

## EXHIBIT B



US005610163A

**United States Patent** [19][11] **Patent Number:** **5,610,163****Banholzer et al.**[45] **Date of Patent:** **Mar. 11, 1997**[54] **ESTERS OF THIENYL CARBOXYLIC ACIDS AND AMINE ALCOHOLS AND THEIR QUATERNIZATION PRODUCTS**[75] **Inventors:** **Rolf Banholzer**, Ingelheim am Rhein; **Rudolf Bauer**, Wiesbaden; **Richard Reichl**, Ingelheim am Rhein, all of Germany[73] **Assignee:** **Boehringer Ingelheim GmbH**, Ingelheim am Rhein, Germany[21] **Appl. No.:** **405,111**[22] **Filed:** **Mar. 16, 1995****Related U.S. Application Data**

[63] Continuation of Ser. No. 254,324, Jun. 6, 1994, abandoned, which is a continuation of Ser. No. 100,822, Aug. 2, 1993, abandoned, which is a continuation of Ser. No. 838,724, Mar. 13, 1992, abandoned.

[30] **Foreign Application Priority Data**

Sep. 16, 1989 [DE] Germany ..... 39 31 041.8

[51] **Int. Cl.<sup>6</sup>** ..... A61K 31/435; C07D 401/00; C07D 451/12[52] **U.S. Cl.** ..... 514/291; 514/304; 546/18; 546/91; 546/125[58] **Field of Search** ..... 546/91, 125; 514/291, 514/304[56] **References Cited****U.S. PATENT DOCUMENTS**

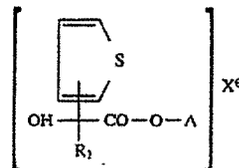
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**OTHER PUBLICATIONS**

The Merck Index, 11th ed (1989), Merck and Co, Inc., pp. 242 and 802-803.

*Primary Examiner*—Philip I. Datlow  
*Attorney, Agent, or Firm*—Robert P. Raymond; Alan R. Stempel; Mary-Ellen M. Devlin[57] **ABSTRACT**

Compounds of the formula



of which, in exemplary compounds, the thienyl group is attached via the 2-position and:

(a) A is 3 $\alpha$ -(6 $\beta$ , 7 $\beta$ -epoxy)-tropanyl methobromide and R<sub>1</sub> is 2-thienyl;(b) A is 3 $\alpha$ -(6, 7dehydro)-tropanyl methobromide and R<sub>1</sub> is 2-thienyl;(c) A is 3 $\beta$ -tropanyl methobromide and R<sub>1</sub> is 2-thienyl; and,(d) A is 3 $\alpha$ -(N-isopropyl)-nortropanyl methobromide and R<sub>1</sub> is cyclopentyl.

These are anticholinergics. Administered by inhalation, they are useful for the treatment of chronic obstructive bronchitis or slight to moderately severe asthma. Administered by the intravenous or oral routes, they are useful for the treatment of vagally induced sinus bradycardia.

**16 Claims, No Drawings**

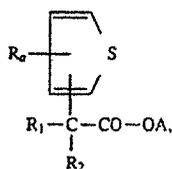
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**ESTERS OF THIENYL CARBOXYLIC ACIDS  
AND AMINO ALCOHOLS AND THEIR  
QUATERNIZATION PRODUCTS**

This is a continuation of application Ser. No. 08/254,324, filed on Jun. 6, 1994, now abandoned which is a continuation of application Ser. No. 08/100,822, filed on Aug. 2, 1993, now abandoned, which is a continuation of application Ser. No. 07/838,724, filed on Mar. 13, 1992, now abandoned.

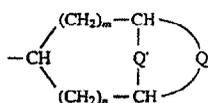
The invention relates to novel thienylcarboxylates of amino alcohols and their quaternary products and to the preparation of the novel compounds and their use as active ingredients in medicaments.

The novel compounds correspond to the formula



in which

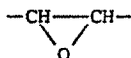
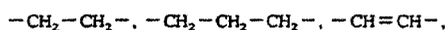
A represents the group



wherein

m and n independently of one another denote 1 or 2,

Q represents one of the double-bonding groups



and

Q' represents the group =NR or the group =NRR', wherein

R denotes H or an optionally halogen-substituted or hydroxy-substituted C<sub>1</sub>-C<sub>4</sub>-alkyl radical, R' denotes a C<sub>1</sub>-C<sub>4</sub>-alkyl radical and R and R' together may also form a C<sub>4</sub>-C<sub>6</sub>-alkylene radical, and wherein, in the case of quaternary compounds, one equivalent of an anion (X<sup>-</sup>) opposes the positive charge of the N atom,

R<sub>1</sub> represents a thienyl, phenyl, furyl, cyclopentyl or cyclohexyl radical, wherein these radicals may also be methyl-substituted, thienyl and phenyl may also be fluoro-substituted or chloro-substituted,

R<sub>2</sub> represents hydrogen, OH, C<sub>1</sub>-C<sub>4</sub>-alkoxy or C<sub>1</sub>-C<sub>4</sub>-alkyl,

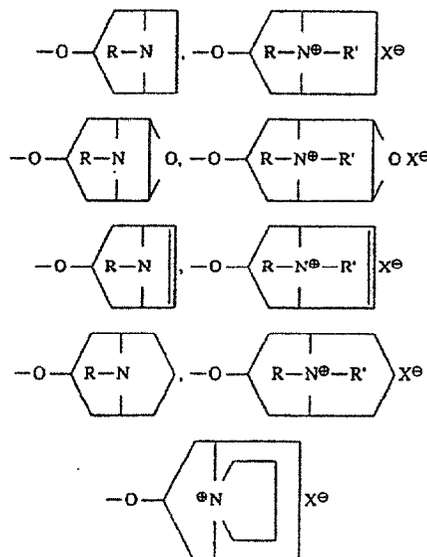
R<sub>a</sub> represents H, F, Cl or CH<sub>3</sub> and, if =NR denotes a secondary or tertiary amino group, also the acid addition salts.

In the compounds of formula I, R<sub>1</sub> preferably represents thienyl, R<sub>2</sub> preferably represents OH. The group -OA preferably has the α-configuration and is derived from, for example scopine, tropine, granatoline or 6,7-dehydrotropine

2

or the corresponding nor-compounds; however, -OA may also have the β-configuration, as in pseudotropine, pseudoscopine.

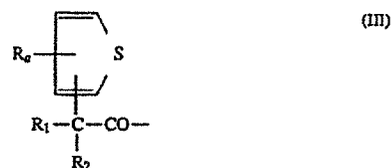
Corresponding radicals are, for example



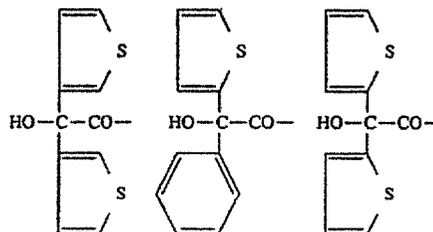
The substituent R is preferably a lower alkyl radical, such as CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub>, n-C<sub>3</sub>H<sub>7</sub>, i-C<sub>3</sub>H<sub>7</sub>, R' is preferably CH<sub>3</sub>. R and R' together are, for example -(CH<sub>2</sub>)<sub>5</sub>-. As halogen substituents for R, F or, as second choice, Cl are suitable.

If R denotes a halogen-substituted or hydroxy-substituted alkyl radical, it is preferably -CH<sub>2</sub>-CH<sub>2</sub>F or -CH<sub>2</sub>-CH<sub>2</sub>OH. Accordingly, the group A represents, for example the radicals of scopine, N-ethylnorscopine, N-isopropyl-norscopine, tropine, N-isopropyl-nortropine, 6,7-dehydrotropine, N-β-fluoroethylnortropine, N-isopropyl-6,7-dehydronortropine, N-methylgranatoline or the corresponding quaternary compounds, wherein the anion is preferably Br<sup>-</sup> or CH<sub>3</sub>SO<sub>3</sub><sup>-</sup>.

As the acid radical

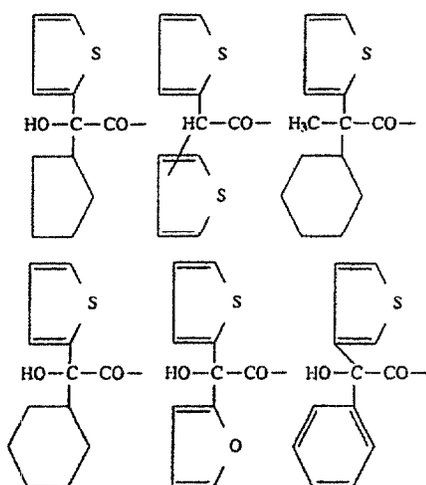


the following are particularly suitable:



3

-continued



The quaternary compounds are particularly suitable for therapeutic application, whereas the tertiary compounds are important not only as active ingredients but also as intermediate products.

The compounds of the invention are strong anti-cholinergic agents and have prolonged action. Action lasting at least 24 hours is achieved at inhaled dosages in the  $\mu\text{g}$  range. In addition, the toxicity is in the same range as the commercial product Ipratropium bromide, while at the same time the therapeutic effect is stronger.

The novel compounds are suitable, in accordance with their anti-cholinergic nature, for example for the treatment of chronic obstructive bronchitis and (slight to moderately severe) asthma, also for the treatment of vagally induced sinus bradycardia.

Whereas application of the novel active ingredients (in particular the quaternary compounds) by inhalation is mainly recommended for respiratory tract diseases, as a result of which side-effects are largely eliminated, the application for sinus bradycardia is preferably carried out intravenously or orally. It has thus proved to be advantageous

4

that the novel compounds leave the gastro/intestinal motility largely unaffected.

For administration the compounds of the invention are processed using known auxiliaries and/or excipients to give conventional galenic preparations, for example inhalation solutions, suspensions in liquified propellants, preparations containing liposomes or proliposomes, injection solutions, tablets, coated tablets, capsules, inhalation powders for use in conventional inhalation apparatus.

Formulation examples (measures in weight per cent):

#### 1. Controlled dosage aerosol

Active ingredient according to the invention	0.005
Sorbitan trioleate	0.1
monofluorotrichloromethane and difluorodichloromethane 2:3	to 100

The suspension is poured into a conventional aerosol container with a dosage valve. 50  $\mu\text{l}$  of suspension are preferably dispensed per actuation. The active ingredient may also be metered in a higher amount if required (for example 0.02 wt. %).

#### 2. Tablets

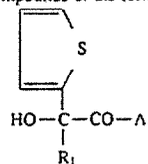
Active ingredient according to the invention	0.05
Colloidal silicic acid	0.95
Lactose	65.00
Potato starch	28.00
Polyvinylpyrrolidone	3.00
Na cellulose glycolate	2.00
Magnesium stearate	1.00

The constituents are processed in conventional manner to give tablets of 200 mg.

The advantageous properties of the novel compounds are shown, for example, in the inhibition of broncholysis in the rabbit (acetylcholine spasms intravenously). After intravenous administration of the novel active ingredients (dosage 3  $\mu\text{g}/\text{kg}$  intravenously), the maximum effect occurred after 10 to 40 minutes. After 5 hours the inhibiting effect had still not been reduced to half, that is to say the half effect time is more, in some cases considerably more, than 5 hours, as made clear by the residual effects after 5 hours listed below:

Compound	Residual effect in %
A	76
B	76
C	81
D	61
E	68
F	73
G	69

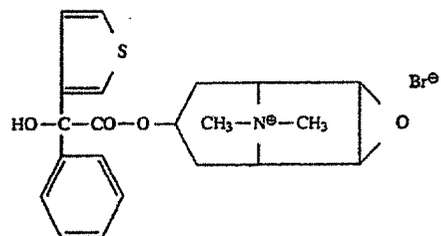
Compounds of the formula



Compound A

Compound A	$\text{Br}^\ominus$	$\text{R}_1$
A		2-thienyl
B		3-thienyl
D		2-thienyl
E		3-thienyl
F		cyclopentyl
G		cyclopentyl

Compound C

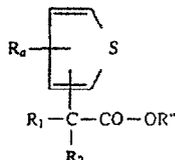


## Notes:

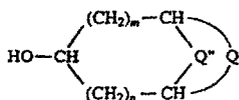
1. The compounds in which  $\text{R}_1$  is not 2-thienyl are racemates.
2. The compounds are  $3\alpha$ -compounds in each case.

Processes known per se are used to prepare the novel compounds.

An ester of the formula



wherein R'' represents a C<sub>1</sub>-C<sub>4</sub>-alkyl radical, preferably a methyl or ethyl radical (R<sub>1</sub>, R<sub>2</sub> and R<sub>4</sub> have the above meanings), is preferably transesterified using an amino alcohol of the formula



wherein m, n and Q have the above meanings, Q'' represents =NR or =NH and the OH group is in the α- or β-position, in the presence of a conventional transesterification catalyst, and the compound obtained is optionally quaternised

a) if Q'' denotes =NR (R ≠ H), using a reactive monofunctionalised derivative Z-(C<sub>1</sub>-C<sub>4</sub>-alkyl) of a corresponding alkane (Z=leaving group) or is optionally quaternised

b) if Q'' denotes =NH, using a terminally disubstituted alkane Z-(C<sub>4</sub>-C<sub>6</sub>-alkylene)-Z without isolation of intermediates.

The transesterification is carried out with heat in an organic solvent, for example toluene, xylene, heptane, or in a melt, strong bases such as sodium methylate, sodium ethylate, sodium hydride, metallic sodium, being used as catalyst. Reduced pressure is used to remove the released lower alcohol from the equilibrium, the alcohol is optionally distilled off azeotropically. The transesterification takes place at temperatures which in general do not exceed 95° C. Transesterification often proceeds more favourably in a melt. If required, the free bases may be obtained in a manner known per se from acid addition salts of the tertiary amines using suitable basic compounds. Quaternisation is carried out in suitable solvents, for example acetonitrile or acetonitrile/methylene chloride, preferably at room temperature; a corresponding alkyl halide, for example alkyl bromide, is preferably used in the process as quaternising agent. Transesterification products wherein Q' represents NH are used as starting materials for those compounds in which R and R' together represent a C<sub>4</sub>-C<sub>6</sub>-alkylene group. Conversion into the tertiary and then quaternary compound then takes place with the aid of suitable 1,4-dihaloalkanes, 1,5-dihaloalkanes or 1,6-dihaloalkanes without isolation of intermediates.

The starting materials may be obtained analogously to known compounds—in as much as they have not already been described.

#### EXAMPLES

methyl di-(2-thienyl)glycolate from dimethyl oxalate and 2-thienyl magnesium bromide;  
ethyl di-(2-thienyl)glycolate from (2-thienyl)glyoxylic acid and 2-thienyl lithium;  
ethyl hydroxy-phenyl-(2-thienyl)acetate from methyl phenylglyoxylate and 2-thienyl magnesium bromide or from methyl (2-thienyl)glyoxylate and phenyl magnesium bromide.

Methyl 2-thienylglyoxylate and cyclohexyl or cyclopentyl magnesium bromide may be reacted in a similar manner.

Several processes are also available for the preparation of the amino alcohols.

(IV) 5 Pseudoscopine may be obtained in accordance with M. Polonovski et al., Bull. soc. chim. 43, 79 (1928). Pseudotrophenol may be removed from the mixture (fractional crystallisation or distillation) which is obtained, for example in accordance with V. Hayakawa et al., J. Amer. Chem. Soc. 1978, 100(6), 1786 or R. Noyori et al., J. Amer. Chem. Soc. 1974, 96(10), 3336.

The corresponding methyl esters may be prepared in a conventional manner starting from 2-furyl glyoxyl nitrile or 3-furyl glyoxyl nitrile via the 2-furyl glyoxylic acid or 3-furyl glyoxylic acid which can be obtained therefrom. The corresponding glycolates are obtained from these as described using the organometallic derivatives of 2-bromothiophene or 3-bromothiophene. The organometallic compounds which can be obtained from 2-, 3- or 4-halopyridine can be reacted with methyl 2-thienylglyoxylate or methyl 3-thienylglyoxylate to give the corresponding glycolates.

Thienylglycolates, in which the thiophene ring contains fluorine in the 2- or 3-position, are prepared, for example starting from 2-fluorothiophene or 3-fluorothiophene (bromination to give 2-bromo-3-fluorothiophene or 2-bromo-5-fluorothiophene), and after conversion to the corresponding organometallic compounds, reaction with suitable glyoxylates to give the glycolates.

2-Fluorothiophene and 3-fluorothiophene can be reacted analogously to give the corresponding glyoxylates Unterhalt, Arch. Pharm. 322, 839 (1989) which in turn, as already described, may be reacted with, for example 2-thienyl or 3-thienyl derivatives, to give glycolates. Symmetrically substituted di-thienylglycolates can be prepared analogously by selecting suitable components.

A further route is available via a process analogous to the benzoin condensation and benzilic acid rearrangement.

The following examples illustrate the invention without limiting it.

#### EXAMPLE 1

#### EXAMPLE 1

#### Scopine di-(2-thienyl)glycolate

50.87 g (0.2 mole) of methyl di-(2-thienyl)glycolate and 31.04 g (0.2 mole) of scopine are dissolved in 100 ml of absolute toluene and reacted at a bath temperature of 90° C. with addition of 1.65 g (0.071 gram atom) of sodium in several portions. The resulting methanol is distilled off at a reaction mixture temperature of 78°-90° C. under a pressure of 500 mbar. After a reaction time of about 5 hours, the reaction mixture is stirred into a mixture of ice and hydrochloric acid. The acid phase is separated off, rendered alkaline using sodium carbonate and the free base is extracted using methylene chloride. After drying over sodium sulphate, the methylene chloride is distilled off under reduced pressure and the residue is recrystallised from acetonitrile; beige-coloured crystals (from acetonitrile), m.p. 149°-50° C.,

Yield: 33.79 g (44.7% of theoretical).

#### EXAMPLE 2

#### Scopine di-(2-thienyl)glycolate

12.72 g (0.05 mole) of methyl di-(2-thienyl)glycolate and 7.76 g (0.05 mole) of scopine are melted in a heating bath

at 70° C. under a water jet vacuum. 2.70 g (0.05 mole) of sodium methylate are introduced into this melt and heated for 1 hour in a heating bath at 70° C. under a water jet vacuum and subsequently for a further hour in a heating bath at 90° C. The solidified melt is taken up in a mixture of 100 ml of water and 100 ml of methylene chloride while monitoring the temperature, and the methylene chloride phase is extracted several times using water. The methylene chloride phase is extracted using the corresponding amount of dilute hydrochloric acid. The scopine di-(2-thienyl)glycolate is extracted from the combined aqueous phases using methylene chloride after adding the corresponding amount of sodium carbonate and dried over sodium sulphate. The hydrochloride is prepared from the dried methylene chloride solution in a conventional manner. The crystals are filtered off under suction, washed using acetone and dried under reduced pressure at 35° C. Pale yellow crystals (from methanol), m.p. 238°-41° C. (decomposition);

Yield: 10.99 g (53.1% of theoretical).

The hydrochloride may be converted to the base in a conventional manner.

### EXAMPLE 3

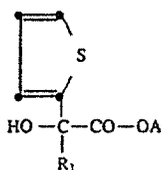
#### Scopine di-(2-thienyl)glycolate

38.15 g (0.15 mole) of methyl di-(2-thienyl)glycolate and 23.28 g (0.15 mole) of scopine are mixed, 0.34 g (0.015 gram atom) of sodium is added and the mixture is melted in a heating bath at 90° C. under a water jet vacuum. The reaction lasts 2.5 hours. 100 ml of absolute toluene are then added and the mixture is stirred at a heating bath temperature of 90° C. until a solution is produced. The reaction solution is cooled to room temperature and stirred into a mixture of ice and hydrochloric acid cooled using ice. The hydrochloride of the basic ester crystallising out is filtered off under suction and washed using a small amount of water and a large amount of diethyl ether. The filtrate phases are separated off and the aqueous phase is extracted using diethyl ether. The hydrochloride filtered off under suction is suspended in the (acid) aqueous phase and converted to the base while monitoring the temperature and adding the corresponding amount of sodium carbonate; the base is extracted using methylene chloride. The combined methylene chloride phases are dried over sodium sulphate. After distilling off the methylene chloride, crystals remain which are purified over active charcoal and recrystallised from acetonitrile. Pale yellow crystals (from acetonitrile), m.p. 148°-49° C.;

Yield: 39.71 g (70.1% of theoretical).

TABLE I

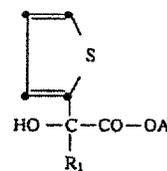
Compounds of the formula



No. A	R <sub>1</sub>	Base	M.p. [°C.] Hydrochloride
1	3α-(6β,7β-epoxy)-tropanyl	2-thienyl	149-50 238-41
2	3α-tropanyl	2-thienyl	167-8 253

TABLE I-continued

Compounds of the formula



No. A	R <sub>1</sub>	Base	M.p. [°C.] Hydrochloride
3	3α-(6,7-dehydro)-tropanyl	2-thienyl	164-5
4	3α-(N-β-fluoroethyl)-nortropanyl	2-thienyl	236
5	3α-(N-isopropyl)-granatanyl	2-thienyl	232
6	3α-(N-isopropyl)-nortropanyl	2-thienyl	256
7	3α-(6β,7β-epoxy)-N-isopropyl-nortropanyl	2-thienyl	206
8	3α-(6β,7β-epoxy)-N-ethyl-nortropanyl	2-thienyl	212-3
9	3α-(N-ethyl)-nortropanyl	2-thienyl	256-7
10	3α-(N-N-methyl)-granatanyl	2-thienyl	241
11	3α-(6β,7β-epoxy)-N-β-fluoroethyl-nortropanyl	2-thienyl	188-90
12	3α-(6β,7β-epoxy)-N-n-propylnortropanyl	2-thienyl	104-6
13	3α-(6β,7β-epoxy)-N-n-butyl-nortropanyl	2-thienyl	225-7
14	3α-(6β,7β-epoxy)-tropanyl	phenyl	246-7
15	3α-tropanyl	phenyl	243-4
16	3α-(N-β-fluoroethyl)-nortropanyl	phenyl	219-20
17	3α-(6,7-dehydro)-tropanyl	phenyl	181-3
18	3α-(N-ethyl)-nortropanyl	phenyl	231-2
19	3α-(N-isopropyl)-nortropanyl	phenyl	246-7
20	3α-tropanyl	cyclohexyl	260
21	3α-(N-β-fluoroethyl)-nortropanyl	cyclohexyl	203-4
22	3α-(6β,7β-epoxy)-tropanyl	cyclopentyl	237
23	3α-tropanyl	cyclopentyl	260
24	3α-(N-β-fluoroethyl)-nortropanyl	cyclopentyl	182-3
25	3α-(N-ethyl)-nortropanyl	cyclopentyl	227-8
26	3α-(N-isopropyl)-nortropanyl	cyclopentyl	174-5
27	3α-(6β,7β-epoxy)-tropanyl	2-thienyl	240-2
28	3β-tropanyl	2-thienyl	217-9
29	3β-(6,7-dehydro)-tropanyl	2-thienyl	233-5
30	3α-(6,7-dehydro)-tropanyl	3-thienyl	247-8
31	3α-(6β,7β-epoxy)-tropanyl	3-thienyl	242-3
32	3α-(6β,7β-epoxy)-tropanyl	2-furyl	
33	3α-(6,7-dehydro)-tropanyl	2-furyl	
34	3α-tropanyl	2-furyl	
35	3α-tropanyl	2-pyridyl	
36	3α-(6β,7β-epoxy)-tropanyl	2-pyridyl	
37	3α-(6,7-dehydro)-tropanyl	2-pyridyl	
38	3α-tropanyl	3-thienyl	
39	3α-(6,7-dehydro)-tropanyl	cyclopentyl	
40	3α-(6β,7β-epoxy)-tropanyl	cyclohexyl	
41	3α-(6,7-dehydro)-tropanyl	cyclohexyl	

65 Note: All hydrochlorides melt with decomposition.

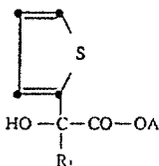
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EXAMPLE 4

## Scopine di-(2-thienyl)glycolate methobromide

10.0 g (0.0265 mole) of scopine di-(2-thienyl)glycolate  
are dissolved in a mixture comprising 20 ml of anhydrous  
methylene chloride and 30 ml of anhydrous acetonitrile and  
treated with 12.8 g (0.1325 mole) of methyl bromide (as  
50% strength solution in anhydrous acetonitrile), and the  
reaction mixture is allowed to stand for 24 hours at room  
temperature in a tightly sealed reaction vessel. Crystals are  
precipitated during this time. They are filtered off under  
suction, washed using methylene chloride and dried at 35°  
C. under reduced pressure. White crystals (from methanol/  
acetone), m.p. 217°-8° C. (decomposition) after drying at  
111° C. under reduced pressure.

TABLE II

Quaternary compounds of the formula

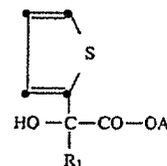


No. A	R <sub>1</sub>	M.p. [°C.]
1	3α-(6β,7β-epoxy)-tropanyl methobromide	2-thienyl 217-18
2	3α-tropanyl methobromide	2-thienyl 263-64
3	3α-(6,7-dehydro)-tropanyl methobromide	2-thienyl 191-92
4	3α-(N-β-fluoroethyl)-nortropanylmethobromide	2-thienyl 242-43
5	3α-tropanyl-β-fluoroethobromide	2-thienyl 214-15
6	3α-(N-isopropyl)-granatanylmethobromide	2-thienyl 229-30
7	3α-(N-isopropyl)-nortropanylmethobromide	2-thienyl 245-46
8	3α-(6β,7β-epoxy)-N-isopropyl-nortropanyl methobromide	2-thienyl 223-24
9	3α-(6β,7β-epoxy)-N-ethylnortropanyl methobromide	2-thienyl 215-16
10	3α-(N-ethyl)-nortropanyl methobromide	2-thienyl 260-61
11	3α-(N-methyl)-granatanylmethobromide	2-thienyl 246-47
12	3α-(6β,7β-epoxy)-N-fluoroethyl-nortropanyl methobromide	2-thienyl 182-83
13	3α-(6β,7β-epoxy)-N-propylnortropanyl methobromide	2-thienyl 209-10
14	3α-tropanyl-β-hydroxyethobromide	2-thienyl 231-32
15	3α-(6β,7β-epoxy)-tropanyl ethobromide	phenyl 217-18
16	3α-tropanyl methobromide	phenyl 273-74
17	3α-(N-β-fluoroethyl)-nortropanylmethobromide	phenyl
18	3α-(6,7-dehydro)-tropanyl methobromide	phenyl 110-71
19	3α-(N-ethyl)-nortropanyl methobromide	phenyl 249-50
20	3α-(N-isopropyl)-nortropanyl methobromide	phenyl 259-60

## 12

TABLE II-continued

Quaternary compounds of the formula



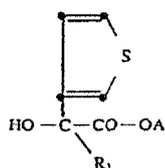
No. A	R <sub>1</sub>	M.p. [°C.]
21	3α-tropanyl ethobromide	phenyl 248-49
22	3α-(N-ethyl)-nortropanyl ethobromide	phenyl 244-45
23	3α-(6β,7β-epoxy)-tropanyl ethobromide	phenyl 226
24	3α-tropanyl-β-fluoroethobromide	phenyl 241
25	3α-tropanyl methobromide	cyclohexyl 278
26	3α-(N-β-fluoroethyl)-nortropanyl methobromide	cyclohexyl 198
27	3α-tropanyl-β-fluoroethobromide	cyclohexyl 233-34
28	3α-tropanyl methobromide	cyclopentyl 260
29	3α-tropanyl ethobromide	cyclopentyl 235-36
30	3α-(N-ethyl)-nortropanyl methobromide	cyclopentyl 251-52
31	3α-(N-isopropyl)-nortropanyl-methobromide	cyclopentyl 244-45
32	3α-tropanyl-β-fluoroethobromide	cyclopentyl 189-90
33	3α-(N-β-fluoroethyl)-nortropanyl-methobromide	cyclopentyl 226-27
34	3α-(6,7-dehydro)-tropanyl metho-methanesulphonate	2-thienyl 225-6
35	3α-(6β,7β-epoxy)-tropanyl methobromide	2-thienyl 218-20
36	3α-tropanyl methobromide	2-thienyl 243-4
37	3α-(6,7-dehydro)-tropanyl methobromide	2-thienyl 211-4
38	3α-(6,7-dehydro)-tropanyl methobromide	3-thienyl 182-3*
39	3α-(6β,7β-epoxy)-tropanyl methobromide	3-thienyl 217-8
40	(+) enantiomer of No. 1	
41	(-) enantiomer of No. 1	
42	3α-(6β,7β-epoxy)-tropanyl methobromide	2-furyl
43	3α-(6,7-dehydro)-tropanyl methobromide	2-furyl
44	3α-tropanyl methobromide	2-furyl
45	3α-(6β,7β-epoxy)-tropanyl methobromide	2-pyridyl
46	3α-(6,7-dehydro)-tropanyl methobromide	2-pyridyl
47	3α-tropanyl methobromide	2-pyridyl
48	3α-tropanyl methobromide	3-thienyl
49	3α-(6,7-dehydro)-tropanyl methobromide	cyclopentyl
50	3α-(6β,7β-epoxy)-tropanyl methobromide	cyclohexyl
51	3α-(6,7-dehydro)-tropanyl methobromide	cyclohexyl
52	3α-(6β,7β-epoxy)-tropanyl methobromide	cyclopentyl

\*contains crystalline methanol

60 Note: All compounds in the table melt with decomposition.

TABLE III

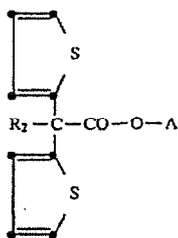
Compounds of the formula



No.	A	R <sub>1</sub>	M.p. [°C.] Hydrochloride
1	3α-(6β,7β-epoxy)-tropanyl	phenyl	246-7
2	3α-(6,7-dehydro)-tropanyl	phenyl	261-2
3	3α-(6β,7β-epoxy)-tropanyl	3-thienyl	
4	3α-(6,7-dehydro)-tropanyl	3-thienyl	
5	3α-tropanyl	3-thienyl	
6	3α-(N-methyl)-granatanyl	3-thienyl	

TABLE IV

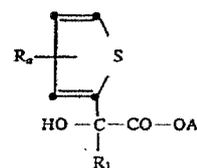
Compounds of the formula



No.	A	R <sub>2</sub>	M.p. [°C.] Hydrochloride
1	3α-(6β,7β-epoxy)-tropanyl	H	
2	3α-(6,7-dehydro)-tropanyl	H	
3	3α-(6β,7β-epoxy)-tropanyl	methyl	210-2.5
4	3α-(6,7-dehydro)-tropanyl	methyl	
5	3α-(6β,7β-epoxy)-tropanyl	methoxy	
6	3α-(6,7-dehydro)-tropanyl	methoxy	

TABLE V

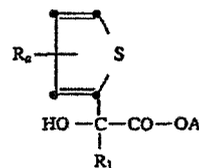
Compounds of the formula



No.	A	R <sub>2</sub>	R <sub>a</sub>	M.p. [°C.]
15	1	3α-(6β,7β-epoxy)-tropanyl	2-thienyl	5-methyl
	2	3α-(6,7-dehydro)-tropanyl	2-thienyl	5-methyl
	3	3α-tropanyl	2-thienyl	5-methyl
	4	3α-(6β,7β-epoxy)-tropanyl	2-(5-methyl)-thienyl	5-methyl
20	5	3α-(6,7-dehydro)-tropanyl	2-(5-methyl)-thienyl	5-methyl
	6	3α-tropanyl	2-(5-methyl)-thienyl	5-methyl
	7	3α-(6β,7β-epoxy)-tropanyl	2-thienyl	5-fluoro
25	8	3α-(6,7-dehydro)-tropanyl	2-thienyl	5-fluoro
	9	3α-tropanyl	2-thienyl	5-fluoro
	10	3α-(6β,7β-epoxy)-tropanyl	2-(5-fluoro)-thienyl	5-fluoro
30	11	3α-(6,7-dehydro)-tropanyl	2-(5-fluoro)-thienyl	5-fluoro
	12	3α-tropanyl	2-(5-fluoro)-thienyl	5-fluoro

TABLE VI

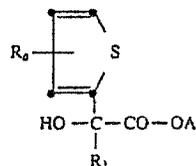
Compounds of the formula



No.	A	R <sub>1</sub>	R <sub>a</sub>	M.p. [°C.]
1	3α-(6β,7β-epoxy)-tropanyl methobromide	2-thienyl	5-methyl	
2	3α-(6,7-dehydro)-tropanyl methobromide	2-thienyl	5-methyl	
3	3α-tropanyl-methobromide	2-thienyl	5-methyl	
4	3α-(6β,7β-epoxy)-tropanyl methobromide	2-(5-methyl)-thienyl	5-methyl	

TABLE VI-continued

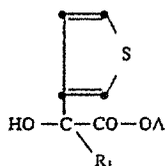
Compounds of the formula



No.	A	R <sub>1</sub>	R <sub>2</sub>	M.p. [°C.]
5	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	2-(5-methyl)-thienyl	5-methyl	
6	3 $\alpha$ -tropanyl methobromide	2-(5-methyl)-thienyl	5-methyl	
7	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	2-thienyl	5-fluoro	
8	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	2-thienyl	5-fluoro	
9	3 $\alpha$ -tropanyl methobromide	2-thienyl	5-fluoro	
10	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	2-(5-fluoro)-thienyl	5-fluoro	
11	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	2-(5-fluoro)-thienyl	5-fluoro	
12	3 $\alpha$ -tropanyl methobromide	2-(5-fluoro)-thienyl	5-fluoro	

TABLE VII

Compounds of the formula

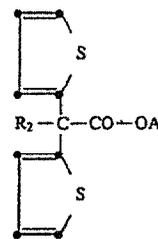


No.	A	R <sub>1</sub>	M.p. [°C.]
1	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	phenyl	211-2
2	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	phenyl	158-60*
3	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	3-thienyl	
4	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	3-thienyl	
5	3 $\alpha$ -tropanyl methobromide	3-thienyl	
6	3 $\alpha$ -(N-methyl)-granatanyl methobromide	3-thienyl	

\*(with crystalline methanol)

TABLE VIII

Quaternary compounds of the formula

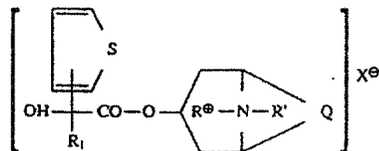


No.	A	R <sub>2</sub>	M.p. [°C.]
1	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	H	
2	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	H	
3	3 $\alpha$ -(6 $\beta$ ,7 $\beta$ -epoxy)-tropanyl methobromide	methyl	
4	3 $\alpha$ -(6,7-dehydro)-tropanyl methobromide	methyl	206-8
5	3 $\alpha$ -tropanyl methobromide	methoxy	
6	3 $\alpha$ -(N-methyl)-tropanyl methobromide	methoxy	

17

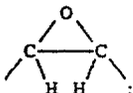
We claim:

1. A compound of the formula

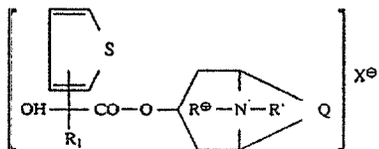


wherein

Q is a group of the formula  $-\text{CH}_2-\text{CH}_2-$ ,  
 $-\text{CH}=\text{CH}-$  or

R and R' are each independently C<sub>1</sub>-C<sub>4</sub>-alkyl;R<sub>1</sub> is thienyl, phenyl, cyclopentyl or cyclohexyl; andX<sup>-</sup> is a physiologically acceptable anion.

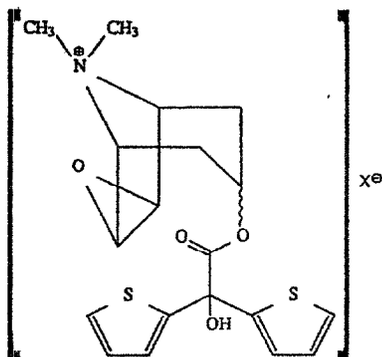
2. A compound in accordance with claim 1, of the formula



wherein

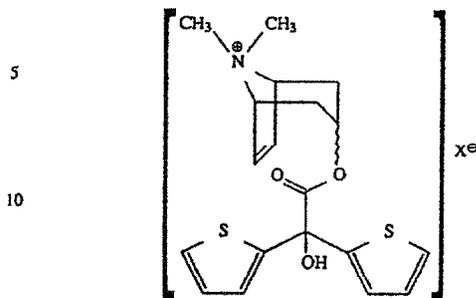
R is CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub>, n-C<sub>3</sub>H<sub>7</sub>, or i-C<sub>3</sub>H<sub>7</sub>;R' is CH<sub>3</sub>; andR<sub>1</sub>, Q and X<sup>-</sup> are as defined in claim 1.3. A compound in accordance with claim 2 wherein R<sub>1</sub> is thienyl.4. A compound in accordance with claim 2 wherein X<sup>-</sup> is Br<sup>-</sup> or CH<sub>3</sub>SO<sub>3</sub><sup>-</sup>.

5. A compound of the formula

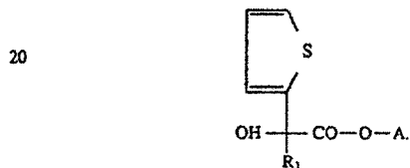
wherein X<sup>-</sup> is a physiologically acceptable anion.

18

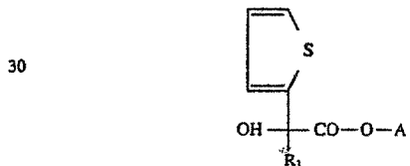
6. A compound of the formula

wherein X<sup>-</sup> is a physiologically acceptable anion.

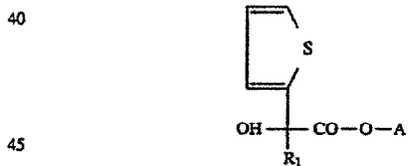
7. A compound of the formula



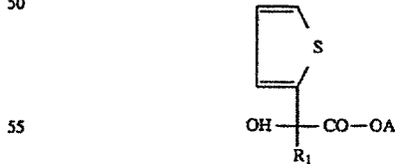
8. A compound of the formula

wherein R<sub>1</sub> is 2-thienyl and A is 3α-(6,7-dehydro)-tropanyl methobromide.

9. A compound of the formula

wherein R<sub>1</sub> is 2-thienyl and A is 3β-tropanyl methobromide.

10. A compound of the formula

wherein R<sub>1</sub> is cyclopentyl and A is 3α-(N-isopropyl)-nortropanyl methobromide.

11. A method for treating chronic obstructive bronchitis which comprises administering, by inhalation, to a subject suffering from the same, a therapeutic amount of a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

12. A method for treating slight to moderately severe asthma which comprises administering, by inhalation, to a subject suffering from the same, a therapeutic amount of a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

13. A method for treating vagally induced sinus bradycardia which comprises administering, by the intravenous or oral routes, to a subject suffering from the same, a therapeutic amount of a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

14. A pharmaceutical composition, for administration by inhalation, suitable for the treatment of chronic obstructive bronchitis or slight to moderately severe asthma, which

comprises a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

15. A pharmaceutical composition, for oral administration, suitable for the treatment of vagally induced sinus bradycardia, which comprises a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

16. A pharmaceutical composition, for intravenous administration, suitable for the treatment of vagally induced sinus bradycardia, which comprises a compound in accordance with claims 1, 2, 3, 4, 6, 7, 8, 9, 10.

\* \* \* \* \*