

Condylox® Gel - Package Insert

1 **Condylox® (podofilox gel) Gel, 0.5%**

2 (con' de lox)

3 **DESCRIPTION**

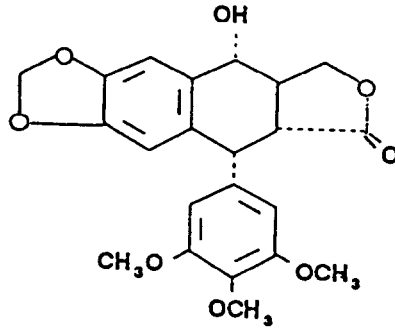
4 Podofilox is an antimitotic drug which can be chemically synthesized or purified from the
5 plant families *Coniferae* and *Berberidaceae* (e.g. species of *Juniperus* and *Podophyllum*).

6 Condylox® Gel 0.5% is formulated for topical administration. Each gram of gel contains
7 5mg of podofilox in a buffered alcoholic gel containing alcohol, glycerin, lactic acid,
8 hydroxypropyl cellulose, sodium lactate, and butylated hydroxytoluene.

9 Podofilox has a molecular weight of 414.4 daltons, and is soluble in alcohol* and sparingly
10 soluble in water. Its chemical name is [5R-(5 α ,5a β ,8a α ,9 α)]-tetrahydro-9-hydroxy-5-
11 (3,4,5-trimethoxyphenyl)furo[3',4':6,7]naphtho-[2,3-d]-1,3-dioxol-6(5aH)-one.

12 Podofilox has the following structural formula:

13



14 **CLINICAL PHARMACOLOGY**

15 **Mechanism of Action**

16 Treatment of anogenital warts with podofilox results in necrosis of visible wart tissue. The
17 exact mechanism of action is unknown.

18 **Pharmacokinetics**

19 In systemic absorption studies in 52 patients, topical application of 0.05mL of an ethanolic
20 solution containing 0.5% podofilox to external genitalia did not result in detectable serum
21 levels. Applications of 0.1 to 1.5mL resulted in peak serum levels of 1 to 17 ng/mL one
22 to two hours after application. The elimination half-life ranged from 1.0 to 4.5 hours. The
23 drug was not found to accumulate after multiple treatments.¹

24 **CLINICAL STUDIES**

25 In the first multicenter clinical study in 326 patients with anogenital warts, Condylor® Gel
26 0.5% and its vehicle were applied in a double-blind fashion to comparable patient groups.
27 Of the 260 patients with efficacy data, 176 were treated with Condylor® Gel 0.5%.
28 Patients applied Condylor® Gel 0.5% twice daily for three consecutive days followed by
29 a 4 day "rest" period.

30 At the end of 4 weeks, 38.4% of the patients had complete clearing of the wart tissue
31 when treated with Condylor® Gel 0.5%.

32 In the second multicenter clinical trial in 108 evaluable patients with anogenital warts,
33 Condylor® (podofilox solution) Solution 0.5% was compared with Condylor® Gel 0.5%
34 for efficacy. As in the first clinical trial, patients applied Condylor® Gel 0.5% twice daily
35 for three consecutive days followed by a four day "rest" period.

36 Similar clearance rates were observed. At the end of 4 weeks, 25.6% of the patients had
37 complete clearing of the wart tissue when treated with Condylor® Gel 0.5%.

38 INDICATIONS AND USAGE

39 Condylor® Gel 0.5% is indicated for the topical treatment of anogenital warts (external

40 genital warts and perianal warts). This product is *not* indicated in the treatment of mucous
41 membrane warts (see **PRECAUTIONS**).

42 **Diagnosis**

43 Although anogenital warts have a characteristic appearance, histopathologic confirmation
44 should be obtained if there is any doubt of the diagnosis. Differentiating warts from
45 squamous cell carcinoma and "Bowenoid papulosis" is of particular concern. Squamous
46 cell carcinoma may also be associated with human papillomavirus which should not be
47 treated with Condyllox® Gel 0.5%.

48 **CONTRAINDICATIONS**

49 Condyllox® Gel 0.5% is contraindicated for patients who develop hypersensitivity or
50 intolerance to any components of the formulation.

51 **WARNINGS**

52 Correct diagnosis of the lesions to be treated is essential. See the **Diagnosis** subsection of
53 the **INDICATIONS AND USAGE** section. Condyllox® Gel 0.5% is intended for
54 cutaneous use only. Avoid contact with the eyes. If contact with the eyes occurs,

55 **patients should immediately flush the eyes with copious quantities of water and seek**
56 **medical advice.**

57 **Drug Product is Flammable. Keep Away From Open Flame.**

58 **PRECAUTIONS**

59 **General**

60 **Data are not available on the safe and effective use of this product for treatment of warts**
61 **occurring on mucous membranes of the genital area (including the urethra, rectum and**
62 **vagina). The recommended method of application, frequency of application, and duration**
63 **of usage should not be exceeded (see DOSAGE AND ADMINISTRATION).**

64 **Information for Patients**

65 **Patients using Condylox® Gel 0.5% should receive the following information and**
66 **instructions. This information is intended to aid in the safe and effective use of this**
67 **medication. It is not intended to disclose all possible adverse or intended effects.**

68 1) **This medication should be used only as directed by the health care provider.**

- 69 Patients should be instructed to wash their hands thoroughly before and after each
70 application. It is for external use only. Avoid contact with the eyes.
- 71 2) Patients should be advised not to use this medication for any disorder
72 other than that for which it was prescribed.
- 73 3) Patients should report any signs of adverse reactions to the health care
74 provider.
- 75 4) If no improvement is observed after 4 weeks of treatment, discontinue
76 the medication and consult the health care provider.

77 **Carcinogenesis, Mutagenesis and Impairment of Fertility**

78 An 80-week carcinogenicity study in the mouse was performed using a 0.5% podofilox
79 solution applied dermally at 0.04, 0.2 and 1.0 mg/kg/day. There were no differences
80 between the podofilox treated mice at any dose level and vehicle control in the incidence
81 of neoplasia. Published animal studies, in general, have not shown the drug substance,
82 podofilox, to be carcinogenic.^{2,3,4,5,6} There are published reports that, in mouse studies,
83 crude podophyllin resin (containing podofilox) applied topically to the cervix produced
84 changes resembling carcinoma *in situ*.⁷ These changes were reversible at five weeks after
85 cessation of treatment. In one reported experiment, epidermal carcinoma of the vagina

86 and cervix was found in 1 out of 18 mice after 120 applications of podophyllin⁸ (the drug
87 was applied twice weekly over a 15-month period).

88 Podofilox was not mutagenic in the Ames plate reverse mutation assay at concentrations
89 up to 5mg/plate, with and without metabolic activation. No cell transformation related to
90 potential oncogenicity was observed in BALB/3T3 cells after exposure to podofilox at
91 concentrations up to 0.008 μ g/mL, without metabolic activation and 12 μ g/mL podofilox
92 with metabolic activation. Results from the mouse micronucleus *in vivo* assay using
93 podofilox 0.5% solution at doses up to 25 mg/kg (75 mg/m²), indicate that podofilox
94 should be considered a potential clastogen (a chemical that induces disruption and
95 breakage of chromosomes).

96 Daily topical application of 0.5% podofilox solution at doses up to the equivalent of
97 0.2mg/kg (1.18 mg/m², approximately equivalent to the human daily dose) to rats
98 throughout gametogenesis, mating, gestation, parturition and lactation for two generations
99 demonstrated no impairment of fertility.

100 **Pregnancy**

101 Pregnancy Category C: 0.5% podofilox solution was not teratogenic in the rabbit
102 following topical application of up to 0.21 mg/kg (2.85 mg/m², approximately 2 times the
103 maximum human dose) once daily for 13 days. The scientific literature contains references
104 that podofilox is embryotoxic in rats when administered intraperitoneally at a dose of
105 5mg/kg (29.5 mg/m², approximately 19 times the recommended maximum human dose).⁹
106 Teratogenicity and embryotoxicity have not been studied with intravaginal application.
107 Many antimitotic drug products are known to be embryotoxic. There are no adequate and
108 well-controlled studies in pregnant women. Condylox[®] Gel 0.5% should be used in
109 pregnancy only if the potential benefit justifies the potential risk to the fetus.

110 **Nursing Mothers**

111 It is not known whether this drug is excreted in human milk. Because of the potential for
112 serious adverse reactions in nursing infants from podofilox, a decision should be made
113 whether to discontinue nursing or to discontinue the drug, taking into account the
114 importance of the drug to the mother.

115 **Pediatric Use**

116 Safety and effectiveness in pediatric patients have not been established.

117 **ADVERSE REACTIONS**

118 In clinical trials with Condylox® Gel 0.5%, the following local adverse reactions were
119 reported during the treatment of anogenital warts. The severity of local adverse reactions
120 were predominantly mild or moderate and did not increase during the treatment period.
121 Severe reactions were most frequent within the first 2 weeks of treatment.

122	Adverse Reaction	Mild	Moderate	Severe
123	Inflammation	32.2%	30.4%	9.3%
124	Burning	37.1%	25.9%	11.5%
125	Erosion	27.0%	20.8%	8.9%
126	Pain	23.7%	20.4%	11.5%
127	Itching	32.2%	16.0%	7.8%
128	Bleeding	19.2%	3.0%	0.7%

129 Other local adverse reactions reported included stinging (7%), and erythema (5%); less
130 commonly reported local adverse events included desquamation, scabbing, discoloration,
131 tenderness, dryness, crusting, fissures, soreness, ulceration, swelling/edema, tingling,

132 rash, and blisters.

133 The most common systemic adverse event reported during the clinical studies was
134 headache (7%).

135 **OVERDOSAGE**

136 Topically applied podofilox may be absorbed systemically (see **CLINICAL**
137 **PHARMACOLOGY** section). Toxicity reported following systemic administration of
138 podofilox in investigational use for cancer treatment included: nausea, vomiting, fever,
139 diarrhea, bone marrow depression, and oral ulcers. Following 5 to 10 daily intravenous
140 doses of 0.5 to 1 mg/kg/day, significant hematological toxicity occurred but was
141 reversible.¹⁰ Other toxicities occurred at lower doses. Toxicity reported following systemic
142 administration of podophyllum resin included: nausea, vomiting, fever, diarrhea,
143 peripheral neuropathy, altered mental status, lethargy, coma, tachypnea, respiratory
144 failure, leukocytosis, pancytosis, hematuria, renal failure and seizures.¹¹ Treatment of
145 topical overdosage should include washing the skin free of any remaining drug and
146 symptomatic and supportive therapy.

147 **DOSAGE AND ADMINISTRATION**

148 The prescriber should ensure that the patient is fully aware of the correct method of
149 therapy and identify which specific warts should be treated.

150 Apply twice daily for 3 consecutive days, then discontinue for 4 consecutive days. This
151 one week cycle of treatment may be repeated until there is no visible wart tissue or for a
152 maximum of four cycles. If there is incomplete response after four treatment cycles,
153 **discontinue treatment and consider alternative treatment. Safety and effectiveness**
154 **of more than four treatment cycles has not been established.**

155 There is no evidence to suggest that more frequent application will increase efficacy, but
156 additional applications would be expected to increase the rate of local adverse reactions
157 and systemic absorption.

158 Condylox® Gel 0.5% should be applied to the warts with the applicator tip or
159 finger. Application on the surrounding normal tissue should be minimized. Treatment
160 should be limited to 10 cm² or less of wart tissue and to no more than 0.5g of the gel
161 per day.

162 Care should be taken to allow the gel to dry before allowing the return of opposing skin
163 surfaces to their normal positions. Patients should be instructed to wash their hands
164 thoroughly before and after each application.

165 **HOW SUPPLIED**

166 Condylor® Gel 0.5% is supplied as 3.5g of clear gel in aluminum tubes with an applicator
167 tip. NDC 55515-102-01. Store at controlled room temperature between 15°-30°C (59°-
168 86°F). Avoid excessive heat. Do not freeze.

169 Caution: Federal law prohibits dispensing without prescription.

170 **REFERENCES**

- 171 1. von Krogh G. Podophyllotoxin in serum: Absorption subsequent to three day
172 repeated applications of a 0.5% ethanolic preparation on condylomata acuminata.
173 Sex Trans Disease 1982; 9: 26-33.
- 174 2. Berenblum I. The effect of podophyllotoxin on the skin of the mouse, with
175 reference to carcinogenic, cocarcinogenic, and anticarcinogenic action. J Cancer
176 Inst 11:839-841, 1951.
- 177 3. Kaminetzky HA, Swerdlow M. Podophyllin and the mouse cervix:assessment of

- 178 carcinogenic potential. Am J Obst Gyn 95:486-490, 1965.
- 179 4. McGrew EA, Kaminetzky HA. The genesis of experimental cervical epithelial
180 dysplasia. Am J Clin Path 35:538-545, 1961.
- 181 5. Roe FJC, Salaman MH. Further studies on incomplete carcinogenesis: triethylene
182 melamine (T.E.M.) 1,2 benzanthracene and beta-propiolactone as initiators of skin
183 tumor formation in the mouse. Brit J Cancer 9:177-203, 1955.
- 184 6. Taper HS. Induction of the deficient acid DNAase activity in mouse interfollicular
185 epidermis by croton oil as a possible tumor promoting mechanism. Zeitschrift fur
186 Krebsforschung and Klinisch Onkologie (Cancer Research and Clinical Oncology,
187 Berlin) 90:197-210, 1977.
- 188 7. Kaminetzky HA, McGrew EA, Phillips RL. Experimental cervical epithelial
189 dysplasia. J Obst Gyn 14:1-10,1959.
- 190 8. Kaminetzky HA, McGrew EA: Podophyllin and mouse cervix: Effect of long
191 term application. Arch Path 73:481-485, 1962.
- 192 9. Thiersch JB. Effect of podophyllin (P) and podophylotoxine (PT.) on the rat litter
193 in utero. Soc Exptl Biol Med Proc 113:124-127, 1963.
- 194 10. Savel H.: Clinical experience with intravenous podophyllotoxin. Proc Amer

195 Assoc Cancer Res, 1964; 5: 56.

196 11. Cassidy DE, Dewry J and Fanning JP: Podophyllum toxicity: A report of a fatal
197 case and a review of the literature. J Toxicol Clin Toxicol 1982; 19: 35-44.

198 Mfd. for

199 Oclassen

200 PHARMACEUTICALS, INC.

201 San Rafael, CA 94901

202 by DPT Laboratories, Inc.

203 San Antonio, TX 78215

204 Revised March 13, 1997

MAR 13 1997

Condylox® Gel 0.5%

(podofilox gel)

Condylox® Gel (podofilox gel) and Anogenital Warts

Patient Information

FPO

1. APPLY CONDYLOX® GEL ONLY ON THE WARTS POINTED OUT BY YOUR DOCTOR.
2. YOU MAY FEEL SOME MILD TO MODERATE DISCOMFORT DURING TREATMENT.
3. STOP TREATMENT AND CALL YOUR DOCTOR IF YOU HAVE BLEEDING, SWELLING, OR EXCESSIVE PAIN, BURNING, OR ITCHING.
4. DO NOT USE MORE THAN TWO TIMES A DAY.
5. DO NOT USE FOR MORE THAN THREE DAYS IN A ROW.
6. DO NOT HAVE SEXUAL INTERCOURSE ON THE DAYS YOU ARE APPLYING CONDYLOX® GEL.
7. WASH HANDS AFTER EVERY USE.

INTRODUCTION

Condylox® Gel slowly kills external anogenital warts. The warts will change from a fleshy skin color to a dry, crusted, dead look, then disappear. Three out of four patients feel some burning or pain after they apply Condylox® Gel. Other side effects may include redness, soreness, tenderness, and small sores. These usually go away within a week after Condylox® Gel is stopped. If pain or other side effects bother you too much, stop applying Condylox® Gel and contact your doctor.

HOW TO USE CONDYLOX® GEL

Follow these and your doctor's instructions carefully. Apply Condylox® Gel only on the warts pointed out by your doctor. Do not use it on any other warts on or inside your body, or for any other skin growth.

1. Unscrew the entire applicator cap. Invert the cap and puncture the tube seal. Replace the applicator cap. To apply Condylox® Gel, remove the protective cap on the applicator tip and apply to the warts using the applicator tip or finger. Make sure to replace the applicator cap tightly after use.

APPLY CONDYLOX® GEL ONLY WHERE YOUR DOCTOR HAS INSTRUCTED YOU.

2. Apply a small amount of Condylox® Gel to the wart(s). Do not get it on normal skin. If a wart is in a skin fold, spread the skin apart so you can reach the wart. A hand mirror can help sometimes. Let Condylox® Gel dry before letting the skin folds return to their normal position. Wash your hands well with soap and water after you use Condylox® Gel.
3. Apply Condylox® Gel once in the morning and once in the evening for three days in a row. Then stop applying Condylox® Gel and wait four days. Using Condylox® Gel like this is called a treatment

week. You should not wash Condyllox® Gel off the wart area unless you experience excessive pain, burning, or itching.

DO NOT APPLY CONDYLOX® GEL MORE THAN TWICE EACH DAY OR FOR MORE THAN THREE DAYS IN A ROW. USING CONDYLOX® GEL MORE OFTEN WILL NOT MAKE IT WORK BETTER BUT MAY INCREASE SIDE EFFECTS.

4. If the warts do not go away, repeat the Condyllox® Gel treatment for another week. You can use Condyllox® Gel up to four treatment weeks (REMEMBER: a treatment week is twice a day for three days, then four days with no treatment). Your doctor may ask you to come back for a check-up visit during treatment. If the warts have not gone away after four treatment weeks, stop applying Condyllox® Gel and contact your doctor.

IF THE AREA YOU ARE PUTTING CONDYLOX® GEL ON IS BLEEDING OR SWOLLEN, OR IF THERE IS EXCESSIVE PAIN, BURNING OR ITCHING, STOP APPLYING CONDYLOX® GEL AND CONTACT YOUR DOCTOR.

5. Anogenital warts can come back. If your warts come back, contact your doctor.

SPECIAL CAUTIONS

- Anogenital warts are contagious. You can give them to or get them from your sexual partner. Make sure your sexual partner has been checked for anogenital warts. Condoms may help prevent giving anogenital warts to your sexual partner. Do not have sexual intercourse for the three days you are applying Condyllox® Gel.
- Women should make sure to use birth control so they will not get pregnant while on Condyllox® Gel. The effects on the unborn baby are not known. Women can use Condyllox® Gel during their menstrual period.
- Condyllox® Gel is prescribed only for your external anogenital warts. Do not let anyone else use it.
- Drug Product is Flammable. Keep Away From Open Flame.

REMEMBER

- Always wash your hands after using Condyllox® Gel.
- Do not get it in your eyes. If you do, immediately flush your eyes with water and contact your doctor.
- Keep the tube cap tightly closed.
- Be sure to keep this and all medications out of the reach of children.

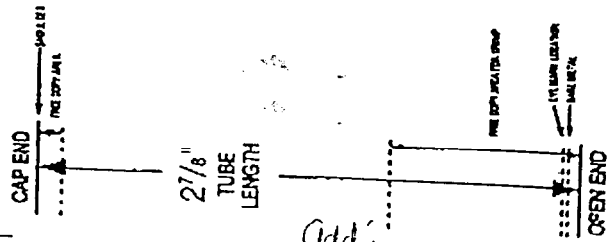
CONTACT YOUR DOCTOR IF YOU HAVE QUESTIONS ABOUT CONDYLOX® GEL.

Mfd. for
Oclassen
PHARMACEUTICALS, INC.
San Rafael, CA 94901
by DPT Laboratories, Inc.
San Antonio, TX 78215

126862-1096

Revised October 31, 1986.

TUBE LABEL (ACTUAL SIZE)



For topical use only. Not for use in the eyes.
 Keep out of the reach of children.
 Usual Dosage: See enclosed literature.
 Keep tightly closed.
 See crimp for expiration date and lot no.
 Mfd. for Occlusen Pharmaceuticals, Inc.
 San Rafael, CA 94901 by DPT Laboratories, Inc.
 San Antonio, TX 78215

Add:
 Drug Product is flammable.
 Replenish from open flame.

Condylax[®] Gel 0.5%
 (podofilox gel)

*- delete line
 add line*

Caution: Federal (FDA) for products
 dispensing without a prescription.

Phillips
 3-3-97

CARTON LABELING

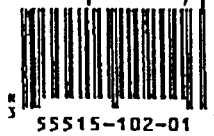
(ACTUAL SIZE)

Acid: Drug product is Flammable. Keep away from open flame.

WARNING: FOR TOPICAL USE ONLY.
NOT FOR USE IN THE EYES.
KEEP OUT OF THE REACH OF CHILDREN.

Oclassen
Pharmaceutical, Inc.
San Antonio, TX 78209

See end of carton for
expiration date and lot number.



B
74

3.5 g NDC 55515-102-01

Condylox[®] Gel 0.5%
(podofilox gel)

Oclassen **FOR TOPICAL USE ONLY** Caution: Federal (U.S.A.) law prohibits dispensing without a prescription.



3.5 g

Condylox Gel 0.5%
PODOFILOX GEL



LOT 88

Each g contains 5 mg podofilox in a buffered alcoholic gel containing alcohol, glycerin, lactic acid, hydroxypropyl cellulose, sodium stearate, and butylated hydroxytoluene.

Instructions For Use: See accompanying patient information for precautions and complete instructions for use.

Invert cap to puncture seal in tube. After each use recap the tube tightly. Always wash hands after use.

Store at controlled room temperature 15° to 30°C (59° to 86°F). Avoid excessive heat. Do not freeze.

Alcohol 78%

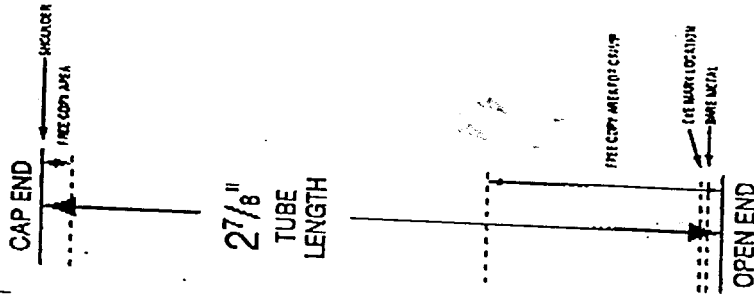
110822-0396

IN

R. Davis

3-3-97

TUBE LABEL
(125% X ACTUAL)



For topical use only. Not for use in the eyes.
Keep out of the reach of children.

Usual Dosage: See enclosed literature.

Keep tightly closed.

See crimp for expiration date and lot no.

Mfd. for Oclassen Pharmaceuticals, Inc.
San Rafael, CA 94901 by DPT Laboratories, Inc.
San Antonio, TX 78215

3.6 g

103758 - 0236
NDC 65515-182-01

Condylox[®] Gel 0.5%
(podofilox gel)

Oclassen
www.oclassen.com

Caution: Federal (N.S.A.) law prohibits
dispensing without a prescription.

Drug Product is Flammable. Keep away from open flame.

delete line
add line

R. Davis
3-3-97

CARTON LABELING

(125% x ACTUAL SIZE)

Add: Drug product is flammable.
Keep away from open flame.

WARNING: FOR TOPICAL USE ONLY.
NOT FOR USE IN THE EYES.
KEEP OUT OF THE REACH OF CHILDREN.

Oclassen
PHARMACEUTICALS, INC.
San Rafael, CA 94901
by DPT Laboratories, Inc.
San Antonio, TX 78216

See end of carton for
expiration date and lot number.



74
NDC 000

3.5 g NDC 55515-102-01

Condylox® Gel 0.5%
(podofilox gel)

Oclassen FOR TOPICAL USE ONLY Caution: Federal (U.S.A.) law prohibits dispensing without a prescription.
PHARMACEUTICALS, INC.



3.5 g

Condylox Gel 0.5%
NDC 55515-102-01

Each g contains 5 mg podofilox ^{ik} as a buffered alcoholic gel containing alcohol, glycerin, lactic acid, hydroxypropyl cellulose, sodium lactate, and butylated hydroxytoluene.

Instructions For Use: See accompanying patient information for precautions and complete instructions for use.

Invert cap to puncture seal in tube. After each use recap the tube tightly. Always wash hands after use.

Store at controlled room temperature 15° to 30°C (59° to 86°F). Avoid excessive heat. Do not freeze.
Alcohol 79% 110622-0396

LOT

EXP



R. D. Miller
3-3-97