A pharmacodynamic evaluation of midazolam as an antiepileptic compound

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SUMMARY Midazolam is a water soluble 1,4 benzodiazepine which is suitable for intramuscular administration. It is currently used for pre-medication and the induction of anaesthesia. Its antiepileptic properties have been evaluated by studying its effect on interictal spikes on the EEG of six adult epileptic patients. The results indicate that intramuscular midazolam 15 mg is more effective than intramuscular diazepam 10 mg in abolishing interictal spikes and as effective as intravenous diazepam 20 mg five minutes after administration.

Midazolam is a new 1,4 benzodiazepine drug belonging to the group of 1,2 annelated benzodiazepines, which have in common a five membered heterocyclic ring fused on position 1,2 of the diazepine nucleus, and have a high affinity to benzodiazepine receptors. It is unique in the basicity of its molecule which permits the preparation of water soluble salts, giving a very stable aqueous injectable solution. It has a short duration of action due to rapid metabolic inactivation, and has been used successfully for the intravenous induction of anaesthesia.

Animal data have suggested that midazolam might be useful as an intravenous or intramuscular antiepileptic drug.

Ahmad et al reported that EEG spike counting could be used for the rapid assessment of new antiepileptic compounds given in single intravenous doses. Milligan et al modified the technique and successfully demonstrated the efficacy of rectal diazepam and intravenous phenytoin in suppressing interictal spikes. We have used this method to assess the comparative efficacy of midazolam and diazepam.

Methods

Six adult epileptic patients (four male, two female, age range 32–57 years), resident at the Chalfont Centre for Epilepsy, gave their signed consent to take part in this study. Their routine drug therapy, which consisted of carbamazepine (one subject), carbamazepine and primidone (one subject), carbamazepine and phenytoin (two subjects), car-

bamazepine and sodium valproate (one subject) and phenytoin and sodium valproate (one subject), was kept unchanged throughout. None suffered from severe seizures requiring the use of benzodiazepines for at least three weeks before and during the study. The subjects were selected on the basis of having frequent spontaneous interictal spikes, seen as isolated spikes, polyspikes and spike and wave discharges, in their routine EEGs. All the intramuscular injections were given rapidly in the upper outer quadrant of the right buttock, while the intravenous injections were given at a rate of 0·5 ml/min into an antecubital vein by one experimenter (SJ).

The comparative evaluation of midazolam and diazepam was carried out in three parts.

Part 1

In this study the effects of intramuscular midazolam 10 mg and 15 mg, intravenous diazepam 10 mg and 20 mg and placebo were compared. Each patient received randomly the following treatments separated by intervals of 10 days:

- T1: intravenous normal saline + intramuscular normal saline
- T2: intravenous normal saline + intramuscular midazolam 15 mg
- T3: intravenous normal saline + intramuscular midazolam 10 mg
- T4: intravenous diazepam 10 mg + intramuscular normal saline
- T5: intravenous diazepam 20 mg + intramuscular normal saline

A 10 minute baseline EEG was recorded with the subjects’ eyes closed immediately before drug administration and this was used as a control. Five minutes after drug administration, nine subsequent 10 minute recordings were made every 20 minutes over a 3 hour period. EEG recordings were achieved using an eight channel SLE (Galileo Model E88) electroencephalograph recorder and bipolar electrodes positioned according to the 10–20 system. All the recordings...
were made from the lateral ring of electrodes. The EEGs were then code numbered and subsequently analysed blind and in random order using the method described by Ahmad et al. A spike was defined as a waveform distinguished from background activity and having a duration less than 70 ms (Cooper et al.). Sharp waves were included in the counting where these were clearly of epileptic origin, that is accompanied by spikes or sharp waves in adjacent channels or where they were phase reversing. In general, the duration of sharp waves included did not exceed 150 ms (accepted range 70–200 ms). Spike counting was confined to the one EEG channel in which the waves were most clearly defined. The total number of spikes in each 10 minute recording was compared for different treatments using a computerised three-way analysis of variance.

**Part 2**

The same EEG assessment and analysis procedures were used in the same six subjects to compare directly the effects of intramuscular midazolam 15 mg (T6) and intramuscular diazepam 10 mg (T7) over a 3 hour period. No placebo was used as the effectiveness of intramuscular midazolam against placebo had already been demonstrated in Part 1.

**Part 3**

As the timing of the first EEG recording was set at 5 minutes after injection this did not allow adequate differentiation of the speed of onset of action between the intravenous and intramuscular preparations. As it was anticipated that a benzodiazepine which could be given intramuscularly might be of use in the emergency treatment of seizures, the immediate effects of intramuscular midazolam 15 mg and intravenous normal saline (T8) and intramuscular normal saline and intravenous diazepam 20 mg (T9) were compared in the same six subjects. The EEG was recorded for 10 minutes prior to and after parental injection. The EEGs were coded and analysed later in a blind fashion. The number of spikes in each minute was counted and the results analysed by computerised analysis of variance.

**Results**

**Part 1**

All the patients fell asleep 1½–3 minutes following midazolam 15 mg and diazepam 20 mg, 2–4 minutes following midazolam 10 mg and 3–4 minutes following diazepam 10 mg. All the subjects fell asleep 30–45 minutes following placebo. All the patients were left undisturbed and woke up spontaneously at the end of the recording. No adverse effects were noted during the period of the study, apart from amnesia produced by midazolam.

The mean spike counts for different treatments at different times are given in table 1 and illustrated graphically in the figure.

All the main effects, that is difference between patients, time and treatments, were highly significant on an analysis of variance (p < 0.01).

In addition, the two-way interactions were all highly significant, that is the patients had significantly different time courses for their drug effects and significantly different degrees of response. There were also significant differences between the time courses of the effects of the treatments. Since the latter was significant, the main effects were re-tested using the pooled residual and two-way interaction sums of squares to make sure that the main effects were not due to the two-way interactions. Again, all main effects were still highly significant. Thus, there was a highly significant difference between drug treatments. An estimate of the standard error of a treatment mean at each time point was 7.353.

In order to detect which treatments were significantly different, two range tests on the five treatment means were carried out. The first was the least significant difference test at p < 0.05, and the

**Table 1**  
Mean number of spikes [± SD] in each 10 min EEG recording following treatments T1–T7

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>T4</th>
<th>T5</th>
<th>T6</th>
<th>T7</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-10</td>
<td>81.8 ± 23.6</td>
<td>83.5 ± 21.3</td>
<td>77.0 ± 29.2</td>
<td>78.3 ± 24.7</td>
<td>70.8 ± 13.1</td>
<td>136.5 ± 68.8</td>
<td>130.8 ± 61.1</td>
</tr>
<tr>
<td>[baseline]</td>
<td>78.5 ± 22.0</td>
<td>2.0 ± 3.6</td>
<td>8.2 ± 9.4</td>
<td>16.5 ± 14.8</td>
<td>2.8 ± 2.9</td>
<td>18.3 ± 18.2</td>
<td>75.0 ± 66.3</td>
</tr>
<tr>
<td>25-35</td>
<td>95.0 ± 15.8</td>
<td>4.3 ± 0.0</td>
<td>12.2 ± 16.5</td>
<td>10.5 ± 7.9</td>
<td>3.0 ± 3.2</td>
<td>16.5 ± 20.0</td>
<td>72.8 ± 64.6</td>
</tr>
<tr>
<td>45-55</td>
<td>78.3 ± 22.5</td>
<td>12.5 ± 12.9</td>
<td>15.5 ± 29.5</td>
<td>11.7 ± 14.0</td>
<td>6.7 ± 7.9</td>
<td>8.0 ± 9.0</td>
<td>54.2 ± 50.5</td>
</tr>
<tr>
<td>55-75</td>
<td>95.0 ± 15.8</td>
<td>15.0 ± 25.4</td>
<td>13.2 ± 16.7</td>
<td>6.5 ± 7.5</td>
<td>9.8 ± 11.6</td>
<td>63.7 ± 54.3</td>
<td></td>
</tr>
<tr>
<td>85-95</td>
<td>64.2 ± 16.3</td>
<td>16.7 ± 25.4</td>
<td>26.8 ± 25.8</td>
<td>8.7 ± 7.1</td>
<td>13.7 ± 14.8</td>
<td>70.0 ± 61.7</td>
<td></td>
</tr>
<tr>
<td>105-115</td>
<td>58.8 ± 12.3</td>
<td>13.0 ± 14.2</td>
<td>16.2 ± 20.6</td>
<td>31.8 ± 31.0</td>
<td>17.5 ± 17.5</td>
<td>19.2 ± 22.0</td>
<td>71.8 ± 61.2</td>
</tr>
<tr>
<td>125-135</td>
<td>53.2 ± 11.9</td>
<td>19.8 ± 26.5</td>
<td>16.5 ± 18.1</td>
<td>37.8 ± 32.3</td>
<td>12.2 ± 14.4</td>
<td>12.2 ± 17.0</td>
<td>76.7 ± 68.3</td>
</tr>
<tr>
<td>145-155</td>
<td>66.3 ± 20.7</td>
<td>20.3 ± 25.7</td>
<td>20.5 ± 18.2</td>
<td>39.3 ± 37.7</td>
<td>17.3 ± 12.8</td>
<td>16.3 ± 18.3</td>
<td>63.2 ± 52.0</td>
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<tr>
<td>165-175</td>
<td>86.5 ± 8.8</td>
<td>20.0 ± 24.3</td>
<td>19.2 ± 17.3</td>
<td>39.2 ± 35.2</td>
<td>17.2 ± 11.0</td>
<td>15.2 ± 16.7</td>
<td>64.8 ± 53.6</td>
</tr>
</tbody>
</table>

T1: intravenous normal saline + intramuscular normal saline.
T2: intravenous normal saline + intramuscular midazolam 15 mg.
T3: intravenous normal saline + intramuscular midazolam 10 mg.
T4: intravenous diazepam 10 mg + intramuscular normal saline.
T5: intravenous diazepam 20 mg + intramuscular normal saline.
T6: intramuscular midazolam 15 mg.
T7: intramuscular diazepam 10 mg.
second the Student Newman-Keuls test at \( p < 0.05 \), which avoids certain problems of over sensitivity in the least significant difference test. In this case, however, both tests produced the same result. Treatment 1 (placebo) was significantly different from all other treatments. Treatment 4 (intramuscular normal saline and intravenous diazepam 10mg) was also significantly different from all other treatments. All the other treatments, that is treatment 2 (intravenous normal saline and intramuscular midazolam 15mg), treatment 3 (intravenous normal saline and intramuscular midazolam 10mg) and treatment 5 (intramuscular normal saline and intravenous diazepam 20mg) did not differ.

**Part 2**
All the subjects slept following intramuscular midazolam. None of the subjects slept following intramuscular diazepam and all of them complained of moderate pain at the site of injection.

Table 1 shows the mean number of spikes following intramuscular midazolam 15 mg (T6) versus intramuscular diazepam 10 mg (T7). The analysis of variance showed significant interaction terms for both subject-drug and drug-time. The main effects were retested against the combined residual and interaction variances, and “F value” for the differences between drugs was still highly significant \( p < 0.001 \). This indicated the superiority of intramuscular midazolam 15 mg over intramuscular diazepam 10 mg in suppressing interictal spikes.

**Part 3**
Table 2 gives the mean number of spikes at each minute following intramuscular midazolam 15 mg + intravenous normal saline (T8) and intravenous diazepam 20 mg + intramuscular normal saline (T9). A quantitative analysis of the difference in rate of onset of drug action proved difficult from the data obtained. It involved fitting the data to some empirical model. A simple approach was adopted, assuming that the drug effect builds exponentially, so a plot of log response against time might be approximately linear up to the attainment of maximum response. The mean data were treated in this way because of presence of zeros in the individual patient data. The scatter of the diazepam data followed a log linear fit, but for the midazolam data a sigmoid curve.
was obtained, suggesting that the model is not entirely suitable and that delay of 2–3 min may occur before the response builds up at a rate similar to that of diazepam. These conclusions are qualitative and do not permit more complex numerical analysis.

Discussion

The assessment in humans of new compounds with antiepileptic potential has been dependent traditionally on clinical trials. However, attempts have been made recently to devise techniques which permit new drugs to be evaluated at an earlier stage but only when a large number of compounds have been assessed will it be possible to validate these methods as predictors of antiepileptic efficacy in patients. The technique of interictal spike counting, following a single IV or oral dose, has so far been used to examine proven antiepileptic drugs, namely rectal diazepam, phenytoin, and an experimental compound, lamotrigine, which is currently undergoing clinical trials and frusemide. Interestingly, intravenous frusemide which is not generally regarded as an antiepileptic drug significantly reduced interictal spikes and was subsequently shown to be an effective antiepileptic agent when given as add-on therapy. Sodium valproate, on the other hand, when given intravenously at a dose of 15 mg/kg had no effect on interictal spikes (Milligan, unpublished). This latter finding may reflect a different mode of action of valproate in suppressing seizure activity.

The results of this study show that intramuscular midazolam at doses of 10 mg and 15 mg produced a significant pharmacodynamic effect as measured by a reduction in spike counts in the EEG. This effect was comparable at 5 minutes after injection to that produced by intravenous diazepam 20 mg. Intramuscular midazolam 15 mg was, however, more effective than diazepam 10 mg given intramuscularly or intravenously.

It is highly unlikely that suppression of interictal spikes is only a non-specific effect due to sedation. In Part 1 of the study all subjects fell asleep after each treatment and although a non significant fall in spikes was seen after placebo the mean spike counts were still significantly lower after all active treatments. In Part 2 of the study subjects remained awake for 45 minutes after diazepam 10 mg intramuscularly but nevertheless a significant fall in spike counts was observed.

Although in the first part of this study intramuscular midazolam and intravenous diazepam appeared to be equally effective in abolishing interictal spikes, it should be noted that the first measurement was not made until 5 minutes after injection, and it is possible that intravenous diazepam produced a greater effect within this time. Similarly, the nature of the assessment of drug action, that is by counting EEG spikes, was unsuitable for determining the speed of onset of action of the intravenously administered drug. However, a qualitative assessment of the data obtained would indicate a delay in the onset of action of midazolam compared with diazepam of 2–3 minutes.

A drug which can be given intramuscularly and which effectively and rapidly suppresses interictal spikes may be of considerable clinical value in the emergency treatment of seizures. Although intravenous diazepam is generally regarded in the UK as the treatment of choice in this situation, it may be that intramuscular midazolam will have a comparable effect in suppressing seizures and the anticipated delay in onset of action of 2–3 minutes may well be acceptable in clinical situations where it is not possible for an intravenous injection to be given. The results of this study and preliminary clinical experience indicate that intramuscular midazolam should be evaluated further in a controlled trial against intravenous diazepam.

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References
