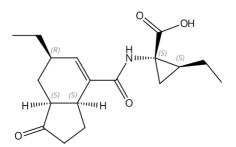
# CORONATINE

# XXX



Common name	Coronatine
Chemical name	(1S,2S)-2-Ethyl-1-[[[(3aS)-6α-ethyl-2,3,3aβ,6,7,7aβ- hexahydro-1-oxo-1H-inden-4- yl]carbonyl]amino]cyclopropanecarboxylic acid
CAS No.	62251-96-1
Empirical formula	$C_{18}H_{25}NO_4$
RMM	319.4
<i>m.p</i> .	163.2 °C~165.9 °C
<i>v.p</i> .	2.45×10 <sup>-7</sup> Pa at 20 °C 5.84×10 <sup>-7</sup> Pa at 25 °C
Solubility	In water 1.177 g/l; in methanol >250 g/l; acetone 160-200 g/l; ethyl acetate 50-57g/l; n-hexane <48.74 mg/l; toluene 294.87 mg/l; 1,2-dichloroethane 50-57g/l (all at 20 $\$ )
Stability	Stable in acidic and neutral conditions, stable at room temperature
Description	White powder
Formulations	Soluble concentrate

## **CORONATINE TECHNICAL**

## XXX/TC/M/-

**1. Sampling.** Take at least 100 g.

#### 2. Identity tests

**2.1 HPLC.** Use the HPLC method below. The relative retention time of the coronatine peak in the sample solution should not deviate by more than 1.5% from that of the calibration solution.

**2.2 Infrared.** Prepare potassium bromide discs for the technical sample and coronatine reference substance. Scan the discs from 4000 to 400 cm<sup>-1</sup>. The spectrum from the sample should not differ significantly from that of the reference substance.

### **3.** Coronatine

## **OUTLINE OF METHOD**

Coronatine is determined by high performance liquid chromatography on a reversed phase column (C18) with UV detection at 220 nm and external standardization.

## REAGENTS

Coronatine reference standard of known purity.

Methanol HPLC grade.

Acetonitrile HPLC grade.

*Water* ultrapure quality or distilled in glass.

Phosphoric acid analytical grade, minimum 85%.

0.1% Phosphoric acid aqueous solution Dissolve 1 ml phosphoric acid into 1000 ml water.

**Calibration solution**. Weigh in duplicate (to the nearest 0.02 mg) 19 mg of coronatine reference standard (*s* mg) into separate volumetric flasks (50 ml). Dissolve with 30 ml mobile phase and place the flask in an ultrasonic bath for 3 minutes. Allow to cool to ambient temperature and fill to the mark with mobile phase. Mix thoroughly. Filter the solution through a 0.45  $\mu$ m filter before injection (calibration solutions C<sub>A</sub> and C<sub>B</sub>).

## **APPARATUS**

*High performance liquid chromatograph* equipped with a UV detector capable for operation at 220 nm, a constant-temperature column compartment and an injection system capable of injecting 5µl.

Column stainless steel 250 ×4.6 mm (i.d), packed with  $C_{18}$ , 5.0 µm, or equivalent with the same selectivity.

Filtering apparatus disposable plastic syringes (or equivalent) fitted with  $0.45 \ \mu m$  filters

Electronic integrator or data system

Ultrasonic bath

#### PROCEDURE

#### (a) Liquid chromatographic conditions (typical):

Column	stainless steel, 250 ×4.6 mm (i.d.), packed with $C_{18}$ , 5.0 µm, or equivalent with the same selectivity
Mobile phase	acetonitrile: 0.1% phosphoric acid aqueous solution, 30:70 (v/v)
Flow rate	1.0 ml/min
Column temperature	35°C
Detector wavelength	220 nm
Injection volume	5 µl
Retention time	approximately 21.5 min
Run time	28 min

(b) System equilibration. Inject 5  $\mu$ l portions of calibration solution C<sub>A</sub> until the response factors (*fi*) obtained for two consecutive injections differ by less than 1.5%. Then inject 5  $\mu$ l portions of calibration solution C<sub>B</sub>. The response factor (*fi*), for two consecutive injections should not deviate by more than 1.5% from that of solution C<sub>A</sub>, otherwise prepare new calibration solutions.

(c) Sample preparation. Prepare solutions in duplicate for each sample.

Weigh (to the nearest 0.02 mg) sufficient sample (w mg) to contain about 19 mg of coronatine into a volumetric flask (50 ml). Dissolve with 30 ml mobile phase and place the flask in an ultrasonic bath for 3 minutes. Allow to cool to ambient temperature and fill to the mark with mobile phase. Mix thoroughly. Filter the solution through a 0.45 µm filter before injection (sample solutions S<sub>1</sub> and S<sub>2</sub>).

(d) **Determination.** Inject in duplicate 5  $\mu$ l portions of each sample solution bracketing them by injections of the calibration solutions as follows:

 $C_A, S_1, S_1, C_B, S_2, S_2, C_{A, \dots}$ 

(e) Calculation. Calculate the mean value of each pair of calibration response factors *f*, bracketing the two injections of a sample, and use this value for calculating the coronatine contents of the bracketed sample injections.

$$f_{i} = \frac{S \times P}{H_{s}}$$
  
Coronatine content (g/kg) =  $\frac{H_{w} \times f}{W}$ 

where:

 $f_i$  = individual response factor

f = mean response factor

 $H_s$  = peak area of coronatine in the calibration solution

 $H_w$  = peak area of coronatine in the sample solution

S = mass of coronatine reference standard in the calibration solution (mg)

W = mass of sample taken (mg)

P = purity of coronatine reference standard (g/kg)

**Repeatability**  $\mathbf{r} = g/kg$  at an active ingredient content of g/kg**Reproducibility**  $\mathbf{R} = g/kg$  at an active ingredient content of g/kg

#### CORONATINE SOLUBLE CONCENTRATE

#### XXX/SL/M/-

**1. Sampling.** Take at least 200 ml.

## 2. Identity tests.

# **2.1 HPLC.** As for coronatine technical XXX/TC/M/2.1

**3. Coronatine.** As for coronatine technical XXX/TC/M/3 except:

## REAGENTS

**Calibration solution.** Weigh in duplicate (to the nearest 0.02 mg) 19 mg of coronatine reference standard (*s* mg) into separate volumetric flasks (25ml). Dissolve with 5 ml methanol and place the flask in an ultrasonic bath until the sample has been dissolved completely. Allow to cool to ambient temperature and fill to the mark with methanol and mix thoroughly as stock solution. Then transfer by pipette 2.0 ml of each stock solution into separate 25 ml volumetric flask, dilute to volume with mobile phase and mix thoroughly. Filter the solution through a 0.45 µm filter before injection (calibration solutions  $C_A$  and  $C_B$ ).

# PROCEDURE

## (a) Liquid chromatographic conditions (typical):

*Injection volume* 20 µl

(c) Sample preparation. Prepare solutions in duplicate for each sample. Add to the mark of each 25 ml volumetric flask with coronatine sample, accurately weigh the sample mass (to the nearest 0.2 mg). Filter the solution through a 0.45  $\mu$ m filter before injection (solutions S<sub>1</sub> and S<sub>2</sub>).

(e) Calculation. Calculate the mean value of each pair of calibration response factors f, bracketing the two injections of a sample, and use this value for calculating the coronatine contents of the bracketed sample injections.

$$f_{i} = \frac{S \times P}{H_{s}}$$
  
Coronatine content  $(g/kg) = \frac{H_{w} \times f}{W \times 12.5}$ 

where:

 $f_i$  = individual response factor

f = mean response factor

 $H_s$  = peak area of coronatine in the calibration solution

 $H_w$  = peak area of coronatine in the sample solution

S = mass of coronatine reference standard in the calibration solution (mg)

W = mass of sample taken (mg)

P = purity of coronatine reference standard (g/kg) 12.5 = dilution ratio of the standard relative to the sample

**Repeatability r** = g/kg at an active ingredient content of g/kg**Reproducibility R** = g/kg at an active ingredient content of g/kg

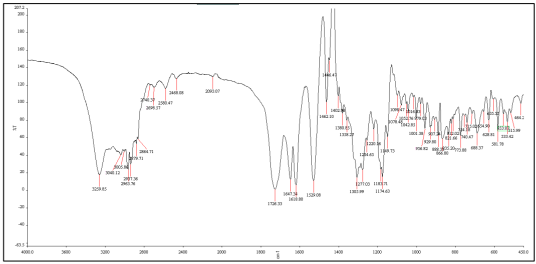


Fig. 1 FTIR spectrum of coronatine

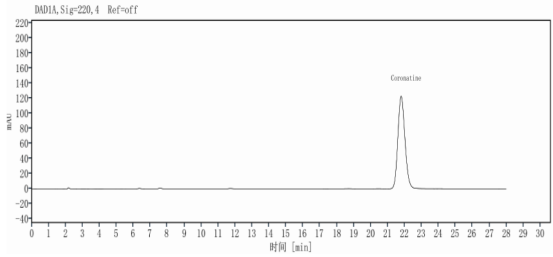


Fig. 2 HPLC Chromatogram of coronatine standard

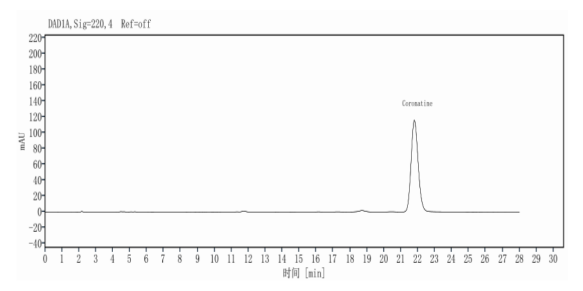


Fig. 3 HPLC Chromatogram of coronatine TC

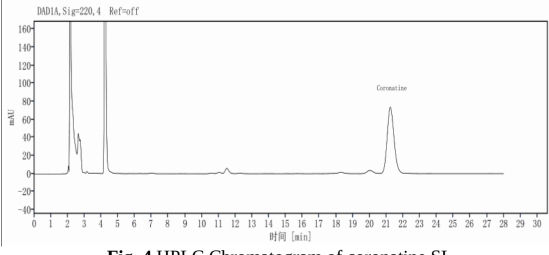


Fig. 4 HPLC Chromatogram of coronatine SL