In vitro Evaluation of Tinidazole Bioadhesive Vaginal Gels
Zainab T. Salih*1

Abstract
Many trials were made to prepare Tinidazole 2% as bioadhesive vaginal gels using different gel bases including hydroxypropyl methyl cellulose (3 and 4% w/w), methylcellulose (3 and 4% w/w) and carboxymethyl cellulose (2 and 3% w/w). Swelling index of the polymers, pH, viscosity, bioadhesive force, and in-vitro drug release to the simulating vaginal fluid (S.V.F.) were investigated for all the prepared bioadhesive gels. The mechanism of drug release from the gel bases was also investigated. The results revealed that CMC 3% gave the highest viscosity and bioadhesive strength with the lowest release rate while lowest viscosity and bioadhesive force was obtained with HPMC gel base with the highest release rate. The mechanism of drug release was affected by the type of gel base. Fickian diffusion was obtained with all gel bases.

Key words: Tinidazole, bioadhesive polymers, gels

This medication1 patients tolerate gels better than vaginal inserts or ointments. This study was conducted to formulate TND in a suitable mucoadhesive vaginal gel through studying different variables affecting the physicochemical properties and the in vitro release of the drug from these preparations.

Materials and Methods

Materials
Tinidazole (TND) was supplied by Sigma chemical co., methylcellulose (MC) and carboxymethylcellulose (CMC) from BDH limite (HPMC) from Shin-Etsu chemical co.- Germany. All other materials used were of analytical grade.

Methods
Preparation of simulating vaginal fluid (S.V.F)
Simulating vaginal fluid (S.V.F) was prepared by dissolving 3.51gm of sodium chloride, 0.22gm of calcium chloride, 0.018gm of bovine serum albumin, 2 gm of lactic acid, 1gm of acetic acid, 0.16gm of glycerol, 0.4gm of urea and 5gm of glucose in 1 liter distilled water then the pH was adjusted at 4.2.

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Preparation of TND vaginal aqueous gel bases

The tested aqueous gel bases were prepared by dispersing the polymers powder of 3 and 4%(w/w) MC, 2 and 3%(w/w) CMC and 3 and 4%(w/w) HPMC in cold freshly distilled water with the aid of a high speed stirrer, until a solutions were complete. The products were refrigerated for hour before use to obtain a suitable consistency. TND gels were prepared by dispersing 2% drug in different gel bases as shown in table(1).

### Table 1: Composition of TND bioadhesive vaginal gel

<table>
<thead>
<tr>
<th>Materials</th>
<th>F1</th>
<th>F2</th>
<th>F3</th>
<th>F4</th>
<th>F5</th>
<th>F6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tinidazole</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Methylcellulose(Mc)</td>
<td>3</td>
<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Carboxymethylcellulose (CMC)</td>
<td>-</td>
<td>-</td>
<td>2</td>
<td>3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Hydroxypropylmethylcellulose (HPMC)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3</td>
<td>4</td>
<td>-</td>
</tr>
<tr>
<td>H C Q S</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

In vitro dissolution of TND from gel bases

The release study was carried out in a USP apparatus type I at 100 rpm, a basket of 2.5 cm in diameter was enclosed with a multifold filter paper (dialysis cell) in order to be filled with 1 gm of each base containing 2% w/w TND. After connecting to a stirrer motor, the dialysis cell was immersed to about 1 cm of its surface in 500 ml of simulating vaginal fluid pH4.2 at 37°C for 5 hours. Samples were withdrawn after 0.5, 1, 2, 3, 4 and 5 hours and replaced with an equal volume of the same fluid solution. The samples were analyzed for their drug content at its λmax 320nm.

Mechanism of TND release from gel bases

To understand the mechanism of the drug release, the release profiles from different gel bases, the % fraction drug released verses time profiles were plotted using linear fitted equations proposed by Koresmeyer - Peppas:

\[ F = \frac{Mt}{Mo} = kt^n \]  
\[ \ln F = \ln k + n \ln t \]

The equations express the fraction (F) release of drug from gel. Where Mt is the amount released at time t, Mo is the initial amount of drug, K is the rate constant and n is the releasing exponent value indicative the mechanism of drug release (if n is 0.5 or less (fickian) holds the release, while when n greater than 0.5 this indicate non-fickian (anomalous) releasing mechanism), n value could be result from the slope obtained by the drawing of ln. Fraction released verses ln. Time.

Viscosity measurement

A Brook Field digital viscometer with a suitable sample adaptor was used to measure the viscosity in centipoises of the bioadhesive prepared gel.

Determination of pH

The pH of the bioadhesive gels were determined by digital pH meter. One gram of gel was dissolved in 25 ml of distilled water and the electrode dipped into gel formulation for 30 minute. The measurements of pH of each system were replicated three times.
Determination of mucoadhesive force

The mucoadhesive potential of each system was determined by measuring the force required to detach the gel from sheep vaginal mucosal tissue by using a modified chemical balance. A section of vaginal mucosa was cut from the sheep's vaginal cavity and instantly fixed with mucosal side out onto each glass vial using a rubber band. The vials with vaginal mucosa were stored at 37°C for 5 minutes. Then next vial with a section of mucosa was connected to the balance in inverted position, while first vial was placed on a height adjustable pan. Fixed amount of sample of each gel system were placed onto the vaginal mucosa of first vial. Then the height of second vial was adjusted so that mucosal surfaces of both vials come in intimate contact. Five minutes contact time was given to ensure intimate contact between tissues and the sample. Then weight was kept rising in the pan until vials get detached. The bioadhesive forces as the detachment stress in dyne/cm², was determined from the minimal weights that detached the tissues from the surface of each gel using the following equation\(^{(19)}\):

\[
\text{Detachment stress (dyne/cm}^2) = \frac{m \times g}{A}
\]

Where, \(m\) = weight required for detachment of two vials in grams.
\(g\) = Acceleration due to gravity [980cm/s²].
\(A\) = area of tissue exposed.

The vaginal mucosa was changed for each measurement. Measurements were repeated three times for each of the gel system.

Results and Discussion

In order to fortify the adhesion of administered drug onto the mucosal surface, TND was formulated in a suitable mucoadhesive polymers, MC, CMC and HPMC bases to take a full advantageous of the contact time, the drug can be dispersed in the gel giving a concentration that is higher than corresponding to the solubility of the drug\(^{(20)}\), as well as it can be considered as a way for sustaining the release of the drugs from gel dosage form. MC, CMC and HPMC were used as a gel base because of their safety use for vaginal application as well as there compatibility with TND, with other vaginal secretion.

The effect of polymer type and concentration on the releasing of TND from the gel bases

Figures 1, 2 and 3 showed the amount of TND released from gel bases in S.V.F medium. It was found that the amount of TND released from gel bases were significantly influenced (\(p<0.05\)) by the type of the base and the concentration the polymer used\(^{(21)}\). In general it was found that highest amount of TND was released from CMC gel base and decreased with an increasing in polymer concentration\(^{(22)}\), also the drug release rate was decreased with the increase in the polymer concentration\(^{(23)}\).

Estimation of swelling capacity

The swelling index (capacity) for 1gm dry MC and CMC are 50% and 80% respectively. HPMC is soluble and non-swollen polymer at the concentration used. The high swelling % of CMC in S.V.F indicates that the swelling of this polymer takes place at a slower rate. The straight lines obtained from plotting of \(\frac{t}{v}\) (\(t\): time in hour, \(v\): volume of aqueous S.V.F absorbed per volume of dry polymer) against time \(t\) in hour as shown in figures (4and 5) were utilized to calculate the kinetic of swelling including initial rate of swelling (reciprocal of the intercept) and swelling equilibrium (reciprocal of slope)\(^{(24)}\), as shown in table 2.

<table>
<thead>
<tr>
<th>Polymer type</th>
<th>Swelling to equilibrium size (ml of fluid/ml of dry polymer)</th>
<th>Initial rate of swelling (ml of fluid solution/hr.ml of dry polymer)</th>
<th>Correlation coefficient (r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC 3%</td>
<td>1.797</td>
<td>74.626</td>
<td>0.99</td>
</tr>
<tr>
<td>MC 4%</td>
<td>1.496</td>
<td>12.987</td>
<td>0.9885</td>
</tr>
<tr>
<td>CMC 2%</td>
<td>1.951</td>
<td>11.454</td>
<td>0.9944</td>
</tr>
<tr>
<td>CMC 3%</td>
<td>1.500</td>
<td>14.619</td>
<td>0.995</td>
</tr>
</tbody>
</table>
It was found that formulation with higher swelling index retard the release of drugs more than those with lower swelling indices, this swelling also depends on the pH of the medium and the presence of electrolytes, where the S.V.F pH 4.2 has a lot of electrolytes content such as: sodium chloride, potassium hydroxide, calcium hydroxide (25). This swelling is important since its significantly decreases the releasing rate of the drug and increases in the amount of the drug release after five hours, as shown in table-3-. The reason behind this increasing in the amount released due to erosion of the gel as a result of severe hydration or relaxation of the polymer, so interchain intermolecular force will no longer be able to resist any external forces, once gel erodes, it breaks up into smaller and smaller particles, more surfaces will be exposed to the fresh swelling medium.
Swelling of a gel base is a necessary condition to increase the amount of drug release. Therefore, the swelling ability of CMC gel base is more than other swollen MC gel base after five hours. As a result, CMC gel base hold a Fickian release mechanism.

Table 3: The release rate, amount released and the releasing mechanism of the TND from gel bases

<table>
<thead>
<tr>
<th>Polymer type and concentration</th>
<th>n exponent release value</th>
<th>K % min releasing rate</th>
<th>amount release (mg/5hr) (mean±SD,n=3)</th>
<th>Mechanism of TND release</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC 3%</td>
<td>0.41</td>
<td>5.3974</td>
<td>7.05±0.14</td>
<td>Fickian</td>
</tr>
<tr>
<td>MC 4%</td>
<td>0.49</td>
<td>3.5744</td>
<td>4.466±0.13</td>
<td>Fickian</td>
</tr>
<tr>
<td>CMC 2%</td>
<td>0.509</td>
<td>4.1594</td>
<td>7 ±0.15</td>
<td>Fickian</td>
</tr>
<tr>
<td>CMC 3%</td>
<td>0.52</td>
<td>3.582</td>
<td>4.35±0.12</td>
<td>Fickian</td>
</tr>
<tr>
<td>HPMC 3%</td>
<td>0.29</td>
<td>6.1472</td>
<td>5.2±0.03</td>
<td>Fickian</td>
</tr>
<tr>
<td>HPMC 4%</td>
<td>0.5</td>
<td>3.1864</td>
<td>4.58±0.11</td>
<td>Fickian</td>
</tr>
</tbody>
</table>

Viscosity measurement

Viscosity is an important parameter for characterizing the gel as it affects the spreadibility, extrudability and release of drug. CMC gel base show the highest viscosity, and as the polymer concentration increase, the viscosity will be increased as shown in table 4. On the other hand, minimum level of viscosity was seen with F5 and F6 explain the rapid release of drug from this gel system and this due to non swelling property of HPMC polymer at these concentration.

Table 4: The viscosity and pH of the prepared TND bioadhesive vaginal gel

<table>
<thead>
<tr>
<th>Formula No.</th>
<th>Viscosity (cps)</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>1450</td>
<td>6.02</td>
</tr>
<tr>
<td>F2</td>
<td>3240</td>
<td>5.2</td>
</tr>
<tr>
<td>F3</td>
<td>1700</td>
<td>5.03</td>
</tr>
<tr>
<td>F4</td>
<td>6800</td>
<td>5.17</td>
</tr>
<tr>
<td>F5</td>
<td>14.5</td>
<td>5.02</td>
</tr>
<tr>
<td>F6</td>
<td>21.6</td>
<td>5.2</td>
</tr>
</tbody>
</table>

Determination of pH

The pH of gels were found to be within the range of 5-6.2 which is within the limit of semisolid specification and at this pH the gel will be non irritant to vagina. The physical appearance of the gel was evaluated with naked eye. The gel was smooth without any lumps and of uniform color, no color change /liquification /separation were observed.

Determination of mucoadhesive force

Scheme 1 indicates the vaginal bioadhesive properties of the prepared gels in sheep vagina, and the results showed that all vaginal bioadhesive strengths were found in the following order F2>F4>F3>F1>F6>F5. Indicating that (F2) and (F4) showed the highest bioadhesive properties.

Conclusion

Results of this study confirm that, the physical properties of the prepared gel were affected by the type and concentration of the polymer used in the preparation. The more sustained preparation with highest bioadhesive force and highest viscosity was obtained by the use of 3% CMC as a gel base. In addition to that the mechanism of drug release from tinidazole 3% CMC gel base was diffusion mixed with erosion.

References


5. BNF; British national formulary, British medical association and Royal Pharmaceutical society of Great Britain 2004; 47 ed.: 287, U.K.


