Telenzepine is at least 25 times more potent than pirenzepine – a dose response and comparative secretory study in man

W LONDON, V LONDON, A MEIERL, AND U VODERHOLZER

From the Chirurgische und Medizinische Kliniken Innenstadt, University of Munich, Munich, West Germany

SUMMARY Telenzepine is an analogue of pirenzepine with a higher potency and similar selectivity for M₂-receptors in animals. In this placebo controlled, double blind, randomised study mean peptone stimulated gastric acid secretion (X±SEM) of 10 male healthy subjects (58±6 mmol H⁻³ h for placebo) was significantly and dose dependently inhibited by oral telenzepine (2 mg: 31±5, 3 mg: 23±5, 5 mg: 21±4 mmol H⁻³ h). Telenzepine 3 and 5 mg were significantly stronger than pirenzepine 50 mg orally (37±8 mmol H⁻³ h). Mean percentage acid inhibition was 37% for pirenzepine, and 48, 61, and 64% for 2, 3, and 5 mg telenzepine, respectively. Basal and peptone stimulated gastrin release was unaffected. Mean salivary output per three hours declined moderately from 156±45 g (placebo) to 136±45 g with pirenzepine and significantly to 88±28 g, 95±39 g and 39±13 g with telenzepine 2, 3, and 5 mg, respectively. There was a parallel effect on Na⁺, K⁺, Ca²⁺ and amylase output in saliva. Near point vision was not altered by either drug. Pulse rates were lowered by both substances. Complaints of dry mouth were more frequent with telenzepine 5 mg. On a molar basis telenzepine proved to be a 25 and 50 times more potent inhibitor of gastric and salivary secretion, respectively.

Telenzepine is a new analogue of pirenzepine having an altered tricyclic structure and an unchanged side chain (Fig. 1). In animal studies, telenzepine was four to 10 times more potent than pirenzepine inhibiting gastric acid secretion.¹ It healed experimentally induced gastroduodenal ulcers more effectively and at lower doses than pirenzepine.² Both compounds exhibited a similar selectivity profile differing from that of atropine.¹ Therefore, telenzepine was considered to be a new M₂-receptor antagonist, although receptor binding studies using telenzepine are lacking.

Pirenzepine has more selective inhibitory properties on oxyntic gastric glands and less antimuscarinic side effects than conventional antimuscarinics like atropine.³⁻⁴ In an oral dose of 100 to 150 mg daily pirenzepine proved to be superior to placebo and as effective as cimetidine in the treatment of patients with duodenal ulcer.⁵ Because of more pronounced antisecretory properties of telenzepine in animals it was of interest to investigate its relationship of antimuscarinic effects and side effects in man. Therefore, we undertook this placebo controlled, double blind and randomised study in which the dose response of telenzepine 2, 3, and 5 mg orally was compared with the standard oral dose of pirenzepine 50 mg in 10 healthy male subjects. Peptone stimulated gastric acid secretion, gastrin release, spontaneous salivation, near point vision and peripheral pulse rates were synchronously measured.

Methods

SUBJECTS

Ten healthy male subjects with a median age of 25 years (range 21–31 years), a median body weight of 75 kg (range 64–90 kg) and a median height of 185 cm (range 180–190 cm) participated in this trial. They had no abnormal physical findings including ECG

Address for correspondence: Prof Dr Walter London, Chirurgische Klinik Innenstadt und Chirurgische Poliklinik, University of Munich, Nussbaumstrasse 20, D-8000 Muenchen 2, West Germany.

Received for publication 11 November, 1986.
Telenzepine is at least 25 times more potent than pirenzepine

and no abnormal values of haematology, biochemistry, and urine analysis screen within 14 days before the first experiment. Criteria for exclusion were significant clinical illness and administration of investigational drug within the preceding four weeks, need for concomitant medication, conditions which could modify the absorption of the study medication, history of allergy, cardiac, renal, hepatic or significant gastrointestinal disease, history of drug addiction and excessive alcohol consumption. Each subject underwent a follow up laboratory screen not more than three days after the last experiment. All subjects gave written consent to participate in this study after full explanation by the investigator. The protocol of the study dated 20 July, 1984 was approved by the Ethical Committee of the Medical Faculty of the University of Munich on 19 October, 1984. The trial was conducted according to the Declarations of Helsinki and Tokyo.

Each subject participated in five experiments in which 2, 3, and 5 mg telenzepine, 50 mg pirenzepine and placebo were administered orally in the form of identical tablets (provided by Byk Gulden Company, D-7750 Konstanz, West Germany). The study was performed double blind and according to a predetermined randomisation code. The intervals between any two experiments were not less than four days, but not more than two weeks. Subjects were requested not to eat or to drink anything after 10 pm the previous evening. Alcohol intake was prohibited the evening before.

**EXPERIMENTAL PROCEDURE**

Each experiment started at 7 am. An iv cannula for collection of blood samples was inserted in a forearm vein and secured. Isotonic saline was infused at a rate of 40 ml/h to keep the cannula patent. Blood for determination of gastrin and drug concentrations was taken before (−5) and 45, 90, 120, 150, 180, 210, 240, and 270 minutes after drug intake. At zero minutes tablets were swallowed with 50 ml water. Eighty minutes after drug administration a double lumen gastric tube (Levine-type Ch 16) was placed in the stomach so that the tip of the tube was positioned in the fundus corpus area (in average 58·5 cm (range 56–61 cm) below front teeth – that is, about 10 cm below the cardia. One hundred minutes after medication the gastric secretion test began with an intragastric instillation of 300 ml 10% aqueous peptone solution (pH 5·5). Acid output was determined over a period of three hours by continuous titration according to Fordtran and Walsh using 1 N sodium hydroxide for automatic titration to an endpoint of pH 5·5. During the test the subjects were lying in a left lateral position. To maintain a constant speed of circulation of 250–280 ml/min, smaller volumes of peptone solution were substituted throughout the test. At the end of each gastric secretion test the stomach was emptied by suction. Details of our method have already been described.

During each experiment the saliva of the subjects was collected by a standardised and continuous suction using a perfusor system (B Braun Melsungen AG, Melsungen/FRG, type 371 102) in a reverse direction. The saliva was aspirated through a small standardised tube (original perfusor tube no 872 296/0) positioned in the anterior oral cavity into a 50 ml syringe to avoid evaporation. The speed of suction was adapted to the individual flow rate; on average the speed was 30–60 ml per 30 minutes. The subjects were instructed neither to swallow nor to spit out saliva. Salivary output was measured in 30 minutes intervals by weighing. The concentrations of sodium and potassium were determined by flame photometry, calcium by colorimetric estimation using o- cresolphthalein complexone as indicator and α-amylase by a direct spectrophotometric test using p-nitrophenyl α-maltoside as substrate. Furthermore, accommodation was monitored by determining near point vision at definite intervals during the test; under standardised light conditions a Rodenstock accommodometer was used. We calculated the mean of the measurements of accommodation before medication in all five intraindividual comparisons; this was defined as zero point. The values after drug intake are given as difference from zero point. Peripheral pulse rates were recorded in a standardised manner.

Blood samples for determination of gastrin and drug concentrations were placed on ice. After clotting and centrifugation (at 4°C, 4000 g) the serum phase was separated and stored at −20°C. Serum gastrin concentrations were measured by a sensitive radioimmunoassay using the specific antibody 4562 (kindly provided by Professor J F Rehfeld, Copenhagen, Denmark). Serum samples were analysed for concentrations of telenzepine using a gaschromatography-mass spectrometry assay, while pirenzepine was determined by radioimmunoassay.

![Pirenzepine and Telenzepine](image)

**Fig. 1** Structural formula of pirenzepine and its analogue telenzepine having an altered tricyclic structure and an unchanged side-chain.
Mean drug serum concentrations (\( \bar{X} \pm SEM \)) of 10 healthy male subjects after oral intake of 50 mg pirenzepine (PIR), 3 and 5 mg telenzepine (TEL) at time zero. Serum concentrations of 2 mg telenzepine were not measured, they would have been at the lower detection limit. In this diagram peptone stimulation is shown to illustrate the relationship of drug concentrations and experimental phase.

**Statistical Analysis**

Statistical evaluation was made by using two sided Wilcoxon's matched-pairs rank test. Only values of \( p \leq 0.01 \) were considered significant. Results were expressed in the conventional way as mean± standard error or the mean (\( \bar{X} \pm SEM \)).

**Results**

Figure 2 illustrates the mean drug serum concentrations of 50 mg pirenzepine as well as those of 3 and 5 mg telenzepine. Serum concentrations of 2 mg telenzepine were not measured because the values would have been at the lower detection limit of the method used. Pirenzepine concentrations were in the lower normal range after a standard dose of 50 mg pirenzepine orally.

Mean values of peptone stimulated gastric acid secretion per 15 minutes are shown in Figure 3. There were significant reductions of peptone stimulated acid output after 50 mg pirenzepine as well as after the three doses of telenzepine. Telenzepine proved to be more effective than 50 mg pirenzepine and reduced acid output in a dose dependent manner. Synchronously measured basal and peptone stimulated serum gastrin concentrations were not significantly altered by either drug.

Peptone stimulated acid output per three hours (\( \bar{X} \pm SEM \)) was significantly reduced (\( p \leq 0.01 \)) from 58.1±5.6 mmol H\(^+\) (placebo) to 36.8±7.6 mmol H\(^+\) by 50 mg pirenzepine, to 30.5±5.3 mmol H\(^+\) by 2 mg telenzepine, 22.8±4.5 mmol H\(^+\) by 3 mg telenzepine and to 21.2±3.9 mmol H\(^+\) by 5 mg telenzepine. The mean percentage inhibition was 37% for pirenzepine and 48, 61, and 64% for 2, 3, and 5 mg telenzepine, respectively. The effect of 3 and 5 mg telenzepine was significantly stronger than that of 2 mg telenzepine and of 50 mg pirenzepine (\( p \leq 0.01 \)). On a molar basis telenzepine proved to be about 25 times more potent than pirenzepine to inhibit acid output. Mean peptone volumes which had to be added during the secretory studies did not show significant differences within the test with pirenzepine or telenzepine medication.

Histograms of salivary output per 30 minutes are given in Figure 4. Fifty milligrams pirenzepine had only a minor effect on reducing salivation, whereas the effect of 2, 3, and 5 mg telenzepine was stronger being significant at different time intervals. The most dramatic effect was observed with 5 mg telenzepine. Total saliva per three hours (\( \bar{X} \pm SEM \)) was reduced from 156±45 g (placebo) to 136±45 g by pirenzepine and to 88±28 g, 95±39 g, and 39±13 g by 2, 3, and 5 mg telenzepine, respectively. Mean percentage inhibition of salivation was 13% for 50 mg pirenzepine and 44, 39, and 75% for 2, 3, and 5 mg telenzepine, respectively. The inhibition of salivation by the three telenzepine dosages was significant (\( p \leq 0.01 \)), but not strongly dose dependent. Only the difference between the inhibitory effect of 5 mg proved to be significant (\( p \leq 0.01 \)) versus 50 mg pirenzepine. On a molar basis telenzepine was more than 50 times more potent than pirenzepine in reducing salivation. There was a statistically obvious correlation between acid inhibition and reduction of salivation (\( r=0.858, p \leq 0.05 \)).

Table 1 contains mean salivary output of sodium, potassium, calcium, and amylase per three hours. Whereas 50 mg pirenzepine had negligible effects, more pronounced reductions were documented after telenzepine – being significant (\( p \leq 0.01 \)) for sodium after 3 mg as well as for sodium, potassium and calcium after 5 mg.

When monitoring near point vision (Fig. 5) we

<table>
<thead>
<tr>
<th>Placebo</th>
<th>50 mg PIR</th>
<th>2 mg TEL</th>
<th>3 mg TEL</th>
<th>5 mg TEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium</td>
<td>3.2</td>
<td>2.6</td>
<td>1.3</td>
<td>1.7*</td>
</tr>
<tr>
<td>(mmol/3h) ±1.7</td>
<td>±1.5</td>
<td>±0.47</td>
<td>±0.94</td>
<td>±0.22</td>
</tr>
<tr>
<td>Potassium</td>
<td>2.5</td>
<td>2.4</td>
<td>1.6</td>
<td>1.7</td>
</tr>
<tr>
<td>(mmol/3h) ±0.80</td>
<td>±0.80</td>
<td>±0.49</td>
<td>±0.65</td>
<td>±0.26</td>
</tr>
<tr>
<td>Calcium</td>
<td>0.20</td>
<td>0.20</td>
<td>0.12</td>
<td>0.13</td>
</tr>
<tr>
<td>(mmol/3h) ±0.049</td>
<td>±0.054</td>
<td>±0.031</td>
<td>±0.041</td>
<td>±0.031</td>
</tr>
<tr>
<td>(IU/3h)</td>
<td>±5619</td>
<td>±6088</td>
<td>±2180</td>
<td>±4806</td>
</tr>
</tbody>
</table>

*Significant (\( p \leq 0.01 \)) versus placebo.
Telenzepine is at least 25 times more potent than pirenzepine.

Acid output (mmol H⁺/15 min) was measured during peptone stimulation (10⁹/ₒ pH 5.5) in healthy male subjects. The graph shows the mean peptone stimulated gastric acid secretion per 15 min and synchronously measured serum gastrin release (X ± SEM) of 10 healthy male subjects. Acid inhibitory effect of pirenzepine (PIR) and telenzepine (TEL) was significant (p < 0.01) within 30 min after the start of peptone stimulation.

Peripheral pulse rates (Fig. 6) were lowered by either drug; the effects of 5 mg telenzepine and 50 mg pirenzepine were significant (p < 0.01) at 90 minutes and at 125 and 185 minutes postdosing, respectively. Table 2 summarises symptoms which the subjects were aware of during the double blind studies. Dry mouth occurred most frequently after 5 mg telenzepine. Four of seven subjects receiving 5 mg and two of three receiving 3 mg telenzepine complained of dry mouth up to the afternoon or evening of the test day. In all other subjects who had dry mouth it occurred during an interval of 50 up to 250 minutes after drug intake. Dry
eyes were specified by one subject during all five secretory tests and by two others after 2 and 5 mg telenzepine. Blurred vision was short lasting and noticed only once after pirenzepine application. Loose stools were reported during the afternoon or the evening of the test day.

Discussion

In this placebo controlled, double blind and randomised study telenzepine inhibited dose dependently peptone stimulated gastric acid secretion of healthy male subjects. On a molar basis it proved to be about 25 times more potent than pirenzepine. Our result corresponds with data published by others who calculated a comparative potency. Eltze et al. found in in vitro and in vivo experiments in animals a factor of four to 10. Häcki et al. estimated a factor of more than 10 in basal and pentagastrin stimulated gastric acid secretion in man. Müller et al. using sham feeding in man described a factor of 10 to 30. The higher potency of telenzepine might be partly a consequence of a better bioavailability which has been proven recently to be on average 56% when a single oral dose was administered to healthy subjects (personal communication). Corresponding data for pirenzepine are 20 to 30%. In this study the mean pirenzepine serum concentrations were rather low. This might be the reason of a comparatively low antisecretory effect of pirenzepine. A correlation between plasma concentrations of pirenzepine and its inhibitory effect on peptone stimulated gastric acid secretion has been shown recently in man.

The method of peptone stimulation is suitable for pharmacodynamic studies. First, it has been shown to have a good reproducibility, being essential for...
Telenzepine is at least 25 times more potent than pirenzepine

![Graph showing mean peripheral pulse rates](image)

**Fig. 6** Mean peripheral pulse rates (± SEM) of 10 healthy male subjects receiving either pirenzepine (PIR) or telenzepine (TEL). Significant differences versus placebo (p ≤ 0.01) are indicated by asterisks for 5 mg telenzepine at 90 min and for 50 mg pirenzepine at 125 and 185 min.

Intraindividual comparisons, with a mean coefficient of variation of about 10% in man. In a model for food stimulated gastric acid secretion and, contrary to pentagastrin test, allows simultaneous measurements of endogenous gastrin release. In this study, oral telenzepine had no influence on basal and pepsine stimulated gastrin concentrations (Fig. 3). This is in agreement with findings on basal serum gastrin published by Hӓcki et al. In this respect, telenzepine shows the same behaviour as pirenzepine which does neither affect meal stimulated nor sham feeding induced gastrin release in man. Both differ from atropine which enhances significantly serum gastrin in man after food stimulation and sham feeding.

In agreement with our results, Hӓcki et al showed that 50 mg pirenzepine given orally has less effect on salivary output in man than 3 and 5 mg telenzepine which significantly and dose dependently inhibited salivation. Müller et al also stated that a more pronounced reduction of salivation after 5 mg telenzepine resulted in more frequent complaints of dry mouth, similar to our findings (Table 2). The parallel behaviour of a reduced output of electrolytes and amylase (Table 1) suggests that telenzepine mainly affects volume secretion of salivary glands. The correlation between inhibition of gastric acid secretion and salivation supports the assumption that telenzepine does not discriminate between muscarinic receptor sites at gastric and salivary glands.

Both drugs reduced heart rate; the effect of pirenzepine was more pronounced (Fig. 6). Significant reductions of heart rate have already been reported after parenteral or high oral doses of pirenzepine. Blurred vision as a symptom of impaired accommodation was only once noticed after pirenzepine, but not after telenzepine (Table 2). Near point vision was not significantly altered by either drug (Fig. 5). In this respect, it is worth noting that an increase of mean pirenzepine plasma concentration from 40 ng/ml (normal therapeutic level) to 105 ng/ml (high therapeutic level) results in a significant reduction of near point vision.

Loose stools were reported as symptom in similar

---

**Table 2. Symptoms specified by the subjects (n=10) during and/or after the secretory tests**

<table>
<thead>
<tr>
<th>Symptom</th>
<th>Placebo</th>
<th>50 mg Pir</th>
<th>2 mg Tel</th>
<th>3 mg Tel</th>
<th>5 mg Tel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry mouth</td>
<td>0</td>
<td>2</td>
<td>3</td>
<td>3</td>
<td>7*</td>
</tr>
<tr>
<td>Dry eyes</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Blurred vision</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Loose Stools</td>
<td>2</td>
<td>4</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Others†</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

* p ≤ 0.05 versus placebo (using Fisher's exact test); † such as tiredness, headache, and euphoria.
frequency following placebo as well as drug experiments (Table 2). They are probably caused by transpyloric losses of hyperosmotic peptone solution and regularly observed using this test procedure. Rare symptoms like tiredness, headache and euphoria are unspecific. Central side effects of pirenzepine are very unlikely, as it penetrates blood brain barrier only to a small extent. Corresponding studies in cerebrospinal fluid of man using telenzepine are lacking.

Recent data and our results with telenzepine have shown that structural alterations of the tricyclic structure of pirenzepine result in an increase of potency with regard to gastric acid inhibition in man. It would be interesting to study structure activity relationships of further pirenzepine analogues in order to develop even more potent and selective antimuscarinic drugs. In this connection, it should be mentioned that very recently another pirenzepine analogue with an altered side chain and an identical tricyclic structure (AF-DX 116) has been characterised by in vitro binding studies and by pharmacological studies in animals as a M2-receptor antagonist with cardioselective action.

In conclusion, telenzepine proved to be, on a molar basis, a 25 times more potent gastric antisecretagogue than pirenzepine under the conditions tested. Its closely related inhibition of salivation may limit the clinical application of higher doses of telenzepine as antulcer drug.

The authors like to thank Dr E Sturm, Research Laboratories of Byk Gulden Company, Konstanz/FRG, and Dr P Tanswell, Department of Biochemistry, Dr Karl Thomae GmbH, Biberach/FRG, for measuring drug serum concentrations. This study has been published as abstract no 30 in Trends Pharmacol Sci 1986; suppl February: 89.

References

21 El-Sabbagh HN, Bloom SR, Adrian TE, Prinz RA, Baron JH, Welbourne RB. The effect of pirenzepine on
Telenzepine is at least 25 times more potent than pirenzepine


