Antitussive activity of iodo-resiniferatoxin in guinea pigs

M Trevisani, A Milan, R Gatti, A Zanasi, S Harrison, G Fontana, A H Morice, P Geppetti

Background: Iodo-resiniferatoxin (I-RTX) has recently been described as an ultra potent antagonist of the transient receptor potential vanilloid-1 (TRPV1).

Methods: The ability of I-RTX to inhibit cough induced by inhalation of two putative TRPV1 stimulants (capsaicin and citric acid) was tested in non-anaesthetised guinea pigs. Pretreatment with I-RTX either intraperitoneally (0.03−0.3 μmol/kg) or by aerosol (0.1−3 μM) reduced the number of coughs produced by inhalation of citric acid (0.25 M) and capsaicin (30 μM) in a dose dependent manner. Capsazepine (CPZ) also reduced citric acid and capsaicin induced cough, but the activity of I-RTX was 10−100 times more potent than CPZ in all the experimental conditions tested.

Conclusions: I-RTX is a novel and potent antitussive drug which inhibits cough mediated by agents possibly acting via TRPV1 activation.

Cough is one of the most common reasons for medical consultation. In most cases, however, drugs currently available for antitussive treatment are only partially effective. In the search for safe and effective antitussive drugs, it has been noted that, in some animal models and in humans, tussigenic agents stimulate a recently cloned ion channel belonging to the transient receptor potential family of channels. Since this entity is activated by molecules with a vanilloid moiety it has been termed the vanilloid receptor-1 (TRPV1). It is widely used for this purpose. The observation that capsazepine (CPZ), a relatively selective but low potency TRPV1 antagonist, reduced both capsaicin and citric acid induced cough suggested that TRPV1 is involved in the tussive response induced by these two agents. A lowered threshold to capsaicin induced cough seems to be linked to the sex of the patient and has been seen in patients with chronic inflammatory airway diseases including asthma and chronic obstructive pulmonary disease (COPD). TRPV1 therefore seems to have a role in the physiological activation of the cough reflex, as well as in the exaggerated cough response observed during certain pathological states.

On this basis it can be predicted that TRPV1 antagonists are of therapeutic value not only in the treatment of cough in patients with asthma and COPD, but also in patients with other inflammatory diseases including post viral cough and other inflammatory diseases including gastro-oesophageal reflux where upregulation of TRPV1 sensitivity has been implicated. However, the TRPV1 antagonists currently available suffer from poor specificity or potency. For instance, ruthenium red selectively inhibits TRPV1 in a very narrow range of concentrations, whereas, because of its low potency, CPZ effectively blocks it only at high concentrations. It has recently been reported that iodo-resiniferatoxin (I-RTX), the iodinated form of the ultra potent TRPV1 agonist resiniferatoxin, behaves as a high affinity TRPV1 antagonist at the mouse and rat recombinant TRPV1. We have confirmed this early observation by showing that, in a series of “typical” nocebo or neurogenic inflammatory responses activated by capsaicin, I-RTX behaves as an ultra potent antagonist at the native rat and guinea pig TRPV1 and at the recombinant human TRPV1.

The aim of the present study was to investigate whether I-RTX can reduce capsaicin and citric acid induced cough in guinea pigs and to compare its potency with that of CPZ.

METHODS

Animals

Male Dunkin-Hartley guinea pigs (250−350 g, Pampalonii, Italy) were acclimatised in cages at a mean (SD) temperature of 24 (0.5) °C for 1 week after delivery and were allowed free access to water and standard rodent diet (Morini, Italy). All experiments complied with the national guidelines and were approved by the regional ethics committee.

Experimental set up

After acclimatisation to laboratory conditions, animals were individually placed in a transparent perspex box (20×10×10 cm, Vetrotecnica, Italy) ventilated with a constant airflow of 400 ml/min. The tussive agents (citric acid 0.25 M and capsaicin 30 μM) were nebulised via a mini-ultrasonic nebuliser (Ugo Basile, Italy). The particle size produced had an aerodynamic mass median diameter of 0.9 μm and the output of the nebuliser was 0.4 ml/min. The appearance of cough was detected by means of a tie clip microphone (Sony, Japan) and confirmed by the characteristic posture of the animal. The cough sounds were recorded and digitally stored. The number of elicited cough efforts was subsequently counted by a blinded observer.

Study protocols

All experiments were carried out at the same time of day starting at 09.00 hours. The guinea pigs were exposed to aerosols of either capsaicin or citric acid for 10 minutes to elicit cough. To evaluate the effects of aerosolised I-RTX on experimentally induced cough, guinea pigs inhaled various concentrations of I-RTX corresponding to 0.3, 1, and 3 μM in capsaicin trials and 0.1, 0.3, and 1 μM in citric acid trials. To evaluate the effects of aerosolised CPZ on capsaicin and citric acid induced cough, guinea pigs inhaled various concentrations of CPZ.
concentrations of CPZ (30, 100, and 300 μM) before exposure
to each tussigenic agent. Both the I-RTX and CPZ inhalation
times were set at 10 minutes. In experiments aimed at
evaluating the effects of I-RTX injected intraperitoneally
(ip) on induced cough, guinea pigs were administered
0.03, 0.1, and 0.3 μmol/kg 1-RTX 15 minutes before both
capsaicin and citric acid inhalation. To determine the effects
of ip CPZ on induced cough the animals were given 0.3, 1,
and 30 μmol/kg CPZ before capsaicin inhalation or 0.1, 0.3,
and 1 μmol/kg CPZ before inhalation of citric acid. In all
experiments the effects of ip or aerosol administration of the
I-RTX and CPZ vehicles (control conditions) on induced
cough were also determined. Each animal received only one
dose of antagonist.

To detect any non-specific inhibitory properties of ip I-RTX
on the cough reflex, the ability of hypertonic saline (7% sodium chloride, 1.2 M) to induce cough was also investig-
gated. Hypertonic saline was administered for 10 minutes by
aerosol 15 minutes after administration of the antagonist or
its vehicle.

Drugs
Agents were obtained from the following companies: sodium
carbonate, citric acid, capsaicin, CPZ, (Sigma, Italy); I-RTX,
(Tocris, UK). The stock concentrations of capsaicin (10 mM)
and CPZ (10 mM) were prepared in 100% ethanol. The stock
concentration of I-RTX (1 mM) was prepared in 100%
DMSO.

Data analysis
Values are presented as mean (SE). Comparisons between
groups were made by one way analysis of variance (ANOVA)
and the Student’s t test or the Bonferroni t test when
appropriate. A p value of <0.05 was considered significant.
A minimum of eight guinea pigs was used to test the effect of
the vehicle or of each single dose of the drugs. The inhibitory
potency of I-RTX and CPZ was compared using the dose of
antagonist that produces 50% inhibition (ED50).

RESULTS
Independent of the route of administration, pretreatment
with I-RTX and CPZ caused no obvious cough response in
awake, freely moving guinea pigs. In no instance were
inhalations of aerosolised CPZ or I-RTX followed by the
appearance of even a few cough efforts. In contrast,
inhalation of both capsaicin and citric acid (alone or in the
presence of the CPZ or I-RTX vehicles) consistently caused a
brisk tussive response in all animals tested.

The effects of pretreatment with 1-RTX and CPZ on
experimentally induced cough are shown in figs 1 and 2.
Compared with control conditions, ip and aerosol adminis-
tration of I-RTX and CPZ consistently reduced the number of
coughs provoked by inhalation of both capsaicin and citric
acid (p<0.05 in both sets of experiments). Furthermore, in
all the experiments the inhibitory effects of 1-RTX and CPZ on
induced cough were dose dependent. Table 1 shows the ED50
values calculated after completion of the experiments. 1-RTX
administered either by aerosol or ip was significantly more
potent than CPZ. Inhalation of hypertonic saline (7% sodium
chloride, 1.2 M) provoked a significant increase in the
number of coughs (fig 3). Pretreatment with the highest
dose of I-RTX (0.3 μmol/kg ip) used in this study did not
affect the number of coughs induced by hypertonic saline
inhalation.

DISCUSSION
The results show that intraperitoneal and aerosol adminis-
tration of I-RTX is effective in reducing the number of
evoked by inhalation of both capsaicin and citric acid in
guinea pigs. A previous investigation showed that 1-RTX
is a potent compound in antagonising capsaicin induced
Antitussive activity of iodo-resiniferatoxin

We suggest that the inhibitory effect of I-RTX on capsaicin induced cough in guinea pigs is also caused by its ability to antagonise the activation of TRPV1. It should be emphasised, however, that the pharmacological characterisation of I-RTX as a TRPV1 antagonist is far from complete. Previous evidence indicated that I-RTX is a powerful antagonist at TRPV1 in vitro and in vivo.19 20 However, later studies confirmed the high potency of I-RTX in vitro but not in vivo in capsaicin induced paw flinching in rats.21 In this latter study I-RTX produced some excitatory effects at the highest doses. Likewise, minor excitation was produced by I-RTX when injected intraperitoneally to induce the writhing response.20 It is possible that a minor agonistic component is retained by I-RTX and is unmasked when the drug is administered locally in a circumscribed environment (intraperitoneally, subcutaneously), but not following systemic administration.19 20 Thus, the possibility that I-RTX retains some agonistic activity and that an agonist-dependent desensitising effect on sensory nerves contributes to its sensory neuron blocking activity cannot be completely discounted. In contrast, in none of our experiments did I-RTX cause a tussive response. One possible explanation for this is that inhalation of the drug causes widespread diffusion of the molecules into the airways so that its minor (if any) agonistic activity is minimised while its antagonist potential is maintained.

Citric acid is thought to induce cough by its ability to donate protons that are powerful stimulants of different subpopulations of primary sensory neurons including the rapidly adapting receptors (RARs)—for example, the subtype that plays a major role in the tussive response.22 Protons may stimulate neurons with C and A-δ fibres by activating different channels, two of which have been well characterised: (1) the TRPV14 and (2) the acid sensing ion channels (ASICs).23 It has been suggested that RARs do not express TRPV1,22 which suggests that ASIC ought to be responsible for the cough induced by citric acid. However, a previous study showed that CPZ was able to inhibit citric acid induced cough.24 Since CPZ is not a high affinity antagonist of TRPV1, this finding may raise doubts as to its selectivity in inhibiting the cough response. Nevertheless, the findings of Laloo et al25 suggested that TRPV1 rather than ASIC is responsible for citric acid induced cough. The present findings strongly support this view as I-RTX, a TRPV1 antagonist chemically unrelated to CPZ, is also able to inhibit citric acid induced cough effectively. However, a small component of the cough response to citric acid remained even at the highest doses of both I-RTX and CPZ. This residual response might be due to stimulation by citric acid of a non-TRPV1 channel such as the ASIC. Alternatively, it may still be caused by TRPV1 activation due to partial blockade of the channel by the two antagonists.

Capsaicin induced cough was also not completely inhibited by I-RTX or CPZ. Since it is presumed that the cough response induced by capsaicin is entirely mediated by TRPV1, it is likely that the incomplete abolition of citric acid induced cough by CPZ and I-RTX is caused by the partial blockade of TRPV1 at the doses of the antagonists used in the present study. An alternative explanation is that splice variants such as those described in the rat24 25 also occur in guinea pigs, and these variants could exhibit different affinities for different agonists. In addition, although the expression in recombinant systems suggests that the channel monomer is sufficient for the pore function of TRPV1, there is evidence that TRPV1 is capable of forming a specific ternary complex with phospholipase C and the neurotrophin trkA receptor.26 These differences in assembling the channel may be tissue specific and may result in different affinities of antagonists.

Finally, recent electrophysiological observations27 in single airway neurons of guinea pigs show that nerve activation following exposure to acid occurs in two different ways: a slowly inactivating mechanism present in C fibres which is TRPV1 dependent as it is blocked by CPZ and I-RTX, and a rapidly inactivating mechanism present in A-δ fibres which acts independently of TRPV1. The pharmacological observation that vanilloid receptor antagonism (present findings) inhibits most of the tussive response to citric acid28 leads to the surprising conclusion that the TRPV1 resistant pathway has a minor role in acid induced cough in guinea pigs.

Another important observation of the present study is that I-RTX was several times more potent than CPZ in reducing the cough response to capsaicin and citric acid. In vitro assays in different mammalian species have shown that I-RTX has a very high potency towards TRPV1, being 100–1000 times more potent than CPZ.19 20 This striking difference in vitro between I-RTX and CPZ has been confirmed in vivo in the mouse and rat where neurogenic plasma extravasation and nociceptive responses such as the writhing test20 or the capsaicin pain test have been studied.29 The finding that

Table 1  ED50 values of iodo-resiniferatoxin and capsazepine in inhibiting cough induced by capsaicin and citric acid in guinea pigs

<table>
<thead>
<tr>
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<th>Iodo-resiniferatoxin (nM)</th>
<th>Capsazepine (nmol/kg)</th>
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<tbody>
<tr>
<td>Capsaicin</td>
<td>284 (92)*</td>
<td>29 (9)*</td>
</tr>
<tr>
<td>Citric acid</td>
<td>84 (19)*</td>
<td>16 (4)*</td>
</tr>
</tbody>
</table>

Data are mean (SE) of at least eight experiments. *p<0.01 vs capsaicin.
I-RTX was 10–100 times more potent than CPZ in inhibiting experimentally induced cough is therefore in keeping with the results of previous in vivo investigations. The observation that both CPZ and I-RTX were more effective (2–5-fold) in inhibiting cough induced by citric acid than cough induced by capsaicin further supports the view that the citric acid cough is sensitive to TRPV1 antagonism.

The overall importance of TRPV1 in human cough is not fully understood. Available evidence suggests that TRPV1 is of considerable importance because stimuli that activate these channels are powerful tussive agents. In disease states the sensitivity of TRPV1 may be upregulated by proinflammatory mediators including bradykinin and nerve growth factor which have an important role in cough and asthma. The threshold dose of capsaicin to induce cough in patients with asthma and COPD is lowered, and patients with gastro-oesophageal reflux often suffer from cough. A high affinity and selective TRPV1 antagonist is needed to show the pathophysiological role of TRPV1 in the cough experienced by these different groups of patients.

In conclusion, we have shown that I-RTX, a novel and ultra potent antagonist at the native rat and guinea pig TRPV1 and at the recombinant human TRPV1, inhibits cough mediated by agents possibly acting via TRPV1 stimulation. I-RTX can therefore be considered as an exemplar for the design of novel antitussive agents.

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