Study Abstract

The Priya Sensor is safe and well-tolerated, and the proprietary artificial intelligence algorithm to identify a woman’s fertile window and wirelessly communicates to the Priya App which can be installed on a wide range of smartphones.

Methods:
This study (PT-004) has two principal objectives: to validate the successful temperature transmission rate of the Priya Sensor to the Priya app, and to compare the fertile window predictions of the Priya Sensor to that achieved with a urine LH ovulation prediction kit. The study contains 2 cohorts: Group 1 will include up to 20 subjects and Group 2 will include up to 50 subjects.

Results:
A total of 17 subjects have been enrolled in Group 1 to date for whom a total of 29 cycles were completed with an overall successful temperature transmission rate of 97.5% (SD ± 3.8%). In addition, a total of 17 cycles were evaluable for assessing the correlation between the Priya fertility predictions and LH testing results. In all 17 cycles evaluated, the Priya Sensor predicted the fertile window an average of 2.6 days prior to the LH test. The most commonly reported adverse events, irrespective of a causal relationship to the device, consisted of mild abdominal cramps, vaginitis, and abdominal or pelvic pain. Virtually all adverse events were mild in severity and generally resolved without treatment within a few hours or days.

Conclusions:
The Priya Sensor provides precise, continuous core body temperature measurement which is interpreted by a proprietary artificial intelligence algorithm to identify a woman’s fertile window and wirelessly communicate this information to her smartphone. Early results with the Priya Sensor indicate that it is at least as effective as LH tests for predicting the fertile window and can reliably provide the fertile window predictions days ahead of the positive LH results. The Priya Sensor is safe and well-tolerated, and the ease of use may represent a significant advance in trying to conceive.

Introduction

• Despite recent advancements in fertility prediction, there are still a significant number of women and their partners who are trying to find an effortless yet accurate product to help them predict the ‘fertile window’.

• Many currently available products and tracking tools require a considerable amount of user interaction to record frequent temperature readings, urine test results, and other observations. This can lead to frequent errors, user stress, and frustration.

• Basal body temperature (BBT) charting, urinary luteinizing hormone (LH) testing, salivary ferning and analysis, and external temperature reading have traditionally been used as aids for predicting ovulation. However, the utility of these approaches is limited by:
  — Cumbersome temperature measurement devices.
  — Vagaries of the charting process when left to the patient.
  — Lack of generally-accepted criteria for objectively interpreting BBT graphs; data is prospective and does not predict ovulation.

• Other fertility detection methods are unable to capture circadian rhythm patterns. These patterns provide reliable and consistent fertility indications for predicting ovulation and are most reliable through 24/7 continuous core temperature monitoring.

• The Priya Sensor (Figure 1) was developed to overcome the limitations of earlier BBT sensing and other fertility detection methods, and has the following characteristics:
  — Designed to predict and notify the user when core body temperature readings are consistent with the ‘fertile window,’ that is, the pre-ovulatory period of time most favorable for achieving pregnancy.
  — Thermometric device inside a flexible medical-grade silicone polymer continuously measures and records core body temperature (every 6 minutes), along with cycle length, to identify the most fertile phase of the menstrual cycle.
  — The sensor transmits core body temperature data wirelessly to a mobile device allowing the proprietary algorithm to predict the fertile window.

• The objectives of this study are to evaluate:
  — Successful temperature data transmission rates from the intravaginal Priya Sensor to the Priya App on mobile devices.

Study Design

• Open label adaptive design study.
• Up to 20 participants are to be enrolled in the first group of the study and up to 50 participants are to be enrolled in the second group to complete the trial.
• After signing an institutional review board-approved informed consent form, participants wear the intravaginal Priya Sensor continuously during at least one menstrual cycle.
• Eligible participants assigned to 1 of 2 study groups:
  — Group 1: Designed to assess recent software updates to the Priya App.
  — Group 2: Designed to formally assess the usability, reliability and accuracy of the Priya Sensor.
• After completion of the first menstrual cycle in either group, participants are given the option of continuing to wear the Priya Sensor for up to 2 additional menstrual cycles.

Methods

Study Procedures

• Within 3 days of baseline (first day of menstruation), subjects contacted the Principal Investigator to confirm and verify the Priya Sensor instructions for use.
• Subjects inserted the Priya Sensor on the first day after the end of their periods and started twice daily testing (morning and evening) with an LH test kit and continued until a positive result (LH high) was detected followed by a negative result (LH low).
• Subjects also checked the temperature transmission data on the Priya App at least once a day and notified site personnel if they had any difficulty synchronizing the Sensor and the App.
• At the end of Cycle 1 (within 3 days after the first day of the next period), the subject contacted the Principal Investigator to complete a follow-up assessment for Cycle 1 that included:
  — Adverse events, including pain or discomfort with the use of the Priya Sensor.
The Priya Sensor is safe and well-tolerated, and the fertile window predictions days ahead of the positive LH Sensor indicate that it is at least as effective as LH tests for the identification of patterns associated with temporal rhythms within a few hours or days. Virtually all adverse events were mild in severity and generally resolved without treatment within a few hours or days.

### Preliminary Results

**Participants:**
- To date, 17 participants were enrolled into Group 1 and 13 completed ≥1 cycle.
- Three participants discontinued due to adverse events and 2 withdrew consent prior using the Priya Sensor.
- Nine participants provided a total of 17 evaluable cycles. An evaluable cycle is defined as at least 4 days of temperatures before the LH-high and no gaps (ring removed or no readings) of more than 3 hours before the LH-high.

**The demographic characteristics for the overall population and for subjects with evaluable cycles are summarized in Table 1. Data for the first cycles for 9 women are provided in this preliminary analysis.**

### Table 1. Demographic Characteristics of Group 1 Subjects

<table>
<thead>
<tr>
<th></th>
<th>Overall Group 1 Population to Date (N=17)</th>
<th>Group 1 Subjects with Evaluable Cycles (N = 9)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age (years):</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>34.4</td>
<td>34.2</td>
</tr>
<tr>
<td>Median</td>
<td>35.0</td>
<td>34.5</td>
</tr>
<tr>
<td>Minimum, Maximum</td>
<td>25.0, 45.6</td>
<td>270, 40.6</td>
</tr>
<tr>
<td>Standard Deviation</td>
<td>5.68</td>
<td>4.44</td>
</tr>
<tr>
<td><strong>Race/Ethnicity:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>16</td>
<td>9</td>
</tr>
<tr>
<td>Black or African American</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Asian</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Other</td>
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<td>0</td>
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<tr>
<td>Hispanic</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Non-Hispanic</td>
<td>17</td>
<td>9</td>
</tr>
<tr>
<td><strong>BMI (kg/m²):</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>23.3</td>
<td>22.7</td>
</tr>
<tr>
<td>Median</td>
<td>22.3</td>
<td>22.3</td>
</tr>
<tr>
<td>Minimum, Maximum</td>
<td>17.9, 32.6</td>
<td>19.4, 29.1</td>
</tr>
<tr>
<td>Standard Deviation</td>
<td>4.12</td>
<td>3.20</td>
</tr>
</tbody>
</table>

**Efficacy:**
- Efficacy results are presented for 17 evaluable cycles among the 9 subjects in this analysis.
- Temperature data was successfully transmitted from the Priya Sensor to the App on mobile devices for all evaluable subjects.
- In this preliminary analysis, the Priya Sensor predicted the fertile window an average of 2.6 days prior to the LH test (SD 1.6 days) (See table below).

**Priya Predictions Relative to +LH**

<table>
<thead>
<tr>
<th></th>
<th>N of Cycles (N = 17)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mean</strong></td>
<td>2.61 days</td>
</tr>
<tr>
<td><strong>Median</strong></td>
<td>2.33 days</td>
</tr>
<tr>
<td><strong>Standard Deviation</strong></td>
<td>17.5 days^2</td>
</tr>
</tbody>
</table>

1. Number of cycles meeting evaluability criteria (defined above)
2. No fertility predictions occurred after LH positive test result dates

**Safety:**
- The most commonly reported adverse events, irrespective of a causal relationship to the device, consisted of mild abdominal cramps, vaginitis, and abdominal or pelvic pain.
- Virtually all adverse events were mild in severity and generally resolved without treatment within a few hours or days.

### References