

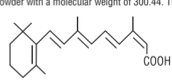


CONTRAINDICATIONS AND WARNINGS
Sotret must not be used by female patients who are or may become pregnant. There is an extremely high risk that severe birth defects will result if pregnancy occurs while taking isotretinoin capsules in any amount, even for short periods of time. Potentially any fetus exposed during pregnancy can be affected. There are no accurate means of determining whether an exposed fetus has been affected.
Birth defects which have been documented following isotretinoin exposure include abnormalities of the face, eyes, ears, skull, central nervous system, cardiovascular system, and thymus and parathyroid glands. Cases of IQ scores less than 85 with or without other abnormalities have been reported. There is an increased risk of spontaneous abortion and premature births have been reported.
Documented external abnormalities include: skull abnormality; ear abnormalities (including anotia, microtia, small or absent external auditory canals); eye abnormalities (including microphthalmia); facial dysmorphism; cleft palate. Documented internal abnormalities include: CNS abnormalities (including cerebral abnormalities, cerebellar abnormalities, hydrocephalus, microcephaly, cranial nerve deficit); cardiovascular abnormalities; thymus gland abnormality; parathyroid hormone deficiency. In some cases death has occurred with certain of the abnormalities previously noted.
If pregnancy does occur during treatment of a female patient who is taking isotretinoin capsules, isotretinoin capsules must be discontinued immediately and she should be referred to an Obstetrician-Gynecologist experienced in reproductive toxicity for further evaluation and counseling.

Special Prescribing Requirements
Because of isotretinoin's teratogenicity and to minimize fetal exposure, Sotret is approved for marketing only under a special restricted distribution program approved by the Food and Drug Administration. This program is called PLEDGE™. Sotret must only be prescribed by prescribers who are registered and activated with the PLEDGE program. Sotret capsules must only be dispensed by a pharmacy registered and activated with PLEDGE, and must only be dispensed to patients who are registered and meet all the requirements of PLEDGE (see PRECAUTIONS).

PRESCRIBER	Female Patients of Childbearing Potential	Male Patients and Female Patients not of Childbearing Potential
Confirms patient counseling	X	X
Enters the two contraception methods chosen by the patient	X	
Enters pregnancy test results	X	
PATIENT		
Answers educational questions before every prescription	X	
Enters two forms of contraception	X	
PHARMACIST		
Contacts system to get an authorization	X	X

DESCRIPTION
Isotretinoin, USP a retinoid, is available as Sotret in 10 mg, 20 mg, 30 mg, and 40 mg soft gelatin capsules for oral administration. Each capsule contains butylated hydroxyanisole, edetate disodium, hydrogenated soybean oil, hydrogenated vegetable oil, iron oxide black, soybean oil and white wax. Gelatin capsules contain glycerin and parabens (methyl and propyl), with the following systems: 10 mg - iron oxide (red) and titanium dioxide; 20 mg - FD&C Red No. 3, FD&C Blue No. 1, and titanium dioxide; 30 mg - FD&C Yellow No. 6, and titanium dioxide; 40 mg - FD&C Yellow No. 6, D&G Yellow No. 10, and titanium dioxide.
Chemically, isotretinoin is 13-*cis*-retinoic acid and is related to both retinoic acid and retinol (vitamin A). It is a yellow to orange crystalline powder with a molecular weight of 300.44. The structural formula is:



Meets USP Dissolution Test 3.
CLINICAL PHARMACOLOGY
Isotretinoin is a retinoid, which when administered in pharmacologic dosages of 0.5 to 1 mg/kg/day (see DOSAGE AND ADMINISTRATION), inhibits sebaceous gland function and keratinization. The exact mechanism of action of isotretinoin is unknown.

Nodular Acne
Clinical improvement in nodular acne patients occurs in association with a reduction in sebum secretion. The decrease in sebum secretion is temporary and is related to the dose and duration of treatment with Sotret, and reflects a reduction in sebaceous gland size and an inhibition of sebaceous gland differentiation.¹

Pharmacokinetics
Absorption
Due to its high lipophilicity, oral absorption of isotretinoin is enhanced when given with a high fat meal. In a crossover study, 74 healthy adult subjects received a single 80 mg oral dose (2 x 40 mg capsules) of isotretinoin capsules under fasted and fed conditions. Both peak plasma concentration (C_{max}) and the total exposure (AUC) of isotretinoin were more than doubled following a standardized high fat meal when compared with isotretinoin capsules given under fasted conditions (see Table 2). The observed elimination half-life was unchanged. This lack of change in half-life suggests that food increases the bioavailability of isotretinoin without altering its disposition. The time to peak concentration (T_{max}) was also increased with food and may be related to a longer absorption phase. Therefore, Sotret capsules should always be taken with food (see DOSAGE AND ADMINISTRATION). Clinical studies have shown that there is no difference in the pharmacokinetics of isotretinoin between patients with nodular acne and healthy subjects with normal skin.

Isotretinoin Capsules 2 x 40 mg Capsules	AUC ₀₋₁₂ (ng•hr/mL)	C _{max} (ng/mL)	T _{max} (hr)	t _{1/2} (hr)
Fed*	10,004 (22%)	862 (22%)	5.3 (77%)	21 (39%)
Fasted	3,703 (46%)	301 (63%)	3.2 (56%)	21 (30%)

*Fasting a standardized high fat meal
Distribution
Isotretinoin is more than 99.9% bound to plasma proteins, primarily albumin.

Metabolism
Following oral administration of isotretinoin, at least three metabolites have been identified in human plasma: 4-oxo-isotretinoin, retinoic acid (tretinoin), and 4-oxo-retinoic acid (4-oxo-tretinoin). Retinoic acid and 13-*cis*-retinoic acid are geometric isomers and show reversible interconversion. The administration of one isomer will give rise to the other. Isotretinoin is also irreversibly oxidized to 4-oxo-isotretinoin, which forms its geometric isomer 4-oxo-tretinoin.
After a single 80 mg oral dose of isotretinoin capsules to 74 healthy adult subjects, concurrent administration of food increased the extent of formation of all metabolites in plasma when compared to the extent of formation under fasted conditions.
All of these metabolites possess retinoid activity that is in some *in vitro* models more than that of the parent isotretinoin. However, the clinical significance of these models is unknown. After multiple oral dose administration of isotretinoin to adult cyclic acne patients (≥ 18 years), the exposure of patients to 4-oxo-isotretinoin at steady-state under fasted and fed conditions was approximately 3.4 times higher than that of isotretinoin.
In vivo studies indicate that the primary P450 isoforms involved in isotretinoin metabolism are C2C, 2D6, 3A4 and 2B6. Isotretinoin and its metabolites are further metabolized into conjugates, which are then excreted in urine and feces.

Elimination
Following oral administration of an 80 mg dose of ¹⁴C-isotretinoin as a liquid suspension, ¹⁴C activity in blood declined with a half-life of 90 hours. The metabolites of isotretinoin and any conjugates are ultimately excreted in the feces and urine in relatively equal amounts (total of 65% to 83%). After a single 80 mg oral dose of isotretinoin to 74 healthy adult subjects under fed conditions, the mean ± SD elimination half-lives (t_{1/2}) of isotretinoin and 4-oxo-isotretinoin were 21 ± 5.2 hours and 24 ± 5.3 hours, respectively. Following single and multiple doses, the observed accumulation ratios of isotretinoin ranged from 0.9 to 5.43 in patients with cyclic acne.

Special Patient Populations
Pediatric Patients
The pharmacokinetics of isotretinoin were evaluated after single and multiple doses in 38 pediatric patients (12 to 15 years) and 19 adult patients (≥ 18 years) who received Sotret for the treatment of severe recalcitrant nodular acne. In both age groups, 4-oxo-isotretinoin was the major metabolite; tretinoin and 4-oxo-tretinoin were also observed. The dose-normalized pharmacokinetics of isotretinoin following single and multiple doses are summarized in Table 3 for pediatric patients. There were no statistically significant differences in the pharmacokinetics of isotretinoin between pediatric and adult patients.

Parameter	Isotretinoin (Single Dose)	Isotretinoin (Steady-State)
C _{max} (ng/mL)	573.25 (278.79)	731.98 (361.86)
AUC ₀₋₁₂ (ng•hr/mL)	3033.37 (1394.17)	5082 (2184.23)
AUC ₀₋₂₄ (ng•hr/mL)	6003.81 (2885.67)	—
T _{max} (hr)†	6 (1 to 24.6)	4 (0 to 12)
C _{5min} (ng/mL)	—	352.32 (184.44)
T _{1/2} (hr)	—	15.69 (5.12)
CL/F (L/hr)	—	17.96 (6.27)

*Single and multiple dose data in this table were obtained following a non-standardized meal that is not comparable to the high-fat meal that was used in the study in Table 2.
†Median (range)
In pediatric patients (12 to 15 years), the mean ± SD elimination half-lives (t_{1/2}) of isotretinoin and 4-oxo-isotretinoin were 15.7 ± 5.1 hours and 23.1 ± 5.7 hours, respectively. The accumulation ratios of isotretinoin ranged from 0.46 to 0.65 for pediatric patients.

INDICATIONS AND USAGE
Severe Recalcitrant Nodular Acne
Sotret is indicated for the treatment of severe recalcitrant nodular acne. Nodules are inflammatory lesions with a diameter of 5 mm or greater. The nodules may be suppurative or hemorrhagic. "Severe," by definition, "means 'many' as opposed to 'few or several' nodules. Because of significant adverse effects associated with its use, Sotret should be reserved for patients with severe nodular acne who are unresponsive to conventional therapy, including systemic antibiotics. In addition, Sotret is indicated only for those female patients who are not pregnant. Sotret can cause severe birth defects (see Boxed CONTRAINDICATIONS AND WARNINGS).

A single course of therapy for 15 to 20 weeks has been shown to result in complete and prolonged remission of disease in many patients.^{1,2,4} If a second course of therapy is needed, it should not be initiated until at least 8 weeks after completion of the first course, because experience has shown that patients may continue to improve while on isotretinoin capsules. The optimal interval before retreatment has not been defined and patients who have not completed skeletal growth (see WARNINGS: Skeletal: Bone Mineral Density, Hypertostosis, and Premature Epiphyseal Closure).

CONTRAINDICATIONS
Pregnancy Category X. See Boxed CONTRAINDICATIONS AND WARNINGS.
Allergic Reactions
Sotret is contraindicated in patients who are hypersensitive to this medication or to any of its components. Sotret should not be given to patients who are sensitive to parabens, which are used as preservatives in the gelatin capsule (see PRECAUTIONS: Hypersensitivity).

WARNINGS
Psychiatric Disorders
Sotret may cause depression, psychosis and, rarely, suicidal ideation, suicide attempts, suicide, and aggressive and/or violent behaviors. No mechanism of action has been established for these events (see ADVERSE REACTIONS: Psychiatric). Prescribers should read the brochure, *Recognizing Psychiatric Disorders in Adolescents and Young Adults: A Guide for Prescribers*, which is available at www.pledgeprogram.com to alert to the warning signs of psychiatric disorders to guide patients to receive the help they need. Therefore, prior to initiation of Sotret therapy, patients and family members should be asked about any history of psychiatric disorder, and at each visit during therapy patients should be assessed for symptoms of depression, mood disturbance, psychosis, or aggression to determine if further evaluation may be necessary. Signs and symptoms of depression, as described in the brochure *Recognizing Psychiatric Disorders in Adolescents and Young Adults*, include sad mood, hopelessness, feelings of guilt, worthlessness or helplessness, loss of pleasure or interest in activities, fatigue, difficulty concentrating, change in sleep pattern, change in weight or appetite, suicidal thoughts or attempts, restlessness, irritability, acting on impulses, and thoughts of suicide. Patients should be alerted to these symptoms. Patients should stop Sotret and the patient or a family member should promptly contact their prescriber if the patient develops depression, mood disturbance, psychosis, or aggression, without waiting until the next visit. Discontinuation of Sotret therapy may be insufficient; further evaluation may be necessary. While such monitoring may be helpful, it may not detect all patients at risk. Patients may report mental health problems or family history of psychiatric disorders. These reports should be discussed with the patient and/or the patient's family. A referral to a mental health professional may be necessary. The physician should consider whether Sotret therapy is appropriate in this setting, for some patients the risks may outweigh the benefits of Sotret therapy.

Pseudotumor Cerebri
Isotretinoin capsule use has been associated with a number of cases of pseudotumor cerebri (benign intracranial hypertension) in some of which involved concomitant use of tetracyclines. Concomitant treatment with tetracyclines should therefore be avoided. Early signs and symptoms of pseudotumor cerebri include papilloedema, headache, nausea and vomiting, and visual disturbances. Patients with these symptoms should be screened for papilloedema and, if present, they should be told to discontinue Sotret immediately and be referred to a neurologist for further diagnosis and care (see ADVERSE REACTIONS: Neurological).

Serious Skin Reactions
There have been postmarketing reports of erythema multiforme and severe skin reactions (e.g., Stevens-Johnson Syndrome (SJS), toxic epidermal necrolysis (TEN)) associated with isotretinoin use. These events may be serious and result in death, life-threatening events, hospitalization or disability. Patients should be monitored closely for severe skin reactions and discontinuation of Sotret should be considered if warranted.
Pancreatitis
Acute pancreatitis has been reported in patients with either elevated or normal serum triglyceride levels. In rare instances, fatal hemorrhagic pancreatitis has been reported. Sotret should be stopped if hypertriglyceridemia cannot be controlled at an acceptable level or if symptoms of pancreatitis occur.

Lipids
Elevations of serum triglycerides in excess of 800 mg/dL have been reported in patients treated with isotretinoin capsules. Marked elevations of serum triglycerides were reported in approximately 25% of patients receiving isotretinoin capsules in clinical trials. In addition, approximately 15% developed a decrease in high-density lipoproteins and about 7% showed an increase in cholesterol levels. In clinical trials, the effects on triglycerides, HDL and cholesterol were reversible upon cessation of isotretinoin capsules therapy. Some patients have been able to reverse triglyceride elevation by reduction in weight, restriction of dietary fat and alcohol, and reduction in dose while continuing isotretinoin capsules.³
Blood lipid determinations should be performed before Sotret is given and then at intervals until the lipid

response to Sotret is established, which usually occurs within 4 weeks. Especially careful consideration must be given when prescribing Sotret therapy (patients with diabetes, hypertension, hypercholesterolemia, increased alcohol intake, lipid metabolism disorder or familial history of lipid metabolism disorder). If Sotret therapy is instituted, more frequent checks of serum values for lipids and/or blood sugar are recommended (see PRECAUTIONS: Laboratory Tests).
The cardiovascular consequences of hypertriglyceridemia associated with Sotret are unknown.

Animal Studies
In rats given 8 or 32 mg/kg/day of isotretinoin (1.3 to 5.3 times the recommended clinical dose of 1 mg/kg/day after normalization for total body surface area) for 18 months or longer, the incidences of focal calcification, fibrosis and inflammation of the myocardium, calcification of coronary, pulmonary and mesenteric arteries, and metastatic calcification of the gastric mucosa were greater than in control rats of similar age. Focal endocardial and myocardial calcifications associated with calcification of the coronary arteries were observed in two dogs after approximately 7 to 7 months of treatment with isotretinoin at a dosage of 60 to 120 mg/kg/day (30 to 60 times the recommended clinical dose of 1 mg/kg/day, respectively, after normalization for total body surface area).

Hearing Impairment
Impaired hearing has been reported in patients taking isotretinoin capsules; in some cases, the hearing impairment has been reported to persist after therapy has been discontinued. Mechanisms of causality for this event have not been established. Patients who experience tinnitus or hearing impairment should discontinue Sotret treatment and be referred for specialized care for further evaluation (see ADVERSE REACTIONS: Special Senses).

Hypotension
Clinical hepatitis considered to be possibly or probably related to isotretinoin capsules therapy has been reported. Additionally, mild to moderate elevations of liver enzymes have been observed in approximately 15% of patients treated during clinical trials, some of which resulted in dosage reduction or continued administration of the drug. If normalization does not readily occur or if hepatitis is suspected during treatment with Sotret, the drug should be discontinued and the etiology further investigated.

Inflammatory Bowel Disease
Isotretinoin capsules have been associated with inflammatory bowel disease (including regional ileitis) in patients without a prior history of intestinal disorders. In some instances, symptoms have been reported to persist after isotretinoin capsules treatment has been stopped. Patients experiencing abdominal pain, rectal bleeding or severe diarrhea should discontinue Sotret immediately (see ADVERSE REACTIONS: Gastrointestinal).

Skeletal
Bone Mineral Density
Effects of multiple courses of Sotret on the developing musculoskeletal system are unknown. There is some evidence that long-term, high dose, or multiple courses of therapy with isotretinoin have more of an effect than short-term, low dose, or single courses of therapy on the musculoskeletal system. In an open-label clinical trial (N = 217) of a single course of therapy with isotretinoin for severe recalcitrant nodular acne, bone density measurements at several skeletal sites were not significantly decreased (lumbar spine change > 4% and total hip change > 5%) or were increased in the majority of patients. One patient had a decrease in lumbar spine bone mineral density > 4% based on unadjusted data. Thym (7.9%) patients had decreases in lumbar spine bone mineral density > 4%, and all the other patients (92%) did not have significant decreases or had increases (adjusted for body mass index). Nine patients (4.5%) had a decrease in total hip bone mineral density > 5% based on unadjusted data. Twenty-one (10.6%) patients had decreases in total hip bone mineral density > 5%, and all the other patients (89%) did not have significant decreases or had increases (adjusted for body mass index). Follow-up studies performed in eight of the patients with decreased bone mineral density for up to 11 months thereafter demonstrated increasing bone density in five patients at the lumbar spine, while the other three patients had lumbar spine bone density measurements below baseline values. Total hip bone mineral densities remained below baseline (range -1.6% to -7.6%) in five of eight patients (62.5%).

In a separate open-label extension study of ten patients, ages 13 to 18 years, who started a second course of isotretinoin 4 months after the first course, two patients showed a decrease in mean lumbar spine bone mineral density up to 3.25% (see PRECAUTIONS: Pediatric Use).

Spontaneous reports of osteoporosis, osteopenia, bone fractures and delayed healing of bone fractures have been reported with isotretinoin capsules. Caution should be exercised when prescribing Sotret to patients who are at increased risk of bone fractures. The risk of bone fractures should be ruled out. Longer term effects have not been studied. It is important that Sotret be given in the recommended doses for no longer than the recommended duration.

Hypertostosis
A high prevalence of skeletal hypertostosis was noted in clinical trials for disorders of keratinization with a mean dose of 2.24 mg/kg/day. Additionally, skeletal hypertostosis was noted in six of eight patients in a prospective study of disorders of keratinization.⁴ Minimal skeletal hypertostosis and calcification of ligaments and tendons have also been observed by x-ray in prospective studies of nodular acne patients treated with a single course of therapy at recommended doses. The skeletal effects of multiple Sotret treatment courses for acne are unknown.

In a clinical study of 217 pediatric patients (12 to 17 years) with severe recalcitrant nodular acne, hypertostosis was not observed after 16 to 20 weeks of treatment with approximately 1 mg/kg/day of isotretinoin capsules over the 16-week study period. Hypertostosis may require a longer time frame to appear. The clinical course and significance remain unknown.

Premature Epiphyseal Closure
There are spontaneous reports of premature epiphyseal closure in acne patients receiving recommended doses of isotretinoin capsules. The effect of multiple courses of Sotret on epiphyseal closure is unknown.

Visual Impairment
Visual problems should be carefully monitored. All Sotret patients experiencing visual difficulties should discontinue Sotret treatment and have an ophthalmological examination (see ADVERSE REACTIONS: Special Senses).

Cornel Opacities
Cornel opacities have occurred in patients receiving isotretinoin capsules for acne and more frequently when higher drug dosages were used in patients with disorders of keratinization. The corneal opacities that have been reported in clinical trials were reversible. Patients with corneal opacities have either completely resolved or were resolving at follow-up 6 to 7 weeks after discontinuation of the drug (see ADVERSE REACTIONS: Special Senses).

Decreased Night Vision
Decreased night vision has been reported during isotretinoin capsules therapy and in some instances the event has persisted after therapy was discontinued. Because the onset in some patients was sudden, patients should be advised of this potential problem and warned to be cautious when driving or operating any vehicle at night.

PRECAUTIONS
Sotret must only be prescribed by prescribers who are registered and activated with the PLEDGE program. Sotret must only be dispensed by a pharmacy registered and activated with PLEDGE, and must only be dispensed to patients who are registered and meet all the requirements of PLEDGE. Registered and activated pharmacies must receive Sotret only from wholesalers registered with PLEDGE. PLEDGE program requirements for wholesalers, prescribers and pharmacists are described below.

Wholesalers
For the purpose of the PLEDGE program, the retail wholesaler refers to wholesaler, distributor and/or chain pharmacy distributor. To distribute Sotret, wholesalers must be registered with PLEDGE and agree to meet all PLEDGE requirements for wholesale distribution of isotretinoin products. Wholesalers must register with PLEDGE by signing and returning the PLEDGE wholesaler agreement that affirms they will comply with all PLEDGE requirements for distribution of isotretinoin. These include:

- Registering prior to distributing isotretinoin and re-registering annually thereafter
- Distributing only FDA approved isotretinoin product
- Only shipping isotretinoin product to:
 - wholesalers registered in the PLEDGE program with prior written consent from the manufacturer or
 - pharmacies licensed in the US and registered and activated in the PLEDGE program
- Notifying the isotretinoin manufacturer (or delegate) of any non-registered and/or non-activated pharmacy or unregistered wholesaler that attempts to order isotretinoin
- Complying with inspection of wholesaler records for verification of compliance with the PLEDGE program by the isotretinoin manufacturer (or delegate)
- Returning to the manufacturer (or delegate) any undistributed product if registration is revoked by the manufacturer or if the wholesaler chooses to not re-register annually

Prescribers
To prescribe isotretinoin, the prescriber must be registered and activated with the pregnancy risk management program PLEDGE. Prescribers can register by signing and returning the completed registration form. Prescribers can only activate their registration by affirming that they meet requirements and will comply with all PLEDGE requirements by attending to the following points:

- Know the risks and severity of fetal/neonatal defects from isotretinoin.
- Know the risk factors for unplanned pregnancy and the effective measures for avoidance of unplanned pregnancy.
- Have the expertise to provide the patient with detailed pregnancy prevention counseling or i will refer her to a specialist for such counseling by the manufacturer.
- Will comply with the PLEDGE program requirements described in the booklets entitled *The Guide to Best Practices for the PLEDGE Program* and *The PLEDGE Program Prescriber Contraception Counseling Guide*.
- Before beginning treatment of female patients of childbearing potential with isotretinoin and on a monthly basis, the patient will be counseled to avoid pregnancy by using two forms of contraception simultaneously and continuously one month before, during and one month after isotretinoin therapy, unless the patient commits to continuous abstinence.
- Will not prescribe isotretinoin to any female patient of childbearing potential until verifying she has a negative screening pregnancy test and monthly negative CLIA-certified (Clinical Laboratory Improvement Amendment) pregnancy tests. Patients should have a pregnancy test at the completion of the entire course of isotretinoin capsules and another pregnancy test one month later.
- Will report any pregnancy case that becomes aware of while the female patient is on isotretinoin or one month after the completion of the course of isotretinoin therapy.

To prescribe isotretinoin, the Prescriber must access the PLEDGE system via the internet (www.pledgeprogram.com) or telephone (1-866-495-0654) to:

- 1) Register each patient in the PLEDGE program.
- 2) Confirm a month that the patient is receiving counseling and education.
- 3) For female patients of childbearing potential:
 - Enter patient's two chosen forms of contraception each month.
 - Enter monthly result from CLIA-certified laboratory conducted pregnancy test.

Isotretinoin must only be dispensed to female patients who are known not to be pregnant as confirmed by a negative screening pregnancy test and monthly negative CLIA-certified pregnancy tests.

Isotretinoin must only be dispensed by a pharmacy registered and activated with the pregnancy risk management program PLEDGE and only when the registered patient meets all the requirements of the PLEDGE program. Meeting the requirements for a female patient of childbearing potential signifies that she has been counseled and activated as a Patient Information/Informed Consent (for female patients who can get pregnant) form that contains warnings about the risk of potential birth defects if the fetus is exposed to isotretinoin. The patient must sign the informed consent form before starting treatment and the prescriber must also document that the patient has used two forms of contraception for one month before receiving the initial isotretinoin prescription. The first test (a screening test) is obtained by the prescriber when the decision is made to pursue qualification of the patient for isotretinoin. The second pregnancy test (a confirmation test) must be done in a CLIA-certified laboratory. The interval between the two tests should be at least 19 days.

For patients with regular menstrual cycles, the second pregnancy test should be done during the first 5 days of the menstrual period immediately preceding the beginning of isotretinoin therapy and after the patient has used two forms of contraception for one month.

For patients with amenorrhea, irregular cycles or using a contraceptive method that precludes withdrawal bleeding, the second pregnancy test must be done immediately preceding the beginning of isotretinoin therapy and after the patient has used two forms of contraception for one month.

Has had a negative urine or serum pregnancy tests with a sensitivity of at least 25 mIU/mL, before receiving the initial isotretinoin prescription. The first test (a screening test) is obtained by the prescriber when the decision is made to pursue qualification of the patient for isotretinoin. The second pregnancy test (a confirmation test) must be done in a CLIA-certified laboratory before receiving each subsequent course of isotretinoin. A pregnancy test must be repeated every month, in a CLIA-certified laboratory, prior to the female patient receiving each prescription.

Egg selection and commitment to use two forms of effective contraception simultaneously, at least one of which must be a primary form, unless the patient commits to continuous abstinence from heterosexual contact, or the patient has undergone a hysterectomy or bilateral oophorectomy, or has been clinically confirmed to be post-menopausal. Patients must use two forms of effective contraception for at least one month prior to initiation of isotretinoin therapy, during isotretinoin therapy and for one month after discontinuing isotretinoin therapy. Counseling about contraception and behaviors associated with an increased risk of pregnancy must be repeated on a monthly basis.

If the patient has unprotected heterosexual intercourse at any time one month before, during or one month after therapy, she should:

1. Stop taking Sotret immediately, if on therapy
2. Have a pregnancy test at least 19 days after the last act of unprotected heterosexual intercourse
3. Start using two forms of effective contraception simultaneously again for one month before resuming Sotret therapy
4. Have a second pregnancy test after using two forms of effective contraception for one month as described above depending on whether she has regular menses or not.

Effective forms of contraception include both primary and secondary forms of contraception:

Primary forms	Secondary forms
• tubal sterilization	• barrier
• partner's vasectomy	• male latex condom with or without spermicide
• intrauterine device	• diaphragm with spermicide
• hormonal (combination oral, injectables, transdermal patch, injectables, implantables or vaginal ring)	• cervical cap with spermicide
	• Ovulation barrier
	• vaginal sponge (contains spermicide)

Any birth control method can fail. There have been reports of pregnancy from female patients who have used oral contraceptives, as well as transdermal patch/injectable/implantable/vaginal ring hormonal birth control products, these pregnancies occurred while these patients were taking isotretinoin capsules. These reports are more frequent for female patients who use only a single method of contraception. Therefore, it is critically important that female patients of childbearing potential use two effective forms of contraception simultaneously. Patients must receive written warnings about the rates of possible contraception failure (included in patient education kits).

Using two forms of contraception simultaneously substantially reduces the chances that a female will become pregnant over the risk of pregnancy with either form alone. A drug interaction that decreases effectiveness of hormonal contraceptives has not been entirely ruled out for Sotret (see PRECAUTIONS: Drug Interactions). Although hormonal contraceptives are highly effective, prescribers are advised to consult the package insert of any medication administered concomitantly with hormonal contraceptives, since some medications may decrease the effectiveness of these birth control products.

Patients should be prospectively cautioned not to self-medicate with the herbal supplement St. John's Wort because a possible interaction has been suggested with hormonal contraceptives based on reports of breakthrough bleeding on oral contraceptives shortly after starting St. John's Wort. Pregnancies have been reported by users of combined hormonal contraceptives who also used some form of St. John's Wort. If a pregnancy does occur during isotretinoin treatment, isotretinoin must be discontinued immediately. The patient should be referred to an Obstetrician-Gynecologist experienced in reproductive toxicity for further evaluation and counseling. Any suspected fetal exposure during or one month after isotretinoin therapy must be reported immediately to the FDA via the MedWatch number 1-800-FDA-1088 and also to the PLEDGE pregnancy registry at 1-866-495-0654 or via the internet (www.pledgeprogram.com).

All Patients
Isotretinoin is contraindicated in female patients who are pregnant. To receive isotretinoin all patients must meet all the following conditions:

- Must be registered with the PLEDGE program by the prescriber
- Must understand that severe birth defects can occur with the use of isotretinoin by female patients
- Must be reliable in understanding and carrying out instructions
- Must sign a Patient Information/Informed Consent (for all patients) form that contains warnings about the risks of childbearing potential
- Must fill and pick up the prescription within 7 days of the date of specimen collection for the pregnancy test for female patients of childbearing potential
- Must fill and pick up the prescription within 30 days of the office visit for male patients and female patients of childbearing potential
- Must not donate blood while on isotretinoin and for one month after treatment has ended
- Must not share isotretinoin with anyone, even someone who has similar symptoms

Female Patients of Childbearing Potential
Isotretinoin is contraindicated in female patients who are pregnant. In addition to the requirements for all patients described above, female patients of childbearing potential must meet the following conditions:

- Must NOT be pregnant or breast-feeding
- Must comply with the required pregnancy tests when taking a CLIA-certified laboratory
- Must fill and pick up the prescription within 7 days of the date of specimen collection for the pregnancy test
- Must be capable of complying with the mandatory contraceptive measures required for isotretinoin therapy, or commit to continuous abstinence from heterosexual intercourse and understand behaviors associated with an increased risk of pregnancy

• Must understand that it is her responsibility to avoid pregnancy one month before, during and one month after therapy (see PRECAUTIONS: Pregnancy Testing).

• Must have signed an additional Patient Information/Informed Consent About Birth Defects (for female patients who can get pregnant) form, before starting isotretinoin, that contains warnings about the risk of potential birth defects if the fetus is exposed to isotretinoin.

• Must access the PLEDGE system via the internet (www.pledgeprogram.com) or telephone (1-866-495-0654), before starting isotretinoin, or a monthly basis during therapy and one month after the last dose to answer questions on the program requirements and to enter the patient's two chosen forms of contraception

• Must have been informed of the purpose and importance of providing information to the PLEDGE program and become pregnant while taking isotretinoin or within one month of the last dose

Pharmacists:
To dispense isotretinoin, pharmacies must be registered and activated with the pregnancy risk management program PLEDGE.

The Responsible Site Pharmacist must register the pharmacy by signing and returning the completed registration form. After registration, the Responsible Site Pharmacist can only activate the pharmacy registration by affirming that they meet requirements and will comply with all PLEDGE requirements by attending to the following points:

- Know the risks and severity of fetal/neonatal defects from isotretinoin.
- Will train all pharmacists who participate in the filling and dispensing of isotretinoin prescriptions, on the PLEDGE program requirements.
- Will comply and seek to ensure all pharmacists who participate in the filling and dispensing of isotretinoin prescriptions comply with the PLEDGE program requirements described in the booklet entitled *The PLEDGE Program*.
- Will obtain Sotret product only from PLEDGE registered wholesalers.
- Will not sell, buy, borrow, loan or otherwise transfer isotretinoin in any manner to or from another pharmacy.
- Will return to the manufacturer (or delegate) any unused product if registration is revoked by the manufacturer or if the pharmacy chooses to not re-activate annually.
- Will not fill isotretinoin for any party other than a qualified patient.

To dispense isotretinoin, the pharmacist must:

- 1) be trained by the Responsible Site Pharmacist concerning the PLEDGE program requirements.
- 2) obtain authorization from the PLEDGE program via the internet (www.pledgeprogram.com) or telephone (1-866-495-0654) for every isotretinoin prescription. Authorization signifies that the patient has met all program requirements and is qualified to receive isotretinoin.

3) write the Risk Management Authorization (RMA) number on the prescription.
Sotret must only be dispensed:
• in more than a 30 day supply
• with a Sotret Medication Guide
• after authorization from the PLEDGE program
• prior to the "do not dispense to patient after" date provided by the PLEDGE system (within 30 days of the office visit for male patients and female patients not of childbearing potential and within 7 days of the date of specimen collection for female patients of childbearing potential)

• with a new prescription for refills and another authorization from the PLEDGE program (No automatic refills are allowed).

A Sotret Medication Guide must be given to the patient each time Sotret is dispensed, as required by law. This Sotret Medication Guide is an important part of the risk management program for the patient. Sotret must not be prescribed, dispensed or otherwise obtained through the internet or any other means outside of the PLEDGE program. Only FDA-approved Sotret products must be distributed, prescribed, dispensed and used. Patients must fill Sotret prescriptions only at U.S. licensed pharmacies.

A description of the PLEDGE program educational materials available with PLEDGE is provided below. The main goal of these educational materials is to explain the PLEDGE program requirements and to reinforce the educational messages.

- 1) *The Guide to Best Practices for the PLEDGE Program* includes: isotretinoin teratogenic potential, information on pregnancy testing, and the method to complete a qualified isotretinoin prescription.
- 2) *The PLEDGE Program Prescriber Contraception Counseling Guide* includes: specific information about effective contraception, the limitations of contraceptive methods, behaviors associated with an increased risk of contraceptive failure and pregnancy and the methods to evaluate pregnancy risk.
- 3) *The Pharmacist Guide for the PLEDGE Program* includes: isotretinoin teratogenic potential and the method to obtain authorization to dispense an isotretinoin prescription.
- 4) *The PLEDGE program* is a systematic approach to comprehensive patient education about their responsibilities and includes education for contraception compliance and reinforcement of educational messages. The PLEDGE program includes information on the risks and benefits of childbearing which is appropriate for male patients and female patients who are registered with PLEDGE and

of alcohol, at least 36 hours should elapse before these determinations are made. It is recommended that these tests be performed at weekly or biweekly intervals until the lipid response to Sotret is established. The incidence of hypertriglyceridemia is one patient in four on isotretinoin therapy (see **WARNINGS: Lipids**).

Liver Function Tests: Since elevations of liver enzymes have been observed during clinical trials, and hepatitis has been reported, pretreatment and follow-up liver function tests should be performed at weekly or biweekly intervals until the response to Sotret has been established (see **WARNINGS: Hepatotoxicity**).

Glucose: Some patients receiving isotretinoin capsules have experienced problems in the control of their blood sugar. In addition, new cases of diabetes have been diagnosed during isotretinoin capsules therapy, although no causal relationship has been established.

CPK: Some patients undergoing vigorous physical activity while on isotretinoin capsules therapy have experienced elevated CPK levels; however, the clinical significance is unknown. There have been rare postmarketing reports of rhabdomyolysis, some associated with strenuous physical activity. In a clinical trial of 212 pediatric patients (12 to 17 years) with severe recalcitrant nodular acne, transient elevations in CPK were observed in 12% of patients, including those undergoing strenuous physical activity in association with reported musculoskeletal adverse events such as back pain, arthralgia, limb injury, or muscle strain. In these patients, approximately half of the CPK elevations returned to normal within 2 weeks and half returned to normal within 4 weeks. No cases of rhabdomyolysis were reported in this trial.

Carcinogenesis, Mutagenesis and Impairment of Fertility

In male and female Fischer 344 rats given oral isotretinoin at dosages of 8 or 32 mg/kg/day (1.3 to 5.3 times the recommended clinical dose of 1 mg/kg/day, respectively after normalization for total body surface area) for greater than 18 months, there was a dose related increased incidence of pheochromocytoma relative to controls. The incidence of adrenal medullary hyperplasia was also increased at the higher dosage in both sexes. The relatively high level of spontaneous pheochromocytomas occurring in the male Fischer 344 rat makes it an equivocal model for study of this tumor; therefore, the relevance of this tumor to the human population is uncertain.

The Ames test was conducted with isotretinoin in two laboratories. The results of the tests in one laboratory were negative while in the second laboratory a weakly positive response (less than 1.6 x background) was noted in *S. typhimurium* TA100 when the assay was conducted with metabolic activation. No dose-response effect was seen and all other strains were negative. Additionally, other tests designed to assess genotoxicity (Chinese hamster cell assay, mouse micronucleus assay, *S. cerevisiae* D5 assay, in vitro clastogenesis assay with human-derived lymphocytes and unscheduled DNA synthesis assay) were all negative.

In rats, no adverse effects on gonadal function, fertility, conception rate, gestation or parturition were observed at oral dosages of isotretinoin of 2, 8 or 32 mg/kg/day (0.3, 1.3 or 5.3 times the recommended clinical dose of 1 mg/kg/day, respectively, after normalization for total body surface area).

In dogs, testicular atrophy was noted after treatment with oral isotretinoin for approximately 30 weeks at dosages of 20 or 60 mg/kg/day (10 or 30 times the recommended clinical dose of 1 mg/kg/day, respectively, after normalization for total body surface area). In general, there was microscopic evidence for appreciable depression of spermatogenesis but some minor spermatids were observed in all testes examined and in no instance were completely atrophic tubules seen. In studies of 66 men, 30 of whom were patients with nodular acne under treatment with oral isotretinoin, no significant changes were noted in the count or motility of spermatozoa in the ejaculate. In a study of 50 men (ages 17 to 35 years) receiving isotretinoin therapy for nodular acne, no significant effects were seen on ejaculate volume, sperm count, total sperm motility, morphology or seminal plasma fructose.

Pregnancy: Category X. See Boxed CONTRAINDICATIONS AND WARNINGS.

Nursing Mothers

It is not known whether this drug is excreted in human milk. Because of the potential for adverse effects, nursing mothers should not receive Sotret.

Pediatric Use

The use of isotretinoin capsules in pediatric patients less than 12 years of age has not been studied. The use of isotretinoin capsules for the treatment of severe recalcitrant nodular acne in pediatric patients ages 12 to 17 years should be given careful consideration, especially for those patients where a known metabolic or structural bone disease exists (see **PRECAUTIONS: General**). Use of Sotret in this age group for severe recalcitrant nodular acne is supported by evidence in a clinical study comparing 103 pediatric patients (13 to 17 years) to 197 adult patients (>18 years). Results from this study demonstrated that Sotret, at a dose of 1 mg/kg/day given in two divided doses, was equally effective in treating severe recalcitrant nodular acne in both pediatric and adult patients.

In studies with isotretinoin capsules, adverse reactions reported in pediatric patients were similar to those described in adults except for the increased incidence of back pain and arthralgia (both of which were sometimes severe) and myalgia in pediatric patients (see **ADVERSE REACTIONS**).

In an open-label clinical trial (N = 217) of a single therapy strategy with isotretinoin capsules for severe recalcitrant nodular acne, bone density measurements at several skeletal sites were not significantly decreased (lumbar spine change > 4% and total hip change > 5%) or were increased in the majority of patients. One patient had a decrease in lumbar spine bone mineral density > 4% based on unadjusted data. Sixteen (7.9%) patients had decreases in lumbar spine bone mineral density in all tests examined and in no instance were there significant decreases or had increases (adjusted for body mass index). Nine patients (4.5%) had a decrease in total hip bone mineral density > 5% based on unadjusted data. Twenty one (10.6%) patients had decreases in total hip bone mineral density > 5% in all tests examined. Ninety percent of patients had increases or had increases (adjusted for body mass index). Follow-up studies performed in eight of the patients with decreased bone mineral density for up to 11 months thereafter demonstrated increasing bone density in five patients at the lumbar spine, while the other three patients had lumbar spine bone density measurements below baseline values. Total hip bone mineral densities remained below baseline (range -1.6% to -7.6%) in five of eight patients (62.5%).

In a separate open-label extension study of ten patients, ages 13 to 18 years, who started a second course of isotretinoin capsules four months after the first course, two patients showed a decrease in mean lumbar spine bone mineral density up to 3.25% (see **WARNINGS: Skeletal: Bone Mineral Density**).

Geriatric Use

Clinical studies of isotretinoin did not include sufficient numbers of subjects aged 65 years and over to determine whether the younger subjects. Although no reported clinical experience has not identified differences in responses between elderly and younger patients, effects of aging might be expected to increase some risks associated with isotretinoin therapy (see **WARNINGS and PRECAUTIONS**).

ADVERSE REACTIONS

Clinical Trials and Postmarketing Surveillance

The adverse reactions listed below reflect the experience from investigational studies of isotretinoin capsules, and the postmarketing experience. The relationship of some of these events to isotretinoin capsules therapy is unknown. Many of the side effects and adverse reactions seen in patients receiving isotretinoin capsules are similar to those described in patients taking very high doses of vitamin A (dryness of the skin and mucous membranes, e.g., of the lips, nasal passage and eyes).

Dose Relationship

Chellitis and hypertriglyceridemia are usually dose related. Most adverse reactions reported in clinical trials were reversible when therapy was discontinued; however, some persisted after cessation of therapy (see **WARNINGS and ADVERSE REACTIONS**).

Body as a Whole

allergic reactions, including vasculitis, systemic hypersensitivity (see **PRECAUTIONS: Hypersensitivity**), edema, fatigue, lymphadenopathy, weight loss

Cardiovascular

palpitation, tachycardia, vascular thrombotic disease, stroke

Endocrine/Metabolic

hypertriglyceridemia (see **WARNINGS: Lipids**), alterations in blood sugar levels (see **PRECAUTIONS: Laboratory Tests**)

Gastrointestinal

inflammatory bowel disease (see **WARNINGS: Inflammatory Bowel Disease**), hepatitis (see **WARNINGS: Hepatotoxicity**), pancreatitis (see **WARNINGS: Lipids**), bleeding and inflammation of the gums, colitis, esophagitis/esophageal ulceration, ileitis, nausea, other nonspecific gastrointestinal symptoms

Hematologic

allergic reactions (see **PRECAUTIONS: Hypersensitivity**), anemia, thrombocytopenia, neutropenia, rare reports of agranulocytosis (see **PRECAUTIONS: Information for Patients**), see **PRECAUTIONS: Laboratory Tests** for other hematological parameters

Musculoskeletal

skeletal hyperostosis, calcification of tendons and ligaments, premature epiphyseal closure, decreases in bone mineral density (see **WARNINGS: Skeletal**), musculoskeletal symptoms (sometimes severe) including back pain, myalgia, and arthralgia (see **PRECAUTIONS: Information for Patients**), transient pain in the chest (see **PRECAUTIONS: Information for Patients**), arthritis, tendinitis, or other types of bone abnormalities, elevations of CPK/rare reports of rhabdomyolysis (see **PRECAUTIONS: Laboratory Tests**)

Neurological

pseudotumor cerebri (see **WARNINGS: Pseudotumor Cerebri**), dizziness, drowsiness, headache, insomnia, lethargy, malaise, nervousness, paresthesias, seizures, stroke, syncope, weakness

Psychiatric

suicidal ideation, suicide attempts, suicide, depression, psychosis, aggression, violent behaviors (see **WARNINGS: Psychiatric Disorders**), emotional instability

Of the patients reporting depression, some reported that the depression subsided with discontinuation of therapy and recurred with reinstatement of therapy.

Reproductive System

abnormal menses

Respiratory

bronchospasms (with or without a history of asthma), respiratory infection, voice alteration

Skin and Appendages

acne fulminans, alopecia (which in some cases persists), bruising, chelitis (dry lips), dry mouth, dry nose, dry skin, epistaxis, erythema xanthomas, erythema multiforme, flushing, fragility of skin, hair abnormalities, hirsutism, hyperpigmentation and hypopigmentation, infections (including disseminated herpes simplex), nail dystrophy, paronychia, peeling of palms and soles, photoallergic/photoenzyming reactions, pruritus, pyogenic granuloma, rash (including facial erythema, seborrhea, and eczema), Stevens-Johnson syndrome, sunburn susceptibility increased, sweating, toxic epidermal necrolysis, urticaria, vasculitis (including Wegener's granulomatosis; see **PRECAUTIONS: Hypersensitivity**) abnormal wound healing (delayed healing of excimer granulation tissue with crusting; see **PRECAUTIONS: Information for Patients**)

Special Senses

Hearing

hearing impairment (see **WARNINGS: Hearing Impairment**), tinnitus

Vision

corneal opacities (see **WARNINGS: Corneal Opacities**), decreased night vision which may persist (see **WARNINGS: Decreased Night Vision**), cataracts, color vision disorder, conjunctivitis, dry eyes, eyelid inflammation, keratitis, optic neuritis, photophobia, visual disturbances

Urinary System

glomerulonephritis (see **PRECAUTIONS: Hypersensitivity**), nonspecific urogenital findings (see **PRECAUTIONS: Laboratory Tests** for other urological parameters)

Laboratory

Elevation of plasma triglycerides (see **WARNINGS: Lipids**), decrease in serum high-density lipoprotein (HDL) levels, elevations of serum cholesterol during treatment

Increased alkaline phosphatase, SGOT (AST), SGPT (ALT), GGT or LDH (see **WARNINGS: Hepatotoxicity**)

Elevation of fasting blood sugar, elevations of CPK (see **PRECAUTIONS: Laboratory Tests**), hyperuricemia

Decreases in red blood cell parameters, decreases in white blood cell counts (including severe neutropenia and rare reports of agranulocytosis; see **PRECAUTIONS: Information for Patients**), elevated sedimentation rates, elevated platelet counts, thrombocytopenia

White cells in the urine, proteinuria, microscopic or gross hematuria

OVERDOSAGE

The oral LD₅₀ of isotretinoin is greater than 4000 mg/kg in rats and mice >800 times the recommended clinical dose of 1 mg/kg/day after normalization of the rat dose for total body surface area and 300 times the recommended clinical dose of 1 mg/kg/day after normalization of the mouse dose for total body surface area and is approximately 1800 mg/kg in rabbits (655 times the recommended clinical dose of 1 mg/kg/day after normalization for total body surface area). In humans, overdosage has been associated with vomiting, facial flushing, cheilitis, abdominal pain, headache, dizziness and ataxia. These symptoms quickly resolve without apparent residual effects.

Isotretinoin causes serious birth defects at any dosage (see **Boxed CONTRAINDICATIONS AND WARNINGS**). Female patients of childbearing potential who present with isotretinoin overdose must be evaluated for pregnancy. Patients who are pregnant should receive counseling about the risks to the fetus, as described in the **Boxed CONTRAINDICATIONS AND WARNINGS**. Non-pregnant patients must be warned to avoid pregnancy for at least one month and receive contraceptive counseling as described in **PRECAUTIONS**. Educational materials for such patients can be obtained by calling the manufacturer. Because an overdose would be expected to result in higher levels of isotretinoin in the serum than found during a normal treatment course, male patients should use a condom, or avoid reproductive sexual activity with a female patient who is or might become pregnant, for one month after the overdose. All patients with isotretinoin overdose should not donate blood for at least one month.

DOSE AND ADMINISTRATION

Sotret should be administered with a meal (see **PRECAUTIONS: Information for Patients**). The recommended dosage range for Sotret is 0.5 to 1 mg/kg/day given in two divided doses with food for 15 to 20 weeks. In studies comparing 0.1, 0.5 and 1 mg/kg/day it was found that all dosages provided initial clearing of disease, but there was a greater need for retreatment with the lower dosages. During treatment, the dose may be adjusted according to response of the disease and/or the appearance of clinical side effects — some of which may be dose related. Adult patients whose disease is very severe with scarring or is primarily manifested on the trunk may require dose adjustments up to 2 mg/kg/day, as tolerated. Failure to take Sotret with food will significantly decrease absorption. Before upward dose adjustments are made, the patients should be questioned about their compliance with food instructions.

The safety of one daily dosing with Sotret has not been established. Once daily dosing is **not recommended**. If the total nodule count has been reduced by more than 70% prior to completing 15 to 20 weeks of treatment, the drug may be discontinued. After a period of 2 months or more off therapy, and if warranted by persistent or recurring severe nodular acne, a second course of therapy may be initiated. The optimal interval before retreatment has not been defined for patients who have not completed skeletal growth. Long-term use of Sotret, even in low doses, has not been studied, and is not recommended. It is important that Sotret be given at the recommended doses for no longer than the recommended duration. The effect of long-term use of Sotret on bone loss is unknown (see **WARNINGS: Skeletal: Bone Mineral Density, Hyperostosis, and Premature Epiphyseal Closure**).

Contraceptive measures must be followed for any subsequent course of therapy (see **PRECAUTIONS**).

Table 4. Sotret Dosing by Body Weight (Based on Administration with Food)

Body Weight		Total mg/day		
kilograms	pounds	0.5 mg/kg	1 mg/kg	2 mg/kg
40	88	20	40	80
50	110	25	50	100
60	132	30	60	120
70	154	35	70	140
80	176	40	80	160
90	198	45	90	180
100	220	50	100	200

*See **DOSE AND ADMINISTRATION**; the recommended dosage range is 0.5 to 1 mg/kg/day.

INFORMATION FOR PHARMACISTS

Access the iPLEDGE system via the internet (www.ipledeprogram.com) or telephone (1-866-495-0654) to obtain an authorization and the **do not dispense to patient alert** date. Sotret must only be dispensed in no more than a 30 day supply.

REFILLS REQUIRE A NEW PRESCRIPTION AND A NEW AUTHORIZATION FROM THE IPLEDGE SYSTEM.

A Sotret Medication Guide must be given to the patient each time Sotret is dispensed, as required by law. This Sotret Medication Guide is an important part of the risk management program for the patient.

HOW SUPPLIED

Sotret (isotretinoin capsules, USP) are available as follows:

- Soft gelatin capsules, 10 mg (light pink), imprinted "30"
- Boxes of 30 containing 3 Prescription Packs of 10 capsules (NDC 10631-584-31)
- Boxes of 100 containing 10 Prescription Packs of 10 capsules (NDC 10631-584-77)
- Soft gelatin capsules, 30 mg (maroon), imprinted "60"
- Boxes of 30 containing 3 Prescription Packs of 10 capsules (NDC 10631-585-31)
- Boxes of 100 containing 10 Prescription Packs of 10 capsules (NDC 10631-585-77)
- Soft gelatin capsules, 30 mg (golden yellow), imprinted "30"
- Boxes of 30 containing 3 Prescription Packs of 10 capsules (NDC 10631-447-31)
- Boxes of 100 containing 10 Prescription Packs of 10 capsules (NDC 10631-447-77)
- Soft gelatin capsules, 40 mg (yellow), imprinted "70"
- Boxes of 30 containing 3 Prescription Packs of 10 capsules (NDC 10631-586-31)
- Boxes of 100 containing 10 Prescription Packs of 10 capsules (NDC 10631-586-77)

Storage

Store at 20° - 25° C (68° - 77° F) (see USP Controlled Room Temperature). Protect from light.

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PATIENT INFORMATION/INFORMED CONSENT ABOUT BIRTH DEFECTS (for female patients who can get pregnant)

To be completed by the patient (and parent or guardian if patient is under age 18) and signed by her doctor.

Read each item below and initial in the space provided to show that you understand each item and agree to follow your doctor's instructions. **Do not sign this consent and do not take isotretinoin if there is anything that you do not understand.**

*A parent or guardian of a minor patient (under age 18) must also read and initial each item before signing the consent.

(Patient's Name)	
1. I understand that there is a very high chance that my unborn baby could have severe birth defects if I am pregnant or become pregnant while taking isotretinoin. This can happen with any amount and even if taken for short periods of time. This is why I must not be pregnant while taking isotretinoin.	Initials: _____
2. I understand that I must not get pregnant one month before, during the entire time of my treatment and for one month after the end of my treatment with isotretinoin.	Initials: _____
3. I understand that I must avoid sexual intercourse completely, or I must use two separate, effective forms of birth control (contraception) at the same time. The only exceptions are: if I have had surgery to remove the uterus (a hysterectomy) or both of my ovaries (bilateral oophorectomy), or if my doctor has medically determined that I am post-menopausal.	Initials: _____
4. I understand that hormonal birth control products are among the most effective forms of birth control. Combination birth control pills and other hormonal products include skin patches, shots, under-the-skin implants, vaginal rings and intrauterine devices (IUDs). Any form of birth control can fail. That is why I must use two different birth control methods at the same time starting one month before, during, and for one month after stopping therapy every time I have sexual intercourse, even if one of the methods I choose is hormonal birth control.	Initials: _____
5. I understand that the following are effective forms of birth control:	Initials: _____
Primary methods:	Secondary forms:
• tubal sterilization (tying my tubes)	Barrier:
• partner's vasectomy	• male latex condom with or without spermicide
• intrauterine device	• diaphragm with spermicide
• hormonal combination birth control pills,	• cervical cap with spermicide
• skin patches, shots, under-the-skin implants or vaginal ring	Other:
	• vaginal sponge (contains spermicide)

A diaphragm and cervical cap must each be used with spermicide, a special cream that kills sperm. I understand that at least one of my two forms of birth control must be a primary method.

6. I will talk with my doctor about any medicines including herbal products I am taking during my isotretinoin therapy. I understand hormonal birth control methods may not work if I am taking certain medicines or herbal products.

7. I may receive a free birth control counseling session from a doctor or other family planning expert. My isotretinoin doctor can give me an isotretinoin Patient Referral Form for this free consultation.

8. I must begin using the birth control methods I have chosen as described above at least one month before I start taking isotretinoin.

9. I cannot get my first prescription for isotretinoin unless my doctor has told me that I have two negative pregnancy test results. The first pregnancy test should be done when my doctor decides to prescribe isotretinoin. The second pregnancy test must be done in a lab during the first 5 days of my menstrual period right before starting isotretinoin therapy treatment or as instructed by my doctor. I will then have one pregnancy test, in a lab:

- every month during treatment
- at the end of treatment
- and 1 month after stopping treatment

10. I must not start taking isotretinoin until I am sure that I am not pregnant, have negative results from two pregnancy tests, and the second test has been done in a lab.

11. I have read and understand the materials my doctor has given to me, including *The IPLEDGE Program Guide for Isotretinoin for Female Patients Who Can Get Pregnant*, *The IPLEDGE Birth Control Workbook* and *The IPLEDGE Program Patient Introductory Brochure*.

My doctor gave me and asked me to watch the DVD containing a video about birth control and a video about birth defects and isotretinoin.

I was told about a private counseling line that I may call for more information about birth control. I have noted this information on emergency line 1-800-495-0654.

11. I must stop taking isotretinoin right away and call my doctor if I get pregnant, miss my expected menstrual period, stop using birth control or have sexual intercourse without using my two birth control methods at any time.

12. My doctor gave me information about the purpose and importance of providing information to the iPLEDGE program should I become pregnant while taking isotretinoin or within one month of the last dose. I understand that if I become pregnant, my pregnancy, my health, and my baby's health may be shared with the makers of isotretinoin, authorized parties who maintain the iPLEDGE program for the makers of isotretinoin and government health regulatory authorities.

13. I understand that being qualified to receive isotretinoin in the iPLEDGE program means that I:

- have had two negative urine or blood pregnancy tests before receiving the first isotretinoin prescription. The second test must be done in a lab. I must have a negative result from a urine or blood pregnancy test done in a lab repeated each month before I receive another isotretinoin prescription.
- have chosen and agreed to use two forms of effective birth control at the same time. At least one method must be a form of birth control, unless I have chosen never to have sexual contact with a male (abstinence), or I have undergone a hysterectomy. I must use two forms of birth control for at least one month before I start isotretinoin therapy, during therapy and for one month after I stop therapy. I must receive counseling, repeated on a monthly basis, about birth control and behaviors associated with an increased risk of pregnancy.
- have signed a Patient Information/Informed Consent About Birth Defects (for female patients who can get pregnant) that contains warnings about the chance of possible birth defects if I am pregnant or become pregnant and my unborn baby is exposed to isotretinoin.
- have interacted with the iPLEDGE program before starting isotretinoin and on a monthly basis to answer questions on the program requirements and to enter my two chosen forms of birth control.

Initials: _____

My doctor has answered all my questions about isotretinoin and I understand that it is my responsibility to get pregnant one month before, during isotretinoin treatment, or for one month after I stop taking isotretinoin.

Initials: _____

I now authorize my doctor _____ to begin my treatment with isotretinoin.

Patient Signature: _____ Date: _____

Parent/Guardian Signature (if under age 18): _____ Date: _____

Please print: Patient Name and Address _____

Telephone _____

I have fully explained to the patient, _____, the nature and purpose of the treatment described above and to the risks to female patients of childbearing potential. I have asked the patient if she has any questions regarding her treatment with isotretinoin and have answered those questions to the best of my ability.

Doctor Signature: _____ Date: _____

PLACE THE ORIGINAL SIGNED DOCUMENTS IN THE PATIENT'S MEDICAL RECORD. PLEASE PROVIDE A COPY TO THE PATIENT.

PATIENT INFORMATION/INFORMED CONSENT (for all patients)

To be completed by patient (and parent or guardian if patient is under age 18) and signed by the doctor.

Read each item below and initial in the space provided if you understand each item and agree to follow your doctor's instructions. A parent or guardian of a patient under age 18 must also read and understand each item before signing the agreement.

Do not sign this agreement and do not take isotretinoin if there is anything that you do not understand about all the information you have received about using isotretinoin.

1. I, _____,

(Patient's Name)

understand that isotretinoin is a medicine used to treat severe nodular acne that cannot be cleared up by any other acne treatments, including antibiotics. In severe nodular acne, many red, swollen, tender lumps form in the skin. If untreated, severe nodular acne can lead to permanent scars.

Initials: _____

2. My doctor has told me about all my choices for treating my acne.

Initials: _____

3. I understand that there are serious side effects that may happen while I am taking isotretinoin. These have been explained to me. These side effects include serious birth defects in babies of pregnant patients. [Note: There is a second Patient Information/Informed Consent About Birth Defects (for female patients who can get pregnant)].

4. I understand that some patients, while taking isotretinoin or soon after stopping isotretinoin, have become depressed or developed other serious mental problems. Symptoms of depression include sad, "anxious" or empty mood; irritability; acting on dangerous impulses; anger; loss of pleasure or interest in social or sports activities; sleeping too much or too little; changes in weight or appetite; school or work performance going down or trouble concentrating. Some patients taking isotretinoin have had thoughts about hurting themselves or putting an end to their own lives (suicidal thoughts). Some people tried to end their own lives to have ended their own lives. There were reports that some of these people did not appear depressed. There have been reports of patients on isotretinoin becoming aggressive or violent. No one knows if isotretinoin caused these behaviors or if they would have happened even if the person did not take isotretinoin. Some people have had other signs of depression while taking isotretinoin (see #7 below).

5. Before I start taking isotretinoin, I agree to tell my doctor if I have ever had symptoms of depression (see #7 below), been psychotic, attempted suicide, had any other mental problems or take medicine for any of these problems. Being psychotic means having a loss of contact with reality, such as hearing voices or seeing things that are not there.

6. Before I start taking isotretinoin, I agree to tell my doctor if, to the best of my knowledge, anyone in my family has had symptoms of depression, been psychotic, attempted suicide or had any other serious mental problems.

Initials: _____

7. Once I start taking isotretinoin, I agree to stop using isotretinoin and tell my doctor right away if any of the following signs and symptoms of depression or psychosis happen:

- Start to feel sad or have crying spells
- Lose interest in activities I once enjoyed
- Sleep too much or have trouble sleeping
- Become more irritable, angry, or aggressive than usual (for example, temper outbursts, thoughts of violence)
- Have a change in my appetite or body weight
- Have trouble concentrating
- Withdraw from my friends or family
- Feel like I have no energy
- Have feelings of worthlessness or guilt
- Start having thoughts about hurting myself or taking my own life (suicidal thoughts)
- Start acting on dangerous impulses
- Start seeing or hearing things that are not real

Initials: _____

8. I agree to return to see my doctor every month I take isotretinoin to get a new prescription for isotretinoin, to check my progress and to check for signs of side effects.

9. Isotretinoin will be prescribed just for me — I will not share isotretinoin with other people because it may cause serious side effects, including birth defects.

Initials: _____

10. I will not give blood while taking isotretinoin or for one month after I stop taking isotretinoin. I understand that if someone who is pregnant gets my donated blood, her baby may be exposed to isotretinoin and may be born with serious birth defects.

Initials: _____

11. I have read *The iPLEDGE Program Patient Introductory Brochure*, and other materials my provider gave me containing important safety information about isotretinoin. I understand all the information I received. Initials: _____

12. My doctor and I have decided I should take isotretinoin. I understand that all the information in the iPLEDGE program to have my prescription filled. I understand that I can stop taking isotretinoin at any time. I agree to tell my doctor if I stop taking isotretinoin.

I now allow my doctor _____ to begin my treatment with isotretinoin.

Patient Signature: _____ Date: _____

Parent/Guardian Signature (if under age 18): _____ Date: _____

Patient Name (print) _____

Patient Address _____ Telephone (_____) _____

I have:

- fully explained to the patient, _____, the nature and purpose of isotretinoin treatment, including its benefits and risks,
- given the patient the appropriate educational materials, *The IPLEDGE Program Patient Introductory Brochure* and asked the patient if he/she has any questions regarding his/her treatment with isotretinoin
- answered those questions to the best of my ability

Doctor Signature: _____ Date: _____

PLACE THE ORIGINAL SIGNED DOCUMENTS IN THE PATIENT'S MEDICAL RECORD. PLEASE PROVIDE A COPY TO THE PATIENT.