Expulsions following 1000 GyneFix insertions

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Abstract
Context. The GyneFix intra-uterine device has been used in our family planning service since 1997. One of the perceived advantages is its low expulsion rate, as reported by clinical trials.

Objective. To calculate expulsion rates in routine clinical use and to look at possible reasons for expulsion.

Design. Retrospective casenote analysis and opportunistic clinician consultation.

Setting. A city centre family planning clinic.

Participants. The first 1000 GyneFix insertions.

Main outcome measures. Parity of client, experience of clinician carrying out insertion, time from device insertion to expulsion.

Results. Overall expulsion rate was 7.6%. There was no significant difference in parity of clients experiencing expulsion. Most (4.7%) expulsions were early, occurring within 3 months of insertion. There was considerable variation in early expulsion rate from one clinician to another. Later expulsions also occurred, up to 28 months after insertion. Increasing experience of the inserting clinician led to lower rates of late expulsion. Unnoticed expulsion led to four unplanned pregnancies.

Conclusions: The GyneFix expulsion rate in our service is higher than quoted in clinical trials. Early expulsions may be related to insertion technique, representing insufficient implantation of the anchoring knot into the fundal myometrium. Late expulsions also occur, often many months after insertion; the reason for these is unclear. Users should be taught to check for the presence of the thread after each menstrual period and unnoticed expulsion should be confirmed by ultrasound and abdomino-pelvic plain X-ray.

Key message points
- Expulsion rate in our service has been found to be higher than quoted in clinical trials.
- Most expulsions occur within 3 months of insertion and may be due to insufficient implantation of the anchoring knot.
- Late expulsions also occur, often many months after insertion. The reason for late expulsions is unclear.
- As a routine, users should be taught to check for the presence of the thread after each menstrual period.
- Unnoticed expulsion should be confirmed by ultrasound and abdomino-pelvic plain X-ray.

Introduction
The frameless GyneFix intra-uterine device (IUD) has been available in the UK since 1997. It was developed to attempt to overcome side effects seen with framed IUDs such as expulsion, bleeding and pain, thought to be due to incompatibility between the uterine cavity and the frame of the device, particularly in nulliparous women. Clinical trials have confirmed that the GyneFix is effective and acceptable to nulliparous as well as parous women, with low expulsion rates, although to date there has not been a randomised trial comparing GyneFix to framed IUDs in nulliparous women.

Two randomised comparative studies using the Flexigard, an earlier model of the GyneFix (identical in all but insertion instrument), found the frameless device to have significantly higher rates of expulsion than the framed device, the TCu380A, in parous women. This was found to be due to shortcomings in the insertion instrument, which was subsequently modified, the resultant device being named ‘GyneFix’.

Studies using the GyneFix have been more encouraging. The 3-year expulsion rate in a non-comparative trial involving both nulliparous and parous women was 0.7%. A randomised comparative trial in parous women in China found the expulsion rate with the GyneFix to be significantly lower (3.0%) than that of the framed device, the T380A (7.4%), at 3 years. The authors state that most expulsions with GyneFix occur within 3 months of insertion, representing ‘insertion failures’ due to insufficient implantation of the anchoring knot into the fundal myometrium through lack of experience with the technique. A recent paper reported no expulsions in a non-comparative study using the device immediately post-abortion in 175 Chinese women, mostly parous. It is not stated which model of GyneFix, the IN (interval) or PT (post-termination) version, was used.

Liverpool was the first clinical site in the UK to routinely offer GyneFix and as such it is a continuing training centre. All insertions are carried out at our city-centre clinic. A protocol has been drawn up in our service to help clinicians decide who might benefit from using a GyneFix over a framed copper device. The GyneFix is available to all women considering IUD use, but is offered mainly to nulliparous women. Clinical studies using the GyneFix have been more encouraging.
been kept. Onto this is entered insertion date, parity (nulliparous meaning never pregnant or no pregnancy over 24 weeks), experience of the clinician carrying out insertion and subsequent events at follow-up, such as removal and expulsion. Clients are advised to re-attend in the event of problems. Clinic staff are regularly reminded to inform the first author of any removals, expulsions or other adverse events, so that these can be noted in the register and kept under review. Data were collected from both the register and by casenote review. Where a client underwent more than one GyneFix insertion, for example if she opted for re-insertion following expulsion, this was counted as two separate insertions and entered onto the register twice.

The clinician carrying out the insertion was classed as ‘inexperienced’ or ‘experienced’, depending on the total number of insertions he or she had carried out. The clinician is considered experienced in GyneFix insertion once he or she has inserted more than 10 devices.

Time in months from insertion to expulsion was noted so that expulsions could be classed as ‘early’, occurring within 3 months of insertion, or ‘late’, more than 3 months after insertion.

**Results**

One thousand devices were successfully inserted from February 1997 to April 2000, by 74 inexperienced and 10 experienced clinicians. Most (799) clients were nulliparous. The remainder (201) were parous.

There were 11 abandoned insertions, either through pain or inability to pass a uterine sound, or failure to anchor the device at the uterine fundus during insertion. Four of these subsequently underwent successful insertion. Only successful insertions have been included in the analysis.

Seventy-six devices are known to have been expelled. There may be further expulsions about which we are unaware, as follow-up is opportunistic. Data were collected up until August 2000. Figure 1 shows time from insertion to expulsion.

Most of the expulsions (47) were early, occurring within 3 months of insertion. The remaining 29 were late, occurring more than 3 months after insertion. Of these, 14 were between four and 6 months, eight between seven and 12 months and seven later than 12 months after insertion. The latest expulsion was 28 months after insertion.

<table>
<thead>
<tr>
<th>Insertions</th>
<th>Nulliparous (%)</th>
<th>Parous (%)</th>
<th>Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expulsions</td>
<td>799 (80)</td>
<td>201 (20)</td>
<td>1000 (100)</td>
</tr>
</tbody>
</table>

Table 1 shows the numbers of expulsions according to parity. The expulsion rate amongst nulliparous users was higher (8.0% vs. 6.0%), although this difference does not reach significance using chi-square testing.

**Experience of clinician carrying out insertion**

Three hundred and sixty-eight of the 1000 insertions were carried out by an inexperienced clinician (Table 2). Thirty-two of these (8.6%) were subsequently expelled. Forty-four (7.0%) of the remaining 632 devices inserted by experienced clinicians were expelled. This difference is not significant using chi-square testing.

**Early and late expulsions**

There was no significant difference in early expulsion rate between inexperienced and experienced clinicians using chi-square testing. The late expulsion rate was higher amongst inexperienced clinicians than experienced. This difference is significant using chi-square testing (p < 0.05).

Within the group of experienced clinicians, early expulsion rate ranged from zero to 14.3%, median 1.6%. For late expulsions, although actual numbers are small (between zero and four per experienced clinician), inter-individual expulsion rates did not vary to the same degree.

There were four pregnancies through unnoticed expulsion, two at 5 months, one at 13 and one at 20 months from insertion. Five women experienced two GyneFix expulsions (therefore appearing twice in Figure 1).
Discussion
The low rates of GyneFix expulsion seen in clinical trials have not been reproduced in routine use in our clinic setting. Expulsion rates may be higher than presented; incomplete follow-up may have been a source of bias if further expulsions occurred but were not reported.

Table 2: Early and late GyneFix expulsion rates for inexperienced and experienced clinicians

<table>
<thead>
<tr>
<th>Expulsion Type</th>
<th>Inexperienced</th>
<th>Experienced</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number</td>
<td>% (range)</td>
<td>Number</td>
</tr>
<tr>
<td>Total inserted</td>
<td>368</td>
<td>100</td>
<td>632</td>
</tr>
<tr>
<td>Early expulsions</td>
<td>16</td>
<td>4.3 (0 – 66.7)</td>
<td>31</td>
</tr>
<tr>
<td>Late expulsions</td>
<td>16</td>
<td>4.3 (0 – 33.3)</td>
<td>13</td>
</tr>
<tr>
<td>Total expulsions</td>
<td>32</td>
<td>8.6</td>
<td>44</td>
</tr>
</tbody>
</table>

Expulsion rates similar to ours have been observed in family planning clinics elsewhere in the UK (personal communication). In the majority of sites offering GyneFix in the UK, ultrasound scanning after insertion is not routinely performed unless the clinician has doubts about correct placement. The possibility of expulsion should be discussed with the client prior to insertion, along with a recommendation that she check for the presence of the thread after each menstrual period. Unnoticed expulsion should be confirmed by ultrasond and plain X-ray of the pelvis and abdomen, to ensure that the device has not perforated. Ongoing experience in the technique of GyneFix insertion is essential to retain expertise. Audit enables continued monitoring of expulsion rates.

In spite of the expulsions, the majority of our users are satisfied with the GyneFix and we continue to receive many requests for insertion. We feel that the GyneFix is a welcome addition to the contraceptive menu offered to our clients.

Conclusion
The GyneFix expulsion rate in our service is higher than quoted in clinical trials. Expulsion appears unrelated to the parity of the user. Early expulsions may be related to insertion technique, representing insufficient implantation of the anchoring knot into the fundal myometrium. Late expulsions also occur, often many months after insertion; the reason for these is unclear. Users should be taught to check for the presence of the thread after each menstrual period and unnoticed expulsion should be confirmed by ultrasonic and abdomino-pelvic plain X-ray.

Statements on funding and competing interests
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Competing interests: The authors work at the Abacus Centres for Contraception and Reproductive Health. In lieu of payment for training other clinicians in the technique of GyneFix insertion, Abacus has received some devices free of charge from Contrel, the manufacturer.

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