

Accutane® (Isotretinoin Capsules USP) 10 mg, 20 mg, 30 mg and 40 mg Capsules

Rx Only
Revised: September 2022

CAUSES BIRTH DEFECTS



CONTRAINDICATIONS AND WARNINGS
Accutane® must not be used by patients who are or may become pregnant. There is an extremely high risk that life-threatening birth defects will result if pregnancy occurs while taking Accutane 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 when an exposed fetus has been affected.
Birth defects which have been documented following Accutane exposure include abnormalities of the face, eyes, ears, skull, central nervous system, cardiovascular system, and thumbs 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 aural, 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 atrophy, hydrocephalus, microcephaly, central nerve defects), cardiovascular abnormalities, thyroid 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 patient who is taking Accutane, Accutane must be discontinued immediately and the patient should be referred to an Obstetrician/Gynecologist experienced in reproductive toxicity for further evaluation and counseling.

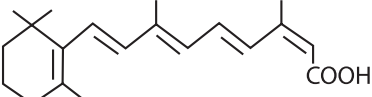


Because of Accutane's teratogenicity and to minimize fetal exposure, Accutane is approved for marketing only under a special restricted distribution program approved by the FDA. The program is called the iPLEDGE® Accutane REMS. Accutane must only be prescribed by and dispensed to patients who are enrolled and activated with iPLEDGE. Accutane must only be dispensed by a pharmacy enrolled and activated with iPLEDGE, and must only be dispensed to patients who are enrolled and meet all the requirements of iPLEDGE (see PRECAUTIONS).

Table 1 Monthly Required iPLEDGE Interactions

	Patients Who Can Become Pregnant	Patients Who Cannot Become Pregnant
PRESCRIBER		
Confirms patient counseling	X	X
Enters the 2 contraception forms chosen by the patient	X	
Enters pregnancy test results	X	
PATIENT		
Answers educational questions before every prescription	X	
Enters 2 forms of contraception	X	
PHARMACIST		
Contacts system to get an authorization	X	X

DESCRIPTION
Isotretinoin USP, a retinoid, is available as Accutane (isotretinoin capsules, USP) in 10 mg, 20 mg, 30 mg and 40 mg soft gelatin capsules for oral administration. Each capsule contains butylated hydroxyanisole, edetate disodium, hydrogenated vegetable oil (Type-I and Type-II), medium chain triglyceride, refined soybean oil and white wax. Gelatin capsules contain ferrous oxide red, ferric oxide black (for 30 mg), gelatin, glycerin, methyl paraben, propyl paraben, blend blue (LB-333 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 10 mg), lake blend red (LB-1774) containing FD&C No. 2, titanium dioxide, and opaque black 5-1-17822 containing iron oxide black, N-butyl alcohol, propylene glycol, ammonium hydroxide and shellac. Chemical structure is 13-*cis*-retinoic acid and is related to both retinoic acid and retinol (vitamin A). It is a yellow to slightly gray crystalline powder with a molecular weight of 300.44. The structural formula is:



US DISCUSSION

Clinical Pharmacology

Isotretinoin is a retinoid, which when administered in pharmacologic dosages of 0.5 to 1 mg/kg/day (see **DOSEAGE 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 Accutane, and reflects a reduction in sebaceous gland size and an inhibition of sebaceous gland differentiation¹.

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 Accutane under fasted and fed conditions. Both peak plasma concentration (C_{max}) and the area under the curve (AUC) of isotretinoin were more than doubled following a standardized high-fat meal when compared with Accutane 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, Accutane capsules should always be taken with food (see **DOSEAGE 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.

Table 2 Pharmacokinetic Parameters of Isotretinoin Mean (SD), N=74

Accutane 2 x 40 mg Capsules	AUC ₀₋₂₄ (ng-hr/mL)	C _{max} (ng/mL)	T _{max} (hr)	t _{1/2} (hr)
Fasted	10,004 (22%)	862 (22%)	5.3 (77%)	21 (39%)
Fasted	3,703 (46%)	301 (63%)	3.2 (56%)	21 (20%)

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

Metabolism
Following oral administration of Accutane, 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 to the other. Isotretinoin is also irreversibly oxidized to 4-*oxo*-isotretinoin, which has been shown to be the major metabolite in human plasma.

After a single 80 mg oral dose of Accutane 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.

These three metabolites possess biological activity that is seen in some *in vitro* models more than that of the parent isotretinoin. However, the clinical significance of these models is unknown. After multiple oral administration of isotretinoin to adult cystic 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 *in vitro* studies indicating that the primary P450 isozymes involved in isotretinoin metabolism are C2C8, 2C9, 3A4, and 2B6, isotretinoin and its metabolites are further metabolized into conjugates, and then excreted in urine and feces.

Elimination
Following oral administration of an 80 mg dose of 13-*cis*-retinoic acid as a liquid suspension, ³C-activity in blood declined with a half-life of 90 hours. The metabolites of isotretinoin and any conjugates of isotretinoin in blood were found to have a half-life of 12 to 15 hours. In a study of 14 healthy adult subjects, the elimination half-life of isotretinoin under fasted conditions was approximately 16 hours. The mean $\pm$ SD elimination half-lives (t_{1/2}) of 4-*oxo*-isotretinoin and 4-*oxo*-tretinoin were 21 $\pm$ 8.2 hours and 24 $\pm$ 5.3 hours, respectively. After a single patient and multiple doses, the observed accumulation ratios of isotretinoin ranged from 0.9 to 5.43 in patients with cystic 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 Accutane for the treatment of severe acne. The pharmacokinetics of isotretinoin were also evaluated. The dose-normalized pharmacokinetic parameters for 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.

Table 3 Pharmacokinetic Parameters of Isotretinoin Following Single and Multiple Dose Administration in Pediatric Patients, 12 to 15 Years of Age Mean (± SD), N=30*

Parameter	Single Dose (n=15)	Multiple Dose (n=15)
C _{max} (ng/mL)	573.25 (278.79)	731.98 (361.86)
AUC ₀₋₂₄ (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 (± 0.24)	4.0 (± 1.2)
C ₁₂₋₁₈ (ng/mL)	—	352.32 (184.44)
t _{1/2} (hr)	—	15.69 (± 1.2)
CL/F (L/hr)	—	17.96 (± 6.27)

*The 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.
‡12 to 15 years, the mean $\pm$ SD elimination half-lives (t_{1/2}) of isotretinoin and 4-*oxo*-isotretinoin were 15.7 $\pm$ 5.1 hours and 23.1 $\pm$ 5.7 hours, respectively. The accumulation ratios of isotretinoin ranged from 0.46 to 3.65 for pediatric patients.

INDICATIONS AND USAGE
Accutane 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 become suppurative or hemorrhagic. "Severe," by definition, means "many" as opposed to "few or several" nodules. Because of significant adverse effects associated with its use, Accutane should be reserved for patients with severe nodular acne who are unresponsive to conventional therapy, including systemic antibiotics. In addition, Accutane is indicated only for those patients who are not pregnant, because Accutane can cause life-threatening 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 off Accutane. The optimal interval before retreatment has not been defined for patients who have not completed skeletal growth (see **WARNINGS: Skeletal: Bone Mineral Density, Hypoparathyroidism, Premature Epiphyseal Closure**).

CONTRAINDICATIONS
Pregnancy: Category X. See Boxed CONTRAINDICATIONS AND WARNINGS.
Isotretinoin REMS: Meeting the requirements for a patient who can become pregnant signifies that the patient:

Accutane is contraindicated in patients who are hypersensitive to this medication or to any of its components (see **PRECAUTIONS: Hypersensitivity**).

WARNINGS
Psychiatric Disorders
Accutane 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 of Isotretinoin*. Prescribers should be alert to the warning signs of psychiatric disorders to guide patients to receive the help they need. Therefore, prior to initiation of Accutane 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 patterns, change in weight or appetite, suicidal thoughts or attempts, restlessness, irritability, acting on dangerous impulses and persistent physical symptoms unresponsive to treatment. Patients should stop Accutane 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 Accutane therapy may be insufficient; further evaluation may be necessary. While not life threatening, changes in sleep patterns, change in weight or appetite, suicidal thoughts or attempts, restlessness, irritability, acting on dangerous impulses and persistent physical symptoms unresponsive to treatment. Patients should stop Accutane 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 Accutane therapy may be insufficient; further evaluation may be necessary. The physician should consider whether Accutane therapy is appropriate in this setting, for some patients the risks may outweigh the benefits of Accutane therapy.

Pseudotumor Cerebri
Accutane has been associated with a number of cases of pseudotumor cerebri (benign intracranial hypertension), some of which involved concomitant use of tetracyclines. Concomitant treatment with tetracyclines should therefore be avoided. Early signs and symptoms of pseudotumor cerebri include papilledema, headache, nausea and vomiting, and visual disturbances. Return to normal vision has been reported in patients who experience tinnitus or hearing impairment should discontinue Accutane immediately and be referred to a neurologist for further diagnosis and care (see **ADVERSE REACTIONS: Neurological**).

Serious Skin Reactions
Decreased vision has been reported for 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 Accutane 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. Accutane should be stopped if hypertriglyceridemia is associated with an acceptable level of if symptoms of pancreatitis occur.

Lipids
Elevations of serum triglycerides in excess of 800 mg/dL have been reported in patients treated with Accutane. Marked elevations of serum triglycerides were reported in approximately 25% of patients receiving Accutane 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 Accutane therapy. Some patients have been able to reverse triglyceride elevation by reduction in weight, reduction in fat and alcohol, and reduction in calories while continuing Accutane.¹

Blood lipid determinations should be performed before Accutane is given and then at intervals until the lipid response to Accutane is established, which usually occurs within 4 weeks. Especially careful consideration must be given to risk/benefit for patients who may be at high risk during Accutane therapy (patients with diabetes, obesity, increased alcohol intake, lipid metabolism disorders, and/or a history of lipid metabolism disorder). If Accutane 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 Accutane are unknown.

12% of individuals treated during clinical trials, some of which normalized with dosage reduction or continued use of Accutane at a lower dose. A single course of therapy for Accutane for severe recalcitrant acne, some dose density measurements at several dosages sites were not significantly decreased (lumber spine change $\sim$ 4%, and total hip change $\sim$ 2%) or were increased in the majority of patients (92% had a decrease in lumber spine bone mineral density $\sim$ 4% based on unadjusted data. Sixteen (7.9%) patients had decreases in lumber spine bone mineral density $\sim$ 4%, and all did not have significant decreases or had increases (adjusted for body mass index). Twenty-one (10.6%) patients had decreases in total hip bone mineral density $\sim$ 4%, and all did not have significant decreases or had increases (adjusted for body mass index). Follow-up studies performed in eight patients with decreased bone mineral density for up to 11 months thereafter demonstrated increasing bone density in five patients at the lumber spine, while the other three patients had lumber spine bone density measurements below baseline values. Total hip bone mineral density remained below baseline (range $\sim$ 1.6% to $\sim$ 6.9% 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 Accutane 4 months after the first course, two patients showed a decrease in mean lumber spine bone mineral density up to 3.25% (see **PRECAUTIONS: Pediatric Use**).

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ADVERSE REACTIONS: Special Senses
Hepatotoxicity
Clinical hepatitis considered to be possibly or probably related to Accutane therapy has been reported. Although, mild to moderate elevations of liver enzymes have been observed in approximately 12% of patients treated during clinical trials, some of which normalized with dosage reduction or continued use of Accutane at a lower dose. The first test is screening test is obtained by the prescriber prior to treatment with Accutane, and the drug should be discontinued and the etiology further investigated.

Inflammatory Bowel Disease
Accutane has 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 Accutane treatment has been stopped. Patients experiencing abdominal pain, rectal bleeding or severe diarrhea should discontinue Accutane immediately (see **ADVERSE REACTIONS: Gastrointestinal**).

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Visual Impairment
Decreased vision has been reported during Accutane 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 danger and warned to be cautious when driving or operating any vehicle at night.

PRECAUTIONS
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Wholesalers:
For the purpose of the iPLEDGE REMS, the term *wholesaler* refers to wholesaler, distributor, and/or chain pharmacy distributor. To distribute Accutane, wholesalers must be enrolled with iPLEDGE, and agree to meet iPLEDGE requirements for wholesaler distribution of isotretinoin products. Wholesalers must enroll with iPLEDGE by signing and returning the iPLEDGE wholesaler agreement that affirms they will comply with all iPLEDGE requirements for distribution of isotretinoin products. These include:

- Distributing only FDA approved isotretinoin product
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Return to normal vision has been reported in patients who experience tinnitus or hearing impairment should discontinue Accutane treatment and be referred for specialized care for further evaluation of the iPLEDGE REMS. Meeting the requirements for a patient who can become pregnant signifies that the patient:

ADVERSE REACTIONS: Special Senses
Hepatotoxicity
Clinical hepatitis considered to be possibly or probably related to Accutane therapy has been reported. Although, mild to moderate elevations of liver enzymes have been observed in approximately 12% of patients treated during clinical trials, some of which normalized with dosage reduction or continued use of Accutane at a lower dose. The first test is screening test is obtained by the prescriber prior to treatment with Accutane, and the drug should be discontinued and the etiology further investigated.

Inflammatory Bowel Disease
Accutane has 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 Accutane treatment has been stopped. Patients experiencing abdominal pain, rectal bleeding or severe diarrhea should discontinue Accutane immediately (see **ADVERSE REACTIONS: Gastrointestinal**).

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Visual Impairment
Decreased vision has been reported during Accutane 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 danger and warned to be cautious when driving or operating any vehicle at night.

PRECAUTIONS
Accutane must only be prescribed by prescribers who are enrolled and activated with the iPLEDGE REMS. Accutane must only be dispensed by a pharmacy enrolled and activated with iPLEDGE, and must only be dispensed to patients who are enrolled and meet all the requirements of iPLEDGE. Enrolled and activated pharmacies must receive Accutane only from wholesalers enrolled with iPLEDGE. iPLEDGE REMS requirements for wholesalers, prescribers, and pharmacies are described below.

Wholesalers:
For the purpose of the iPLEDGE REMS, the term *wholesaler* refers to wholesaler, distributor, and/or chain pharmacy distributor. To distribute Accutane, wholesalers must be enrolled with iPLEDGE, and agree to meet iPLEDGE requirements for wholesaler distribution of isotretinoin products. Wholesalers must enroll with iPLEDGE by signing and returning the iPLEDGE wholesaler agreement that affirms they will comply with all iPLEDGE requirements for distribution of isotretinoin products

