

(Continued from other side)

What should I avoid while taking Accutane?

- Do not get pregnant while taking Accutane and for one month after stopping Accutane. See "What is the most important information I should know about Accutane?"
- Do not breast feed while taking Accutane and for one month after stopping Accutane. We do not know if Accutane can pass through your milk and harm the baby.
- Do not take blood while you take Accutane and for one month after stopping Accutane. If someone who is pregnant gets your donated blood, their baby may be exposed to Accutane and may be born with birth defects.
- Do not take other medicines or herbal products with Accutane unless you talk to your doctor. See "What should I tell my doctor before taking Accutane?"
- Do not drive at night until you know if Accutane has affected your vision. Accutane may decrease your ability to see in the dark.
- Do not have cosmetic procedures to smooth your skin, including waxing, dermabrasion, or laser procedures, while you are using Accutane and for at least 6 months after you stop. Accutane can increase your chance of scarring from these procedures. Check with your doctor for advice about when you can have cosmetic procedures.
- Avoid sunlight and ultraviolet lights as much as possible. Tanning machines use ultraviolet lights. Accutane may make your skin more sensitive to light.
- Do not share Accutane with other people. It can cause birth defects and other serious health problems.

What are the possible side effects of Accutane?

- Accutane can cause birth defects (deformed babies), loss of a baby before birth (miscarriage), death of the baby, and early (premature) births. See "What is the most important information I should know about Accutane?"
- Accutane may cause serious mental health problems. See "What is the most important information I should know about Accutane?"
- serious brain problems. Accutane can increase the pressure in your brain. This can lead to permanent loss of eyesight and, in rare cases, death. Stop taking Accutane and call your doctor right away if you get any of these signs of increased brain pressure:
 - bad headache
 - blurred vision
 - dizziness
 - nausea or vomiting
 - seizures (convulsions)
 - stroke

- skin problems. Skin rash can occur in patients taking Accutane. In some patients a rash can be serious. Stop using Accutane and call your doctor right away if you develop conjunctivitis (red or inflamed eyes, like "pink eye"), a rash with a fever, blisters on legs, arms or face and/or sores in your mouth, throat, nose, eyes, or if your skin begins to peel.
- stomach area (abdomen) problems. Certain symptoms may mean that your internal organs are being damaged. These organs include the liver, pancreas, bowel (intestines), and esophagus (connection between mouth and stomach). If your organs are damaged, they may not get better even after you stop taking Accutane. Stop taking Accutane and call your doctor if you get:
 - severe stomach, chest or bowel pain
 - trouble swallowing or painful swallowing
 - new or worsening heartburn
 - diarrhea
 - rectal bleeding
 - yellowing of your skin or eyes
 - dark urine

- bone and muscle problems. Accutane may affect bones, muscles, and ligaments and cause pain in your joints or muscles. Tell your doctor if you plan hard physical activity during treatment with Accutane. Tell your doctor if you get:
 - back pain
 - joint pain
 - broken bone. Tell all healthcare providers that you take Accutane if you break a bone.

Stop Accutane and call your doctor right away if you have muscle weakness, muscle weakness with or without pain can be a sign of serious muscle damage.

- Accutane may stop long bone growth in teenagers who are still growing.
- hearing problems. Stop using Accutane and call your doctor if your hearing gets worse or if you have ringing in your ears. Your hearing loss may be permanent.
- vision problems. Accutane may affect your ability to see in the dark. This condition usually clears up after you stop taking Accutane, but it may be permanent. Other serious eye effects can occur. Stop taking Accutane and call your doctor right away if you have any problems with your vision or dryness of the eyes that is painful or constant. If you wear contact lenses, you may have trouble wearing them while taking Accutane and after treatment.

- lipid (fats and cholesterol in blood) problems. Accutane can raise the level of fats and cholesterol in your blood. This can be a serious problem. Return to your doctor for blood tests to check your lipids and to get any needed treatment. These problems usually go away when Accutane treatment is finished.
- serious allergic reactions. Stop taking Accutane and get emergency care right away if you develop hives, a swollen face or mouth, or have trouble breathing. Stop taking Accutane and call your doctor if you get a fever, rash, or red patches or bruises on your legs.

- blood sugar problems. Accutane may cause blood sugar problems including diabetes. Tell your doctor if you are very thirsty or urinate a lot.
- decreased red and white blood cells. Call your doctor if you have trouble breathing, faint, or feel weak.
- The common, less serious side effects of Accutane are dry skin, chapped lips, dry eyes, and dry nose that may lead to nosebleeds. Call your doctor if you get any side effect that bothers you or that does not go away.

- These are not all of the possible side effects with Accutane. Your doctor or pharmacist can give you more detailed information.
- Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088 or JG Pharma, Inc. at 1-844-325-3350.

- How should I store Accutane?
 - Store Accutane at 68° to 77°F (20° to 25°C). Protect from light.
 - Keep Accutane and all medicines out of the reach of children.

General Information about Accutane
Medicines are sometimes prescribed for conditions that are not mentioned in Medication Guides. Do not use Accutane for a condition for which it was not prescribed. Do not give Accutane to other people, even if they have the same symptoms that you have. It may harm them.

This Medication Guide summarizes the most important information about Accutane. If you would like more information, talk with your doctor. You can ask your doctor or pharmacist for information about Accutane that is written for healthcare professionals. You can also call iPLEDGE Program at 1-866-495-0654 or visit www.ipledeprogram.com.

What are the ingredients in Accutane?
Active Ingredient: isotretinoin USP
Inactive Ingredients: butylated hydroxyanisole, edetate disodium, hydrogenated vegetable oil (Type-I and Type-II), medium chain triglyceride, refined soybean oil and white wax. Gelatin capsules contain ferric oxide red, ferric oxide yellow (for 30 mg), gelatin, glycerin, methacrylates, propyl paraben, lake blend blue (LB-332) containing D&C Yellow No. 10, FD&C Blue No. 1 (for 10 mg), lake blend red (LB-1574) containing D&C Red No. 27, D&C Red No. 30 (for 20 mg), lake blend green (LB-333) containing D&C Yellow No. 10, FD&C Blue No. 1 (for 40 mg), lake blend white (TLB-1774) containing FD&C Blue No. 2, titanium dioxide, and opacode black S-1-17823 containing iron oxide black, N-butyl alcohol, propylene glycol, ammonium hydroxide and shellac.

This Medication Guide has been approved by the U.S. Food and Drug Administration. Dilant is a registered trademark of Warner-Lambert Company LLC. To reorder additional Medication Guides contact JG Pharma's Customer Service at 1-844-256-4677.

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Neurological
parosmum/cerebr (see **WARNINGS: Pseudotumor Cerebri**), dizziness, drowsiness, headache, insomnia, lethargy, malaise, nervousness, paresthesias, seizures, stroke, syncope, weakness.
Psychiatric
suicidal ideation, suicide attempt, suicide, depression, psychosis, aggression, violent behaviors (see **WARNINGS: Psychiatric Disorders**), emotional instability.
Hepatocholesterol
abnormal mentes.
Respiratory System
bronchospasms (with or without a history of asthma), respiratory infection, voice alteration.
Skin and Appendages
skin lesions, alopecia (which in some cases persists), hirsutism, cheilitis (dry lips), dry mouth, dry nose, dry skin, epistaxis, erythema xanthomas,¹ erythema multiforme, flushing, fragility of skin, hair abnormalities, hirsutism, hyperpigmentation and hyperpigmentation, infections (including disseminated herpes simplex), nail dystrophy, onychia, peeling of palms and soles, photosensitivity/
photosensitizing reactions, pruritis, pruritic eruptions, 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 or exuberant granulation tissue with crusting); see **PRECAUTIONS: Information for Patients**.
Special Senses
Hearing
hearing impairment (see **WARNINGS: Hearing Impairment**, tinnitus).
Vision
contact optics (see **WARNINGS: Contact Optics**), 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 uropathological 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), SGP (ALT), GGT or LDH (see **WARNINGS: Hepatotoxicity**).
Elevation of fasting blood sugar, elevations of CPK (see **PRECAUTIONS: Laboratory Tests**), hyperkalemia.
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.

OVERDOSEAGE
The oral LD50 of isotretinoin is greater than 4000 mg/kg in rats and 1500 times the recommended clinical dose of 1 mg/kg/day after normalization of the rat dose for total body surface area and >200 times the recommended clinical dose of 1 mg/kg/day after normalization of the mouse dose for total body surface area and is approximately 1900 mg/kg in rabbits (833 times the recommended clinical dose of 1 mg/kg/day after normalization for total body surface area). In humans, overdose has been associated with vomiting, facial flushing, cheilitis, abdominal pain, headache, dizziness, and nausea. These symptoms usually resolve without apparent residual effects.
Accutane causes serious birth defects at any dosage (see **Based CONTRAINDICATIONS AND WARNINGS**). Patients who can become pregnant 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 **Based CONTRAINDICATIONS AND WARNINGS**. Non-pregnant patients must be warned to avoid pregnancy for at least one month after receiving 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 women than found during a normal treatment course, male patients should use a condom, or avoid reproductive sexual activity with a 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.
DOSEAGE AND ADMINISTRATION
Accutane should be administered with a meal (see **PRECAUTIONS: Information for Patients**).
The recommended dosage range for Accutane is 0.5 to 1 mg/kg/day given in two divided doses with food for 15 to 20 weeks. In studies comparing 0.5, 1.0, and 1 mg/kg/day, it was found that all dosages provided initial clearing of disease, but that 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 Accutane with food will significantly decrease absorption. Before upward dose adjustments are made, the patient must be questioned about their compliance with food restrictions.
The safety of once daily dosing with Accutane has not been established. Once daily dosing is not recommended. Once daily dosing may be discontinued. After a period of 2 months or more off therapy, and if the total retinol level has returned to or below the level of the first dose, but not been studied, and is not recommended. It is important that Accutane be given at the recommended dose for 15 to 20 weeks and is warranted by persistent or recurring severe nodular acne, a second course of therapy may be initiated. The optimal interval between treatments has not been defined for patients who have not completed their course. Long term use of Accutane, even in low doses, has not been studied, and is not recommended. It is important that Accutane be given at the recommended dose for no longer than the recommended duration. The effect of long-term use of Accutane on bone loss is unknown (see **WARNINGS: Skeletal: Bone Mineral Density, Hypoparathyroidism, and Premature Epiphyseal Closure**).
Contraceptive measures must be followed for any subsequent course of therapy (see **PRECAUTIONS**).
Table 4 Accutane Dosing by Body Weight (Based on Information with Food)

kilograms	Body Weight		Total mg/day	
	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 **DOSEAGE AND ADMINISTRATION**; the recommended dosage range is 0.5 to 1 mg/kg/day.

INFORMATION FOR PHARMACISTS
Access the iPLEDGE Program system via the internet (www.ipledeprogram.com), telephone (1-866-495-0654) or through electronic telecommunication verification (via submission of an isotretinoin prescription claim) to obtain an authorization and the "do not dispense to patient alert" (see **WARNINGS: Information for Patients**).
REFILLS REQUIRE A NEW PRESCRIPTION AND A NEW AUTHORIZATION FROM THE IPLEDGE SYSTEM.
A Accutane Medication Guide must be given to the patient each time Accutane is dispensed, as required by law. This Accutane Medication Guide is an important part of the risk management program for the patient.

HOW SUPPLIED
Accutane (isotretinoin) capsules USP 10 mg are opaque blue elliptical soft gelatin capsules imprinted with black ink, "R133" on one side and are supplied in boxes of 30 containing 3 prescription packs of 10 capsules and unit dose blisters. NDC 2143-231-30
Accutane (isotretinoin) capsules USP 20 mg are opaque pink elliptical soft gelatin capsules imprinted with black ink, "R136" on one side and are supplied in boxes of 30 containing 3 prescription packs of 10 capsules and unit dose blisters. NDC 2143-232-30
Accutane (isotretinoin) capsules USP 30 mg are yellow round biconvex caplets, oval-shaped soft gelatin capsules imprinted with "10" in black colored ink along the length of body on one side and are supplied in boxes of 30 containing 3 prescription packs of 10 capsules and unit dose blisters. NDC 2143-233-30
Accutane (isotretinoin) capsules USP 40 mg are opaque green elliptical soft gelatin capsules imprinted with black ink, "R137" on one side and are supplied in boxes of 30 containing 3 prescription packs of 10 capsules and unit dose blisters. NDC 2143-234-30
STORAGE
Store at 20° to 25°C (68° to 77°F) (see USP controlled room temperature). Protect from light.

REFERENCES
1. Prock CC, Stein TG, Yoder FK et al. Prolonged retention of cyclic and conjugate acene with 13-cis-retinoic acid. *N Engl J Med* 2000;329:333, 1979.
2. Prock CC, Shalit M, Strauss JG, Rivner OJ, Ripstein B, et al. The consensus conference on acne classification. *J Am Acad Dermatol* 1994;49:500, 1991.
3. Farrell LN, Strauss JS, Strazniel AM. The treatment of severe cystic acne with 13-cis-retinoic acid: evaluation of sebium production and the clinical response in a multiple-dose trial. *J Am Acad Dermatol* 1982;6:11, 1982.
4. Jones H, Blanc D, Curth M, et al. 13-cis-retinoic acid and acne. *Lancet* 2:1048-1049, 1980.
5. Katz RA, Jorgensen K, Nigra TP. Elevation of serum triglyceride levels from oral isotretinoin in disorders of keratinization. *Arch Dermatol* 116:1309-1312, 1980.
6. Ellis CH, Madison KE, Frenkel KB, Marone W, Kormanik JJ. Isotretinoin therapy in association with early skeletal radiographic changes. *J Am Acad Dermatol* 1994;30:1204-1208, 1994.
7. Dickson CH, Connelly SM. Enzyme nomenclature associated with isotretinoin (13-cis-retinoic acid). *Arch Dermatol* 116:951-952, 1980.
8. Strauss JS, Rapin RP, Shalit M, et al. Isotretinoin therapy for acne: results of a multicenter dose-response study. *J Am Acad Dermatol* 10:490-496, 1984.

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

- 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.
Initial: _____
- 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.
Initial: _____
- I understand that I must avoid having any sexual contact (penis-vaginal) with a partner who could get me pregnant 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 my doctor has medically confirmed that I am post-menopausal.
Initial: _____
- I understand that hormonal birth control products are among the most effective forms of birth control, and combination birth control pills and other hormonal products include skin patches, shots, under-the-skin implants, vaginal rings, and intrauterine devices (IUDs). Any method of birth control can fail. That is why I must use two different birth control forms at the same time, starting one month before, during, and for one month after stopping therapy every time I have any sexual contact (penis-vaginal) with a partner who could get me pregnant, even if one of the forms I choose is hormonal birth control.
Initial: _____
- I understand that the following are effective forms of birth control:
Primary forms:
 - using my tubes (tubal sterilization)
 - male vasectomy
 - intrauterine device
 - hormonal combination birth control pills, skin patches, shots, under-the-skin implants or vaginal ring)Secondary forms:
 - male latex condom with or without spermicide
 - diaphragm with spermicide
 - cervical cap with spermicide
 - 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 form.
Initial: _____

- I will talk with my doctor about any medicines including herbal products I plan to take during my isotretinoin treatment because hormonal birth control forms may not work if I am taking certain medicines or herbal products.
Initial: _____
- 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 Contraception Referral Form for this free consultation.
Initial: _____
- I must begin using the birth control forms I have chosen as described above at least one month before I start taking isotretinoin.
Initial: _____
- 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 before starting isotretinoin therapy, 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

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.
Initial: _____

- I have read and understand the materials my doctor has provided to me, including the *Guide to Isotretinoin for Patients Who Can Get Pregnant*, *Birth Control Workbook* and *Patient Introductory Brochure*.
I have received information on emergency birth control.
My doctor provided me and asked me to watch a video about birth control and a video about birth defects and isotretinoin.
Initial: _____

- 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 any sexual contact (penis-vaginal) with a partner who could get me pregnant without using my two birth control forms at any time.
Initial: _____
- My doctor provided 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, information about 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.
Initial: _____
- 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 form must be a primary form of birth control, unless I have chosen never to have any sexual contact (penis-vaginal) with a partner who could get me pregnant (abstinence), or I have undergone a hysterectomy or bilateral oophorectomy, or I have been medically confirmed to be post-menopausal. 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 stopping therapy. I must be getting 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 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 been informed of and understand the purpose and importance of providing information to the iPLEDGE Program should I become pregnant while taking isotretinoin or within 1 month of the last dose.
 - 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.

Initial: _____
My doctor has answered all my questions about isotretinoin and I understand that it is my responsibility not to get pregnant one month before, during isotretinoin treatment, or for one month after I stop taking isotretinoin.

I now authorize my doctor _____ Date: _____ 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 the risks to patients who can get pregnant. I have asked the patient if there are any questions regarding treatment with isotretinoin and have answered those questions to the best of my ability.
Date: _____
Doctor Signature: _____

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.

- 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: _____

I have fully explained to the patient, _____, the nature and purpose of the treatment described above and the risks to patients who can get pregnant. I have asked the patient if there are any questions regarding treatment with isotretinoin and have answered those questions to the best of my ability.
Date: _____
Doctor Signature: _____

PLACE THE ORIGINAL SIGNED DOCUMENTS IN THE PATIENT'S MEDICAL RECORD. PLEASE PROVIDE A COPY TO THE PATIENT.
Active Ingredients: isotretinoin USP
Inactive Ingredients: butylated hydroxyanisole, edetate disodium, hydrogenated vegetable oil (Type-I and Type-II), medium chain triglyceride, refined soybean oil and white wax. Gelatin capsules contain ferric oxide red, ferric oxide yellow (for 30 mg), gelatin, glycerin, methacrylates, propyl paraben, lake blend blue (LB-332) containing D&C Yellow No. 10, FD&C Blue No. 1 (for 10 mg), lake blend red (LB-1574) containing D&C Red No. 27, D&C Red No. 30 (for 20 mg), lake blend green (LB-333) containing D&C Yellow No. 10, FD&C Blue No. 1 (for 40 mg), lake blend white (TLB-1774) containing FD&C Blue No. 2, titanium dioxide, and opacode black S-1-17823 containing iron oxide black, N-butyl alcohol, propylene glycol, ammonium hydroxide and shellac.
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- My doctor has told me about my choices for taking my acne.
Initials: _____
- 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 patients who can get pregnant)).
Initials: _____
- I understand that some patients, while taking isotretinoin or soon after stopping isotretinoin, have become depressed or developed other 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 harm on and to their own lives (suicidal thoughts). Some people tried to end their own lives. And some people 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).
Initials: _____
- 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.
Initials: _____
- Before I start taking isotretinoin, I agree to tell my doctor if, to the best of my knowledge, anyone in my family has ever had symptoms of depression, been psychotic, attempted suicide, or had any other serious mental problems.
Initials: _____

- 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. I:
 - Start to feel sad or have crying spells
 - Loss 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 yourself or taking my own life (suicidal thoughts)
 - Start acting on dangerous impulses
 - Start seeing or hearing things that are not real

- 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.
Initials: _____

- 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: _____
- I will not give blood while taking isotretinoin or for 1 month after I stop taking isotretinoin. I understand that if someone who is pregnant gets my donated blood, their baby may be exposed to isotretinoin and may be born with serious birth defects.
Initials: _____

- I have read the *Patient Introductory Brochure*, and other materials my provider provided me containing important safety information about isotretinoin. I understand all the information I received.
Initials: _____
- My doctor and I have decided I should take isotretinoin. I understand that I must be qualified in the iPLEDGE Program to have my prescription filled each month. I understand that I can stop taking isotretinoin at any time. I agree to tell my doctor if I stop taking isotretinoin.
Initials: _____

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
- provided the patient with the appropriate educational materials, such as the *Patient Introductory Brochure* and asked the patient if there are any questions regarding their 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.

Medication Guide ACCUTANE® (ACK-U-TAINE) (isotretinoin) capsules

Read the Medication Guide that comes with Accutane before you start taking it and each time you get a prescription. There may be new information. This information does not take the place of advice from your doctor about your condition or your treatment.
What is the most important information I should know about Accutane?

- Accutane is used to treat a type of severe acne (nodular acne) that has not been helped by other treatments, including antibiotics.
- Because Accutane can cause birth defects, Accutane is only for patients who can understand and agree to carry out all of the instructions in the iPLEDGE Program.
- Accutane may cause serious mental health problems.
- Birth defects (deformed babies), loss of a baby before birth (miscarriage), death of the baby, and early (premature) births. Patients who are pregnant or who plan to become pregnant must not take Accutane. Patients must not get pregnant:
 - for 1 month before starting Accutane
 - while taking Accutane
 - for 1 month after stopping Accutane

Do not get pregnant while taking Accutane, stop taking it right away and call your doctor. Doctors and patients should report all cases of pregnancy to:

- FDA MedWatch at 1-800-FDA-1088, and
- The iPLEDGE Pregnancy Registry at 1-866-495-0654

2. Serious mental health problems. Accutane may cause:

- depression
- psychotic (seeing or hearing things that are not real)
- suicide. Some patients taking Accutane have had thoughts about hurting themselves or putting an end to their own lives (suicidal thoughts). Some people tried to end their own lives. And some people have ended their own lives.

Stop Accutane and call your doctor right away if you or a family member notices that you have any of the following signs and symptoms of depression or psychosis:

- start to feel sad or have crying spells
- lose interest in activities you 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 your appetite or body weight
- have trouble concentrating
- withdraw from your friends or family
- feel like you have no energy
- have feelings of worthlessness or guilt
- start having thoughts about hurting yourself or taking your own life (suicidal thoughts)
- start acting on dangerous impulses
- start seeing or hearing things that are not real

After stopping Accutane, you may also need follow-up mental health care if you had any of these symptoms.
What is Accutane?
Accutane is a medicine taken by mouth to treat the most severe form of acne (nodular acne) that cannot be cleared up by any other acne treatments, including antibiotics. Accutane can cause serious side effects such as "What is the most important information I should know about Accutane?". Accutane can only be used by patients who are registered in the iPLEDGE Program.

- dispensed by a pharmacy that is registered with the iPLEDGE Program
- given to patients who are registered in the iPLEDGE Program and agree to do everything required in the program

What is severe nodular acne?
Severe nodular acne is when many red, swollen, tender lumps form in the skin. These can be the size of pencil erasers or larger. If untreated, nodular acne can lead to permanent scars.
Who should not take Accutane?

- Do not take Accutane if you are pregnant, plan to become pregnant, or become pregnant during Accutane treatment. Accutane causes severe birth defects. See "What is the most important information I should know about Accutane?"
- Do not take Accutane if you are allergic to anything in it. See the end of this Medication Guide for a complete list of ingredients in Accutane. Accutane contains parabens as the preservatives.

What should I tell my doctor before taking Accutane?
Tell your doctor if you or a family member has any of the following health conditions:

- mental problems
- asthma
- liver disease
- diabetes
- heart disease
- bone loss (osteoporosis or weak bones)
- an existing problem called axonal neuropathy (where people eat too little)
- food or medicine allergies

Tell your doctor if you are pregnant or breastfeeding. Accutane must not be used by women who are pregnant or breastfeeding. Accutane and certain other medicines can interact with each other, sometimes causing serious side effects. Especially tell your doctor if you take:

- Vitamin A supplements (Vitamin A in high doses has many of the same side effects as Accutane. Taking both together may increase your chance of getting side effects such as:
- Tetracycline antibiotics. Tetracycline antibiotics taken with Accutane can increase the chances of getting increased pressure in the brain.
- Phenothiaz