2 and 4 respectively in a group of 41 infants.

Preclinical safety data
Toxicological studies did not reveal any effects other than those which can be explained based on the hormonal properties of ENG, regardless of the route of administration.

CLINICAL TRIALS

Three pivotal efficacy and safety studies were performed in healthy, fertile and sexually active women. The single primary endpoint was pregnancy and as secondary endpoints the following parameters were studied: ovulation, weight, time required for Implanon insertion and removal, and laboratory variables. Following the removal of Implanon return to normal ovulation was investigated. In these pivotal studies a total number of 1286 subjects using Implanon were studied. Total Implanon exposure was 2093 Women Years (27322, 28 day cycles). Pregnancies did not occur in the pivotal or supportive studies. The Pearl Index is essentially 0.00 (0.00 – 0.18 95% CI). Contraceptive efficacy is satisfactory for a period of three years. The data demonstrate that Implanon is a highly efficacious contraceptive product mainly by virtue of its very efficient suppression of ovulation evoked by the continuous release
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of the drug substance etonogestrel. The safety profile is consistent with the well-known pharmacological profile of etonogestrel.

One two year study was performed in 46 women receiving Implanon versus 33 women with an IUD to examine the effect of Implanon use on bone mineral density parameters. The changes in bone mineral density parameters were not different from a comparator IUD group and the mean bone mineral density parameters at several sites of the body were generally higher than those reported for a standard reference population. Oestriadiol levels were above the threshold level for maintaining normal bone mass.

A body weight increase of about 1.5% per year was found for Implanon as well as for the IUD comparator. The increase is therefore only partly attributable to the use of Implanon.

In supportive studies return of ovulation after implant removal was assessed by ultrasound measurements and hormone determinations. Ovulation returns after removal of the implant shortly after etonogestrel has disappeared from the body, enabling restoration of fertility. This conclusion is supported by the occurrence of 37 pregnancies that were reported after implant removal.

INDICATIONS

Contraception
(removed and replaced every three years to ensure continued contraceptive efficacy)

CONTRAINDICATIONS

- Known or suspected pregnancy.
- Active thromboembolic disorders.
- Presence or history of severe hepatic disease as long as liver function values have not returned to normal.
- Progestagen dependent tumour (known or suspected).
- Known or suspected carcinoma of the breast.
- Benign or malignant liver tumours.
- Undiagnosed vaginal bleeding.
- Hypersensitivity to the active substance or any of the excipients of Implanon.

PRECAUTIONS

Medical Examination and Follow Up

Prior to the initiation or reinstitution of Implanon a complete medical history (including family medical history) should be taken and pregnancy should be excluded. Blood pressure should be measured and a physical examination should be performed, guided by the CONTRAINDICATIONS and PRECAUTIONS. It is recommended that the woman returns for a medical check-up three months after insertion of Implanon. During this check-up, the blood pressure should be measured and an enquiry should be made after any questions, complaints or the occurrence of undesirable effects. The frequency and nature of further periodic checks should be adapted to the individual woman, guided by clinical judgement, but at least once a year is advised.

Other

Women should be told that Implanon does not offer protection against HIV (AIDS) and other sexually transmitted diseases.

Use with caution in the following circumstances

The user should be informed about the pros and the cons of an implant compared to other contraceptive methods before the insertion of Implanon. If any of the conditions/risk factors mentioned below are present, the benefits of progestagen use should be weighed against the possible risks for each individual case and discussed with the woman before she decides to start with Implanon. In the event of aggravation, exacerbation or the first appearance of any of these conditions, the woman should contact her physician. The physician should then decide whether the use of Implanon should be discontinued.
The risk for breast cancer increases in general with increasing age. During the use of OCs the risk of having breast cancer diagnosed is slightly increased. This increased risk disappears gradually within 10 years after discontinuation of OC use and is not related to the duration of use, but to the age of the woman when using the OC. The expected number of cases diagnosed per 10,000 women who use combined OCs (up to 10 years after stopping) relative to never users over the same period have been calculated for the respective age groups to be: 4.5/4 (16-19 years), 17.5/16 (20-24 years), 48.7/44 (25-29 years), 110/100 (30-34 years), 180/160 (35-39 years) and 260/230 (40-44 years). The risk in users of contraceptive methods which only contain progestagens is possibly of similar magnitude as that associated with combined OCs. However, for these methods, the evidence is less conclusive. Compared to the risk of getting breast cancer ever in life, the increased risk associated with OCs is low. The cases of breast cancer diagnosed in OC users tend to be less advanced than in those who have not used OCs. The increased risk observed in OC users may be due to an earlier diagnosis, biological effects of the OC or a combination of both. Since a biological effect of hormones cannot be excluded, an individual benefit/risk assessment should be made in women with pre-existing breast cancer and in women in whom breast cancer is diagnosed while using Implanon.

Since a biological effect of progestagens on liver cancer cannot be excluded an individual benefit/risk assessment should be made in women with liver cancer.

Epidemiological studies have associated the use of combined oral contraceptives (COCs) with an increased incidence in venous thromboembolism (VTE, deep vein thrombosis and pulmonary embolism). Although the clinical relevance of this finding for etonogestrel (the biologically active metabolite of desogestrel) used as a contraceptive in the absence of an oestrogenic component is unknown, Implanon should be removed in the event of a thrombosis. Removal of Implanon should also be considered for women who are immobilised for a long time because of surgery or a disease. Women with venous thromboembolic disease should be made aware of the possibility of a recurrence.

The following conditions have been reported both during pregnancy and during sex steroid use, but an association with the use of progestagens has not been established: jaundice and/or pruritus related to cholestasis; gallstone formation; porphyria; systemic lupus erythematosus; haemolytic uraemic syndrome; Sydenham’s chorea; herpes gestationis; otosclerosis-related hearing loss.

When acute or chronic disturbances of liver function occur, the woman should be referred to a specialist for examination and/or advice.

Chloasma may occasionally occur, especially in women with a history of chloasma gravidarum. Women with a tendency to develop chloasma should avoid exposure to the sun or ultraviolet radiation whilst using Implanon.

Although progestagens may have an effect on peripheral insulin resistance and glucose tolerance, there is no evidence for a need to alter the therapeutic regimen in diabetics using progestagen-only contraceptives. However, diabetic women should be carefully observed while using progestagen-only contraceptives.

Implanon use should be discontinued when during the use of Implanon there is a constantly elevated blood pressure or when a significant increase in blood pressure does not respond to an antihypertensive therapy.

Expulsion may occur especially if the implant is not inserted according to the instructions given in the section ‘How to insert Implanon’, or as a consequence of a local inflammation.

Occasionally a scar may be formed.

In rare cases, mostly related to either a too deep initial insertion (refer also to ‘How to insert Implanon’) and/or to external forces (e.g. manipulation of the implant or contact sports) the implant may migrate from the insertion site. In these cases localization of the implant may be more difficult and removal may require a larger incision (refer also to ‘How to remove Implanon’). If Implanon cannot be found, contraception and the risk of progestogen-related undesirable effects may continue beyond the time desired by the woman.
• The efficacy of Implanon may be reduced when concomitant medication is used (see ‘Interactions with other medicines’).

• The contraceptive effect of Implanon is related to the plasma levels of etonogestrel, which are inversely related to body weight and decrease with time after insertion. The clinical experience with Implanon in heavier women in the third year of use is limited. Therefore it cannot be excluded that the contraceptive effect in these women during the third year of use may be lower than for women of normal weight. Clinicians may therefore consider earlier replacement of the implant in heavier women.

Changes in vaginal bleeding pattern
During the use of Implanon vaginal bleeding may become more frequent or of longer duration in most women. In others, bleeding may become incidental or be totally absent (approximately 1 out of 5 women).† These changes are reasons for women to reject the method. Acceptance of bleeding pattern can be improved by offering women who have chosen to use Implanon careful counselling on this aspect. Evaluation of vaginal bleeding should be done on an ad hoc basis and may include an examination to exclude gynaecological pathology or pregnancy.

Follicular development
With all low-dose hormonal contraceptives, follicular development occurs and occasionally the follicle may continue to grow beyond the size it would attain in a normal cycle. Generally, these enlarged follicles disappear spontaneously. Often, they are asymptomatic; in some cases they are associated with mild abdominal pain and rarely, they require surgical intervention.

Ectopic pregnancy
The protection with traditional progestagen-only contraceptives against ectopic pregnancies is not as good as with combined OCs, which has been associated with the frequent occurrence of ovulations during the use of these methods. Despite the fact that Implanon consistently inhibits ovulation, ectopic pregnancy should be taken into account in the differential diagnosis if the woman gets amenorrhoea or abdominal pain.

Carcinogenicity and mutagenicity
No drug-related increases in tumour incidences were observed in rats with subcutaneous implants releasing 18 or 36 micrograms of etonogestrel/day. Serum etonogestrel levels in the high-dose group were 3 times higher than those in women with Implanon implants. Mutagenic activity was not observed in bacterial cells (Ames assay), but mutagenic and clastogenic activities in mammalian cells have not been investigated.

Pregnancy: Category B3
The use of Implanon is contraindicated during pregnancy. If pregnancy occurs during use of Implanon, the implant should be removed.†

Animal studies have shown that high doses of progestagens can cause masculinisation of the female fetus. However, in animal studies of etonogestrel, no embryotoxic or fetotoxic effects were seen in rats or rabbits at oral doses up to 2 mg/kg/day. Plasma drug levels were not measured in either study, but it can be estimated that systemic exposure at the high dose level in the rat study was 1.6 to 3 times higher than in women with Implanon implants. There are insufficient data on the use of Implanon during pregnancy in humans to evaluate possible harmful effects during a possible pregnancy. So far there are no indications for an increased risk of birth defects of children born to women using COC or progestagen-only contraceptives prior to pregnancy. Neither is there any indication for teratogenic defects in cases where a progestagen-only contraceptive was used in women not knowing of their pregnancy. The relevance of this to Implanon has not been confirmed yet.

Lactation †
In an open, non-randomised comparative study of Implanon (n=42) vs IUD (n=38) in healthy lactating women, Implanon was shown not to influence the production or the quality (protein, lactose or fat concentrations) of breast milk. However, small amounts of etonogestrel are excreted in breast milk. Based on an average daily milk ingestion of 150 ml/kg, the mean daily infant etonogestrel dose calculated after one month of etonogestrel release is approximately 27 ng/kg/day. This corresponds to
approximately 0.2% of the estimated absolute maternal daily dose (2.2% when values are normalized per kg body weight). Subsequently the milk etonogestrel concentration decreases with time during the lactation period. Long-term data are available on 38 children, whose mothers started using Implanon during the 4th to 8th week postpartum. They were breast-fed for a mean duration of 14 months and followed up to 36 months of age. Evaluation of growth, and physical and psychomotor development did not indicate any differences in comparison to nursing infants whose mothers used an IUD (n=33). Nevertheless, development and growth of the child should be carefully followed. Based on the available data, Implanon may be used during lactation.

Interaction with other medicines†
Interactions between hormonal contraceptives and other medicinal products may lead to breakthrough bleeding and / or contraceptive failure. No specific interaction studies have been performed with Implanon. The following interactions have been reported in the literature (mainly with combined contraceptives but occasionally also with progestogen-only contraceptives).

Hepatic metabolism: Interactions can occur with medicinal products that induce microsomal enzymes, specifically cytochrome P450 enzymes, which can result in increased clearance of sex hormones (e.g., phenytoin, barbiturates, primidone, carbamazepine, rifampicin, and possibly also oxcarbazepine, rifabutin, topiramate, felbamate, ritonavir, nelfinavir, griseofulvin and the herbal remedy St. John's wort). Women on treatment with any of these drugs should temporarily use a barrier method in addition to Implanon. With microsomal enzyme-inducing drugs, the barrier method should be used during the time of concomitant drug administration and for 28 days after their discontinuation. In women on long-term treatment with hepatic enzyme-inducing drugs, it is recommended to remove Implanon and to advise a contraceptive method that is unaffected by the interacting drug.

Hormonal contraceptives may interfere with the metabolism of other drugs. Accordingly, plasma and tissue concentrations may be affected (e.g., cyclosporin). Note: The prescribing information of concomitant medications should be consulted to identify potential interactions.

Effects on laboratory tests
Data on COCs have shown that contraceptive steroids may influence the results of certain laboratory tests, including biochemical parameters of liver, thyroid, adrenal and renal function, serum levels of (carrier) proteins, e.g., corticosteroid binding globulin and lipid/lipoprotein fractions, parameters of carbohydrate metabolism and parameters of coagulation and fibrinolysis. The changes were generally within the normal range. To what extent this also relates to progestagen-only contraceptives is not known.

Effects on ability to drive and use machines
No observed effects.

ADVERSE REACTIONS
Serious undesirable effects
Refer to Precautions

Other possible undesirable effects
The following adverse effects have been reported during the use of Implanon. An association has been neither confirmed nor refuted. Some of these effects have also been occasionally reported with progestagen-only contraceptives.
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**Table:** Percentages of subjects with at least one experience classified by body system and reported as related to the study drug pre-marketing in clinical trials performed by Schering-Plough\(^1\).  

<table>
<thead>
<tr>
<th>Body System (WHO System Organ Class)</th>
<th>Implanon(^2) Related AE’s (&gt; 2.5%) N= 1326</th>
<th>Norplant(^2) (levonorgestrel releasing implants) Related AE’s (&gt; 5%) N= 184</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reproductive disorders, female</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast Pain</td>
<td>9.8</td>
<td>11.4</td>
</tr>
<tr>
<td>Dysmenorrhea</td>
<td>4.4</td>
<td>3.8</td>
</tr>
<tr>
<td>Skin appendage disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acne</td>
<td>14.8</td>
<td>21.1</td>
</tr>
<tr>
<td>Alopecia</td>
<td>2.5</td>
<td>3.3</td>
</tr>
<tr>
<td>Central and peripheral system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>13.3</td>
<td>20.1</td>
</tr>
<tr>
<td>Dizziness</td>
<td>3.8</td>
<td>7.1</td>
</tr>
<tr>
<td>Gastrointestinal system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>4.7</td>
<td>8.2</td>
</tr>
<tr>
<td>Nausea</td>
<td>2.5</td>
<td>5.4</td>
</tr>
<tr>
<td>Flatulence</td>
<td>2.6</td>
<td>1.1</td>
</tr>
<tr>
<td>Psychiatric disorders</td>
<td></td>
<td></td>
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<tr>
<td>Emotional lability</td>
<td>5.2</td>
<td>7.6</td>
</tr>
<tr>
<td>Depression</td>
<td>3.2</td>
<td>4.9</td>
</tr>
<tr>
<td>Nervousness</td>
<td>2.9</td>
<td>3.3</td>
</tr>
<tr>
<td>Libido decreased</td>
<td>3.2</td>
<td>5.4</td>
</tr>
<tr>
<td>Metabolic and nutritional system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight increase</td>
<td>10.4</td>
<td>7.1</td>
</tr>
<tr>
<td>Application site disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Injection site pain</td>
<td>4.1</td>
<td>7.1</td>
</tr>
<tr>
<td>Injection site reaction</td>
<td>2.9</td>
<td>4.3</td>
</tr>
</tbody>
</table>

\(^1\) Some subjects may experience more than one AE.  
\(^2\) Only AEs for Implanon with an incidence higher than 2.5% are included in the table. The data are derived from studies in the US, Europe, Singapore & Thailand.

Adverse experiences were included as drug-related when they were considered possibly, probably or definitely related to study drug administration either by the investigator or by Schering-Plough.

**During market use of Implanon,** in rare cases, a clinically relevant rise in blood pressure has been observed during the use of Implanon. Urticaria and (aggravation of) angioedema and/or aggravation of hereditary angioedema may occur. Insertion or removal of Implanon may cause some bruising, slight local irritation, pain or itching. Fibrosis at the implant site may occur, a scar may be formed or an abscess may develop. Paresthesia or paresthesia-like events may occur and expulsion or migration of Implanon may be possible (refer PRECAUTIONS). Surgical intervention might be necessary when removing Implanon.\(^†\)

A number of undesirable effects have been reported in women using oral contraceptives, which are discussed in more detail in the PRECAUTIONS section.\(^†\)

**DOSAGE AND ADMINISTRATION**

**How to use Implanon**

*Pregnancy should be excluded before insertion of Implanon.*\(^†\)

Prior to inserting Implanon it is strongly recommended to carefully read the instructions for insertion and removal of the implant in ‘How to Insert Implanon’ and ‘How to remove Implanon.’\(^†\)
Implanon is a long-acting contraceptive. One implant is inserted subdermally. The user should be informed that she can request the removal of Implanon at any time but the implant should not be left in place more than three years. Clinicians may consider earlier replacement of the implant in heavier women (see PRECAUTIONS). Only a physician who is familiar with the removal technique should perform, on request or at the end of the 3 years of use, the removal of Implanon. After the removal of the implant, immediate insertion of another implant will result in continued contraceptive protection.†

To ensure uncomplicated removal it is necessary that Implanon is inserted correctly, directly under the skin. The risk of complication is small if the provided instructions are followed.†

Some cases have been reported in which the implant was not inserted on the correct day or was not properly inserted or was not inserted at all.† Incidentally, this has resulted in unintended pregnancy. The occurrence of such incidents can be minimised when the instructions for insertion (see section ‘How to insert Implanon’ and ‘When to insert Implanon’†) are strictly followed. The presence of the implant should be verified by palpation directly after insertion. If the implant cannot be palpated or when the presence of the implant is in doubt, other methods must be applied to confirm its presence (see section “How to insert Implanon”). Until the presence of Implanon has been verified, a contraceptive barrier method must be used.

It is strongly recommended that physicians, prior to practising the insertion or removal of Implanon, participate in training sessions organised by Schering-Plough.

Physicians who have little experience with subdermal insertion are advised to acquire the correct technique under surveillance of a more experienced colleague. Additional information and more detailed instructions concerning the insertion and removal of Implanon will be sent on request free of charges C/- Schering-Plough Pty Limited - Sydney; telephone Toll Free No.: 1800 818 553.

The Implanon package contains a User Card intended for the user, and an adhesive label intended for the physician’s user record. The User Card records the batch number of the provided implant and helps in remembering the date of insertion, the arm of insertion and the intended day of removal. The adhesive label records the batch number, date of insertion, first day of the last normal menstrual period and verifies that the implant was palpable by both physician and user.
HOW TO INSERT IMPLANON

- Insertion of Implanon should be performed under aseptic conditions, and only by a physician who is familiar with the procedure.

- Insertion of Implanon is performed with the specially designed applicator. The use of this applicator differs substantially from that of a classical syringe. A drawing of a dismantled applicator and its individual components (e.g. cannula, obturator and needle with double angled bevel) is shown in this leaflet to illustrate their specific functions. †

- The procedure used for insertion of Implanon is opposite to giving an injection. When inserting Implanon, the obturator must remain fixed while the cannula (needle) is retracted from the arm. This is in contrast to normal injections where the plunger is pushed and the body of the syringe remains fixed.

- Allow the subject to lie on her back with her non-dominant arm (the arm the woman does not use for writing) turned outwards and bent at the elbow.

- To minimize the risk of neural or vascular damage, Implanon should be inserted at the inner side of the non-dominant upper arm about 8-10 cm above the medial epicondyle of the humerus. †

- Implanon should be inserted subdermally, i.e. just under the skin (subcutaneously). †

- When Implanon is inserted too deeply (intramuscularly or in the fascia) this may cause neural or vascular damage. Too deep insertions have been associated with paraesthesia (due to neural damage) and migration of the implant (due to intramuscular or fascial insertion), and in rare cases with intravascular insertion. Moreover, when the implant is inserted too deeply, it may not be palpable and the localization and/or removal can be difficult later on. †

- Mark the insertion site.

- Clean the insertion site with a disinfectant.

- Anaesthetise with an anaesthetic spray, or with 2 mL of lidocaine (1%) applied just under the skin along the ‘insertion canal’.

- Remove the sterile disposable applicator carrying Implanon from its blister. Keep the needle and the implant sterile. If contamination occurs, a new package with a new sterile applicator must be used.

- While keeping the needle shield on the needle visually verify the presence of the implant, seen as a white body inside the needle tip. If the implant is not seen tap the top of the needle shield against a firm surface to bring the implant in to the needle tip. Following visual confirmation, the implant should be lowered back into the needle by tapping it back in to the needle tip. The needle shield can now be removed. †

- Please note that the implant can fall out of the needle prior to insertion. Therefore, always hold the applicator in the upward position (i.e. with the needle pointed upwards) until the time of insertion. This is to prevent the implant from dropping out. Keep the needle and the implant sterile. If contamination occurs, a new package with a new sterile applicator must be used. †

- Stretch the skin around the insertion site with thumb and index-finger (Figure 1).

- Insert only the tip of the needle, slightly angled (~20°) (Figure 2).

- Release the skin.

- Lower the applicator to a horizontal position (Figure 3).

- While lifting the skin, gently insert the needle to its full length. Do not exert force. The needle should be inserted parallel to the skin to ensure that Implanon is inserted superficially just under the skin without using excessive force, to ensure superficial insertion (Figure 4). †

- Keep the applicator parallel to the surface of the skin.

- When the implant is placed too deeply paraesthesia and migration of the implant may occur. Improper insertions
have been associated with rare cases of intravascular insertion. Moreover, removal can be difficult later on.†

- Break the seal of the applicator by pressing the obturator support (Figure 5).
- Turn the obturator 90° with respect to the cannula (Figure 6).
- Fix the obturator with one hand against the arm and with the other hand slowly retract the cannula (needle) out of the arm (Figure 7).
- Never push against the obturator.
- Check the needle for the absence of the implant. After retraction of the cannula, the grooved tip of the obturator should be visible (Figure 8). Do not confuse the protruding end of the obturator with the implant (same colour).
- Always verify the presence of the implant by palpation and also have the woman palpate it herself. †
- If the implant cannot be palpated or when the presence of the implant is in doubt, other methods must be applied to test its presence. Suitable methods to locate the implant are first of all ultrasound (USS) and secondly magnetic resonance imaging (MRI). Prior to the application of USS or MRI for the localisation of Implanon it is recommended that physicians consult Schering-Plough for guidance. If these imaging methods fail, it is advised to verify the presence of the implant by measuring the etonogestrel level in a blood sample of the subject. In this case Schering-Plough will also provide the appropriate procedure.
- Until the presence of the implant has been confirmed, a contraceptive barrier method must be used. †
- Apply a sterile gauze with a pressure bandage to prevent bruising.
- Fill out the User Card and give it to the patient to facilitate removal of the implant later on. Complete the adhesive label and affix it to the patient file.
- The applicator is for single use only and must be adequately disposed of, in accordance with local regulations for the handling of biohazardous waste.
**WHEN TO INSERT IMPLANON**

No additional contraceptive measures are required if the timing of Implanon insertion is as instructed below.

**NO PRECEDING HORMONAL CONTRACEPTIVE USE**

Implanon should be inserted on day 1-5, but at the latest on day 5 of the woman’s natural cycle (day 1 being the first day of her menstrual bleeding).

**CHANGING FROM A COMBINED ORAL CONTRACEPTIVE (COC), VAGINAL RING OR TRANSDERMAL PATCH †**

Implanon should be inserted preferably on the next day following intake of the last (active) tablet of her COC. At the latest it should be inserted on the day following the usual (active) tablet-free interval or last placebo tablet of her COC. *In case a vaginal ring or transdermal patch has been used, Implanon should be inserted preferably on the day of removal, but at the latest when the next application would be due.* †

**CHANGING FROM A PROGESTAGEN-ONLY METHOD (MINIPILL, INJECTABLE, A DIFFERENT IMPLANT OR FROM A PROGESTAGEN-RELEASING INTRAUTERINE SYSTEM (IUS)) †**

Implanon may be inserted on any day when the woman is switching from a minipill from another implant or an IUS † on the day of its removal, and from an injectable when the next injection would be due.

**FOLLOWING FIRST-TRIMESTER ABORTION**

Implanon should be inserted immediately.

**FOLLOWING CHILDBIRTH OR SECOND-TRIMESTER ABORTION**

*For breastfeeding women see “Use in Lactation”†*

Implanon should be inserted on day 21-28 after delivery or a second trimester abortion. When the implant is inserted later, the woman should be advised to use a barrier method in addition for the first 7 days following insertion. However, if intercourse has already occurred, pregnancy should be excluded or the woman’s first natural period should be awaited before the actual insertion of the implant.
HOW TO REMOVE IMPLANON

- Removal of Implanon should only be performed by a physician who is familiar with the removal technique.

  Fig. A

- The location of the implant is indicated on the User Card.

- Locate the implant by palpation and mark the distal end (fig. A).

- A non-palpable implant should always first be localized by either ultrasound (USS) or magnetic resonance imaging (MRI) before removal is attempted and subsequently be removed under guidance of USS. In case of doubt the presence of Implanon can be verified by etonogestrel determination. Please contact Schering-Plough for further guidance. Exploratory surgery without knowledge of the exact localization of the implant is strictly discouraged. Removal of deeply inserted implants should be conducted with caution in order to prevent damage to deeper neural or vascular structures in the arm and be performed by healthcare providers familiar with the anatomy of the arm.†

- Wash the area and apply a disinfectant.

  Fig. B

- Anaesthetise the arm with 0.5-1 mL lignocaine (1%) at the site of incision, which is just below the distal end of the implant (Fig. B). Note: Apply the anaesthetic under the implant. Application above the implant makes the skin swell, which may cause difficulties in locating the implant.

  Fig. C

- Push down the proximal tip to fix the implant; a bulge may appear indicating the distal end of the implant. Starting below the distal tip of the implant, make a longitudinal incision of 2 mm towards the distal tip of the implant (Figure c).†

  Fig. D

- Gently push the implant towards the incision until the tip is visible. Grasp the implant with forceps (preferably ‘mosquito’ forceps) and remove it (Fig. D).

  Fig. E

  Fig. F
If the tip of the implant is not visible, there might be formation of fibrotic tissue around the implant. The fibrotic tissue can be split by continuing to cut towards the distal tip, until the tip is clearly visible. Remove the implant with a forceps. (Figs. E and F).

If the tip of the implant is not visible, gently insert a forceps into the incision and grasp the implant (Figs. G and H). With second forceps carefully dissect the tissue around the implant. The implant can then be removed (Fig. I).

Be sure to remove the implant entirely. Confirm that the entire rod, which is 4 cm long, has been removed by measuring its length.

Close the incision with a butterfly closure.

Apply sterile gauze with a pressure bandage to prevent bruising.

There have been occasional reports of displacement of the implant (refer PRECAUTIONS): usually this involves minor movement relative to the original position. This may somewhat complicate localisation of the implant by palpation, USS, and/or MRI and may require a somewhat larger incision and more time.

If the woman would like to continue using Implanon, a new implant may be inserted immediately after the old implant is removed (refer HOW TO REPLACE IMPLANON).

If the woman does not wish to continue using Implanon and does not want to become pregnant, another contraceptive method should be recommended.

**HOW TO REPLACE IMPLANON**

- Replacement of Implanon should only be performed under aseptic conditions and only by a physician who is familiar with the removal and insertion procedure.
- Immediate replacement can be done after removal of the previous implant as described in 'HOW TO REMOVE IMPLANON'.
- The procedure to replace Implanon is similar to the insertion procedure described in 'How to insert Implanon'. The new implant can be inserted in the same arm, and frequently through the same incision from which the previous implant was removed. If the same incision is being used, the instructions below must also be taken into account.
- The small incision of the removal procedure can be used as the entrance for the needle of the new applicator.
- Anaesthetize the insertion site with 2 ml lidocaine (1%) applied just under the skin commencing at the removal incision along the 'insertion canal'.
- During replacement inserting the needle to its full length is crucial; failure to do so will result in a partly visible implant in the removal incision in the skin.
- Close the incision with a butterfly closure.
- Apply a sterile gauze with a pressure bandage to prevent bruising. Let the woman keep the bandage in place for at least 48 hours to allow the removal incision to heal.

**OVERDOSAGE**

An implant should always be removed before inserting a new one. There are no data available on overdose with etonogestrel. There have been no reports of serious deleterious effects from an overdose of contraceptives in general.
PRESENTATION

Nature and contents of containers
The pack contains one implant, 4 cm in length and 2 mm in diameter, which is placed in the cannula of a disposable sterile applicator. The sterile applicator containing the implant is packed in a blister pack made of transparent polyester sealed with coated paper. The blister pack is packed in a box together with the package leaflet. AUST R No. 70855
Each implant contains 68 mg of etonogestrel.

Shelf-life and Storage
The shelf-life of Implanon is 5 years when stored below 30°C, protected from moisture.

Name and Address of the Sponsor
Schering-Plough Pty Limited
Level 4, 66 Waterloo Road,
North Ryde NSW 2113
Australia
AUST R No. 70855.
Poison Schedule 4

Applicator Figure

Date of TGA approval: 19 March 2008
Date of most recent amendment: 11 February 2011

† Please note changes in Product Information