Antitussive Effect of RU-20201—Central and Peripheral Actions

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Abstract—The antitussive effect of the new compound 1, 2, 3, 4a, 9b-hexahydro-8, 9b-dimethyl-4-[3-(4-methyl-piperazine-1-yl) propionamide] dibenzofuran-3-one dihydrochloride (RU-20201) was investigated in dogs and guinea pigs, including its sites of action. The antitussive effect of RU-20201 was about 1/10 as potent as that of codeine phosphate in dogs with the puncture electrode-induced cough (PEC) method and about 1/12 and 1/4 as potent as that of codeine phosphate in guinea pigs with the PEC and chemical stimulation methods, respectively. When RU-20201 was administered in a dose range of 1 to 10 mg into the vertebral artery toward the brain in lightly anesthetized dogs, no antitussive effect was observed against the coughing elicited by electrical stimulation of the central cut end of the superior laryngeal nerve. However, a stimulative effect on respiration, especially on respiratory rate occurred. The peripheral effect of RU-20201 on the cough was investigated using the in situ upper trachea perfusion preparation which allows a direct drug administration to the local site around the tracheal mucosa, this site being electrically stimulated to induce coughing. A close i.a. infusion of RU-20201 in doses of 1 and 3 mg/min into the tracheal vascular bed for 5 min inhibited the cough response elicited by mucosal stimulation. The above findings suggest that RU-20201 has a significant antitussive activity, the site of action being probably, at least, at the cough receptor level.

1, 2, 3, 4, 4a, 9b-Hexahydro-8, 9b-dimethyl-4-[3-(4-methyl-piperazine-1-yl) propionamide] dibenzofuran-3-one dihydrochloride (RU-20201) (Fig. 1) is an antitussive agent originally found from dibenzofuran derivatives by Matharu et al. (1). They reported that RU-20201 possesses an antitussive effect as potent as that of codeine phosphate in guinea pigs when administered orally. Korpas et al. (2) also demonstrated its antitussive effect in cats when given by inhalation, and Pickering and James (3) demonstrated the effect in guinea pigs and rabbits when given orally and by inhalation.

Codeine phosphate, widely used as an antitussive drug, has an inhibitory effect on respiration as well as on cough (4). On the other hand, dextromethorphan, a non-narcotic antitussive agent, does not inhibit respiration, but rather stimulates it (5). Antitussive agents which produce no inhibitory effect or rather produce a stimulatory effect on respiration are clinically preferred. Korpas et al. (2) and Pickering and James (3) considered that RU-20201 exerts an antitussive effect through a peripheral mechanism, although no sufficient experimental evidence has been available to support this.

In the present study, the antitussive and respiratory effects of RU-20201 were investigated in dogs and guinea pigs, being compared with results obtained for codeine

Fig. 1. Chemical structure of RU-20201.
phosphate. Whether the site of action of its antitussive effect is on the central level or peripheral level was also studied using methods devised by the authors.

Materials and Methods

1) Antitussive effect in the dog
The puncture electrode-induced cough (PEC) method previously described (6) was used with a partial modification. Unanesthetized and unrestrained male mongrel dogs weighing 9–15 kg were used. After the trachea caudal to the thyroid cartilage of the dog was palpated, a stimulation electrode was inserted into the trachea through a guiding cannula (stainless steel needle for s.c., size 18, Terumo) and advanced to the bifurcation tracheae. As the electrode, a stainless steel wire (0.2 mm in diameter and 18 cm in length) coated with a cashew resin for insulation was used. Before use, the insulating layer was removed from the tips of either side of the electrode over a length of 2–3 cm, so that electricity could be transferred. This modification done in the present study ensured a more tight insulation and easier preparation, compared to the conventional method in which a stainless steel wire is wrapped with thin polyvinyl film. This cashew-coated electrode was sheathed with a stainless steel pipe (10 cm in length and 0.8 mm in outer diameter) and fixed with an adhesive substance, Alonalpha, at either end of the pipe. The tip that should be inserted into the trachea was bent at an angle of approximately 120° so that it easily had contact with the mucous membrane of the trachea. A stainless steel needle placed into a subcutaneous site of the back was used as an indifferent electrode. For inducing the cough reflex, square-wave negative pulses (20 Hz, 1 m/sec, 6–8 V) were applied for 10 sec. When coughing was induced too excessively, the puncture electrode was then drawn towards the larynx so that coughing occurred 5 to 9 times per stimulus. Prior to drug administration, electrical stimulations were given at an interval of 10 min until almost the same number of coughs could be evoked by each of three consecutive stimulations, from which an average pre-drug control value was calculated. Sessions of stimulations for inducing coughs were given 5, 15, 30, 45, 60 and 90 min after either RU-20201 or codeine phosphate (as a reference drug) was administered i.v. into the cephalic vein. The antitussive effect was evaluated as an inhibitory ratio in terms of the percent reduction in the number of coughings compared to the average pre-drug control value.

2) Antitussive effect in the guinea pig
Male Hartley guinea pigs weighing 350–450 g were used. Chemical stimulation was given according to the method of Takagi et al. (7), and electrical stimulation was according to the method of Yanaura et al. (8). Both tests were slightly modified for the purpose of the present experiment.

a) Chemical stimulation (SO2) method:
In the method of Takagi et al., sulfur dioxide gas for inducing coughs is obtained by adding conc. sulfuric acid to the saturated solution of sodium bisulfite. Instead, we used a gas cylinder containing highly purified liquid sulfur dioxide to simplify the method. The test apparatus used is shown in Fig. 2. The sulfur dioxide gas detector (A), desiccator (C) and gas cylinder (D) were connected to each other with silicone tubes. The cock (b) was first closed, and the pressure regulator gauge (c) was set at a given position so that sulfur dioxide gas at a given pressure entered the silicone tube located between (b) and (c). The valve (d) was then closed in order to confine the gas in the silicone tube. The cock (b) was then opened to introduce the confined gas into the desiccator. The air pump (B) was used to make a uniform

Fig. 2. Apparatus for the chemical stimulation method used in this study. A: Sulfur dioxide gas detector, B: Air pump, C: Desiccator, D: Sulfur dioxide gas cylinder, a, b: Glass cock with stopper, c: Pressure regulator gauge, d: Gas cylinder valve.
concentration of sulfur dioxide gas throughout
the desiccator. The cock (a) was opened in
case of need to check the concentration of
sulfur dioxide gas in the desiccator with the
gas detector. A gas concentration at 600 to
800 ppm was adequate to evoke coughing.
A guinea pig was placed on the perforated
plate inside the desiccator. Sulfur dioxide gas
was then introduced into the desiccator
according to the method outlined above.
After one-minute exposure to the gas, the
animal was taken out and placed in a cage.
Observation was made for 5 min to check if
the animal was coughing or not. Animals
which coughed twice or more during 5 min
in a preliminary test were used in this ex-
periment. The exposure was repeated 15 min
before (pre-drug control) and 15 and 30 min
after drug administration. When no cough
occurred in either of the exposures given 15
and 30 min after drug administration, the
drug was regarded as effective. ED50 was
calculated with the up and down method of
Brownlee et al. (9).

b) PEC method: Electrical stimulation was
given to guinea pigs as mentioned in the
previous paper (7). Unanesthetized guinea
pigs were fixed in a supine position. A
puncture electrode was introduced into the
trachea through a guiding cannula, the tip
of the electrode being placed near the
bifurcation tracheae. For inducing the cough
reflex, square-wave pulses (40 Hz, 1 msec, 
2 V) were applied for 5 sec. When no
coughing occurred, stimulation was added
again at an interval of 5 sec. Four sessions of
stimulation to induce the cough reflex were
given: twice before drug administration and
twice (15 and 30 min) after drug adminis-
tration. The effects of drugs were evaluated
in the same manner as in the case of the
chemical stimulation method.

3) Effect on cough reflex and respiration
when administered into the vertebral artery

Male mongrel dogs weighing 9-15 kg
anesthetized with pentobarbital Na (30 mg/
kg i.v.) were used in the experiment for
determining the central effect of RU-20201.
The test method used was described in the
previous paper (10). The drug solution was
injected into the vertebral artery at a site
5 cm above the clavicle towards the brain.

Sessions of electrical stimulation for inducing
the cough reflex were given twice at an
interval of 5 min before drug administration
and were also given 1, 3, 5, 10, 15 and 30
min after drug administration.

4) Effect on the cough receptor site

The in situ canine upper trachea perfusion
preparation (11) was used. Electrical stimu-
lation was given to induce the cough reflex
with a silver disc electrode placed on the
upper tracheal mucosa where drug perfusion
was made. For inducing the cough reflex,
square-wave pulses (20 Hz, 1 msec, 8-10 V)
were applied for 10 sec. RU-20201 or
procaine hydrochloride was infused with an
infusion pump (Natsume, KN-202) for 5 min
at a rate of 0.04 ml/min via a rubber tube
into the perfused cranial thyroid artery.
Stimulation was repeated 1, 3 and 5 min
after the start of drug infusion and 3, 5, 10,
15, 30, 45 and 60 min after the cessation of
infusion. The effect of drug was evaluated
with the same criteria as in the experiment in
which the antitussive effect of i.v. adminis-
tration of drugs was investigated in the dog
(as described above).

5) Drugs

RU-20201 (Roussel Uclaf), codeine phos-
phate (Sankyo) and procaine hydrochloride
(Sanko) were used. All drugs were dissolved
in physiological saline before use. Doses are
shown in terms of base quantity.

Results

1) Antitussive effects in the dog: The
antitussive effect of RU-20201 in dogs is
shown in Fig. 3, and that of codeine phos-
phate is shown in Fig. 4. Almost no anti-
tussive effect was observed with 10 mg/kg
of RU-20201. With 20 and 30 mg/kg,
however, the number of coughs evoked was
reduced 5 min after each administration by
about 50% and 90%, respectively (P<0.01).
When 30 mg/kg of RU-20201 was used, the
antitussive effect lasted for 45 min or more
after administration. On the other hand, 1
and 3 mg/kg of codeine phosphate showed a
significant antitussive effect (P<0.05) which
lasted for about 60 min. The potency of RU-
20201 was found to be between one tenth
and one twentieth of that of codeine phos-
phate in terms of cough-inhibitory ratio
calculated 5 min after each administration.

2) Antitussive effects in the guinea pig:
The results of the chemical stimulation method are shown in Table 1a. The 50% antitussive dose of RU-20201 was 66.1 mg/kg i.p., and that of codeine phosphate was 18.2 mg/kg i.p. The potency of RU-20201 was found to be about one fourth that of codeine phosphate. The results of the electrical stimulation method are shown in Table 1b. The 50% antitussive dose of RU-20201 was 119.2 mg/kg i.p., and that of codeine phosphate was 10.0 mg/kg i.p. In this case, the potency of RU-20201 was found to be about 1/12 that of codeine phosphate.

3) Effect on cough reflex and respiration when administered into the vertebral artery:
The number of coughs (N.C.) and the amplitude of cough (A.C.) were used as indices of the cough reflex. When codeine phosphate was administered via the vertebral artery, N.C. decreased dose-dependently (Fig. 5). When 1.0 mg of the drug was given, N.C. was inhibited by a maximum of 75% (P<0.01), and A.C. was inhibited by a maximum of 60% (P<0.05). The inhibitions of N.C. and A.C. lasted for at least 15 min after administration. In the case of RU-20201 injected into the vertebral artery, almost no
A change was observed in both parameters in a dose range of 1.0 to 10.0 mg (Fig. 6). Respiratory rate (R.R.), respiratory amplitude (R.A.) and respiratory volume (R.V.) were used as indices of respiration. In the case of codeine phosphate, no significant change in R.A. was observed for any of the doses of 0.1, 0.3 and 1.0 mg (Fig. 7). When 0.3 and 1.0 mg of the drug were used, however, 20% inhibition was observed both in R.R. and R.V. In the case of RU-20201, no significant change in R.A. was observed for any of the following doses: 1.0, 3.0 and 10.0 mg (Fig. 8). When 10.0 mg of RU-20201 was injected, however, it was observed that R.R. increased by about 40% (P<0.05) and R.V. by about 45%. Neither RU-20201 nor codeine phosphate affected systemic blood pressure at the doses used in this study. The results of the experiment with the drugs administered into the vertebral artery are summarized in Table 2.

**Table 1.** Fifty percent antitussive doses (AtD50s) of RU-20201 and codeine: a) with the chemical stimulation (SO2) method and b) with the puncture electrode-induced cough (PEC) method in conscious guinea pigs

<table>
<thead>
<tr>
<th>Drug</th>
<th>AtD50 (mg/kg)</th>
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<tr>
<td>RU-20201</td>
<td>66.1</td>
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<tr>
<td>Codeine</td>
<td>18.2</td>
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<th>Drug</th>
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<tr>
<td>RU-20201</td>
<td>119.2</td>
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<tr>
<td>Codeine</td>
<td>10.0</td>
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Drugs were administered i.p. Each value represents the average value for two up and down experiments.

![Fig. 5. The effect of codeine administered intravertebroarterially on the cough reflex induced by electrical stimulation given to the central cut end of the superior laryngeal nerve in dogs. Each point represents the mean percent change with S.E. (N=3-6, *P<0.05 and **P<0.01 vs. the control).](image-url)
Fig. 6. The effect of RU-20201 administered intravertebrally on the cough reflex induced by electrical stimulation given to the central cut end of the superior laryngeal nerve in dogs. Each point represents the mean percent change with S.E. (N=3-6).

Fig. 7. The effect of codeine administered intravertebrally on respiration in dogs. Graphs show changes in rate, amplitude and volume of respiration for 3 min after drug administration. Each column represents the mean percent change with S.E. (N=3-8, *P<0.05 and **P<0.01 vs. the control).

4) Effect on the cough receptor site: When RU-20201 (1 mg/min) was infused for 5 min into the cranial thyroid artery, about 30% cough inhibition was observed (P<0.05). The inhibitory effect lasted for 5–30 min after the cessation of infusion. When 3 mg/min of RU-20201 was infused, about 40% cough inhibition was observed (P<0.05) 3 min after the cessation of administration (Fig. 9a). When 3 mg/min of procaine was infused, a maximum of 80% cough inhibition (P<0.05) was seen (Fig. 9b). The inhibitory effect lasted for 15–30 min after the cessation of infusion.

Discussion

The antitussive effect of RU-20201 in guinea pigs has already been demonstrated
in oral administration tests by Matharu et al. (1) and Pickering and James (3). It is reported that the antitussive potency of RU-20201 is almost equal to that of codeine phosphate. RU-20201 in the form of an aerosol agent has been tested in cats and rabbits by Korpas et al. (2) and Pickering and James (3). These experiments also confirmed the antitussive effect of RU-20201. Concerning general pharmacological actions, RU-20201 has no action other than a weak local anesthetic action (12).

Based on the fact that the gastrointestinal absorption of RU-20201 is relatively greater than that of codeine phosphate (13, 14), we tried to compare the effects of both drugs by injecting them intravenously or intraperitoneally. In order to examine and assess their antitussive effects, in the present study, we employed the puncture electrode-induced cough (PEC) method and the chemical stimulation (SO₂) method. In the chemical stimulation method of Takagi et al., sulfur dioxide gas is obtained by adding conc. sulfuric acid to the saturated solution of sodium bisulfite. However, the procedure is rather complicated. We therefore used a simpler method employing a gas cylinder of

![Graph showing the effect of RU-20201 administered intravertebroarterially on respiration in dogs](image)

**Table 2.** The effects of codeine and RU-20201 given intravertebroarterially on respiration and cough reflex in dogs

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<td>RU-20201</td>
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highly pure liquefied sulfur dioxide. In the present study, the antitussive effects of RU-20201 administered intravenously or intraperitoneally were found to be between ten to twenty times less than those of codeine phosphate in the case of dogs using the electrical stimulation method and in the case of guinea pigs, about four times less in the chemical stimulation method and about twelve times less in the electrical stimulation method. It has been reported by Matharu et al. (1), however, that RU-20201 and codeine phosphate have a practically equipotent antitussive effect in the case of oral administration. The main reason for this discrepancy seems to lie in the fact that the gastrointestinal absorption of codeine phosphate is poor as compared with RU-20201 (13, 14).

The arc of the cough reflex consists of the cough receptors, afferent pathway, cough center, efferent pathway and respiratory organs (15, 16). Theoretically, coughing can be suppressed by blocking or inhibiting this arc at any point. Yamatsu et al. and Kase et al. (17, 18) administered some antitussive drugs through various routes including the femoral vein, common carotid artery, vertebral artery and cerebellomedullary cistern in order to disclose the central action. In the present study, the central actions of the two drugs were compared by administering them into the vertebral artery. The results with the intra-arterial administration of codeine phosphate revealed that the agent had an excellent antitussive effect, whereas no antitussive effect was found when RU-20201 was administered by the same route, even in a dose 10 times greater than that of codeine phosphate. This result therefore suggests that RU-20201 might not directly affect the cough center. This statement is also supported by the finding that the amount of RU-20201 entering the brain was considerably lower than that entering other organs when the drug was administered to rats intravenously (13).

The respiratory action of both drugs was simultaneously studied along with their antitussive one. Codeine phosphate was found to have an inhibitory effect on respiration. There is a report (12) which states that a transient stimulation of respiration was observed when RU-20201 was intravenously administered in rabbits. In the present study, a similar stimulating effect on respiration was confirmed in dogs.

Kito et al. (19) and Yanaura et al. (20) found that fominoben, one of the antitussive drugs, stimulates respiration in spite of depressing coughs. In our previous experiment (20), fominoben was administered into the vertebral artery of the dog in the same manner as in the present study. An i.a. injection of fominoben (3 mg) caused a significant but transient stimulation of respiration, and it depressed coughs by about 50% 1 to 3 min after administration. RU-20201 and fominoben injected intra-arterially
have an almost equipotent stimulatory effect on respiration, but in contrast to fominoben, RU-20201 has no antitussive effect when administered by this route. The stimulating action on respiration of fominoben is considered to be based on a direct effect on the central respiratory neuronal mechanisms in the brain stem (21). The mechanism for the stimulatory effect of RU-20201 on respiration remains to be clarified.

Kase et al. (18) were the first who described a method to clarify the peripheral (receptor site) actions of antitussive drugs. They recorded the afferent impulses from the superior laryngeal nerve that were produced by electrical stimulation of the tracheal mucosa by means of a puncture electrode. Then, they assessed the action on the receptor level by observing a quantitative change in the impulses evoked. With this method, however, the effects of the drug on other sites than cough receptors may also be involved in the change because the drug is given systemically by intravenous administration. In the present study, therefore, the drug was applied locally to the receptor level where electrical stimulation was given to induce coughs, enabling the peripheral action of the drug to be observed in the region directly (11). One of the drugs which inhibit coughing by acting on the receptor level is a local anesthetic. Yanaura (22) state that coughing is inhibited by applying procaine to the tracheal mucosa. They also mention that coughing is inhibited by injecting procaine into the anterior thyroid artery which perfuses the stimulated area (23). The same result was obtained in the present study. This means that procaine shows an antitussive effect through its action on the cough receptor level. It was observed that RU-20201 also showed an inhibitory effect on the cough response by acting on the cough receptor level, but that this effect was weaker than that of procaine. A definite local anesthetic action has not been observed with RU-20201 (12). Benzonatate is said to show an antitussive effect through its selective anesthetic action on the stretch receptors in the lung (24, 25). It has been confirmed by Yanaura et al. (26) that benzonatate produces its antitussive effect through the action on the stretch receptors. It has also been confirmed by Yuizono (27) that the agent decreases the number of impulses from the stretch receptors. When benzonatate is applied to the cough receptor level, however, it does not cause an antitussive effect (23) nor decreases afferent impulses from the receptors (27). Thus, an action of an antitussive drug at the cough receptor level is not necessarily explained by the local anesthetic action. It is therefore unreasonable to discuss the control of coughing only in terms of a general local anesthetic action. The mechanism of the peripheral action shown by RU-20201 has not yet been clarified. RU-20201 is, however, a very promising antitussive drug that has a different mechanism of action from those of previous and existing antitussive drugs.

Generally, antitussive drugs which work on the cough center often inhibit the respiratory center too. Clinically, however, a drug is considered to be a useful antitussive preparation if it does not inhibit respiration but in fact stimulates it. RU-20201 may be very useful from this aspect because it stimulates respiration.

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