AACC Guidance Document on
the Use of Point-of-Care Testing in
Fertility and Reproduction

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INTRODUCTION
Point-of-care testing (POCT) is defined as clinical laboratory testing conducted close to the site of patient care, typically by clinical personnel whose primary training is not in the clinical laboratory sciences or by patients (self-testing) (1). POCT is an increasingly popular means of bringing laboratory testing closer to the patient. POCT can provide faster turnaround of results compared to core laboratory testing, because POCT eliminates transportation of blood specimens and considerably reduces processing and preanalytical steps required for laboratory tests. Faster test results offer the potential for rapid medical decisions. The convenience, ease of use, and speed are driving POCT growth. POCT described in this guideline can be performed in the hospital or clinic setting or even purchased over the counter and performed at home.

Despite the apparent simplicity, many factors can negatively affect POCT quality. Failure to follow manufacturer’s instructions, such as storing reagents improperly, incorrect sample collection, insufficient amounts of sample, and over- or undertiming test development, can lead to false positives and false negatives. Failure to analyze controls or troubleshoot when controls are out of range can impact accuracy. Seemingly minor interruptions, such as moving test devices during analysis, can affect POCT results. POCT is methodologically distinct, with different interferences and limitations compared to laboratory methods. Thus, POCT results are not necessarily harmonized with laboratory methods, and test results may not agree.

Although POCT has great potential to improve patient care, when inappropriately utilized or incorrectly performed, POCT can lead to unnecessary follow-up tests and procedures with considerable clinical and financial implications for the patient. Using POCT to improve the speed of care delivery in time-sensitive settings may also put the patient at risk if errors occur. Clinicians/POCT operators need guidance in utilizing POCT and in implementing good laboratory practices to obtain reliable results. The AACC Academy (formerly known as the National Academy of Clinical Biochemistry) developed best practice recommendations for use of POCT in patient care. The Laboratory Medicine Practice Guidelines (LMPG): Evidence-Based Practice for Point-of-Care Testing were published in 2007 (1). Those guidelines critically reviewed the peer literature, graded the evidence that links POCT to clinical outcomes, and provided recommendations for optimizing POCT utilization. They offer a comprehensive, systematic review of the POCT literature. However, recent POCT studies have been published, and the original 2007 LMPG consequently needs updating. Instead of revising the entire LMPG, sections of the document were prioritized for revision to narrow the focus of the revision process and expedite publication of updated recommendations. The first section to be revised was published last year as the AACC Guidance Document on Management of POCT (2). This manuscript addresses revision of the Reproductive Testing section of the 2007 POCT guidelines.

MATERIALS AND METHODS
The AACC Academy formed a committee of experts with interest and experience in POCT and laboratory testing for fertility and reproduction. The expert committee was composed of AACC Academy members and was supplemented with clinicians having Emergency Medicine and Obstetrics/Gynecology training. The committee divided into several subgroups to formulate clinical questions related to the use of POCT in the assessment of ovulation, pregnancy, premature rupture of membranes, and evaluation of fetal distress. Literature searches were conducted for peer literature that could address each clinical question. Common search engines were utilized such as PubMed, Cochrane, Embase, and Web of Science. Searches were limited to manuscripts published in the English language with emphasis on recent literature published since the previous 2007 guidelines. Publications on test performance, sensitivity, and potential interferences were also included. Guidance documents from other professional organizations such as the American College of Obstetrics and Gynecology (ACOG) were reviewed to not
duplicate recommendations. Once drafted by the committee, prospective guidelines were distributed among the AACC membership and to the wider laboratory community for public comment. Revisions were made to address each comment, and the final guidance document was approved by the AACC Academy Council prior to publication. This guidance document represents a consensus opinion of the expert committee and is not intended to be a comprehensive evidence-based review of the literature. This guidance provides best practice recommendations for use of reproductive POCT in patient management.

GUIDANCE RECOMMENDATIONS

Point-of-Care Ovulation Testing

Ultrasoundography, luteinizing hormone (LH), estradiol, and progesterone testing are the main methods employed to detect, predict, and monitor ovulation in clinical practice. They are also valuable tools to evaluate ovarian function and optimize the timing of fertilization in assisted reproductive procedures. The development of POCT such as basal body temperature (BBT) measurement, salivary and vaginal mucus ferning, and urinary LH/estrone-3-glucuronide (E3G) testing provides simple and convenient ways for women to identify their fertile windows (3).

Is the diagnostic accuracy of over-the-counter urine LH tests sufficient for detecting and predicting ovulation when compared to ultrasound as the gold standard for confirming ovulation?

Multiple studies have demonstrated that urine LH tests are accurate and reliable predictors of ovulation with the urinary LH surge preceding ovulation by approximately 1 day.

Follicle-stimulating hormone is secreted by the pituitary and promotes the growth of multiple follicles inside the ovary. These follicles produce estrogen to induce the abrupt increase of LH, which in turn triggers the dominant follicle(s) to rupture and release the mature oocyte(s). The menstrual cycle is divided into 2 phases—the follicle phase starts from the first day of menstruation until ovulation, and after ovulation the luteal phase begins until the onset of next menstruation. The average length of a woman’s menstrual cycle is considered to be 28 days; however, it has been long known that there exists considerable inter- and intra-individual menstrual cycle variability.

Urinary LH point-of-care (POC) tests were first performed in clinics by professional staffs, but now, over-the-counter urinary LH testing for fertility monitor are available for at-home testing without prescription.

Earlier reports established that commercially available urinary LH POC kits have excellent diagnostic sensitivity (85%–100%, median 100%) and predictive value for ovulation (85%–100%, median 93%) with ovulation occurring within approximately 1 day of the LH surge (1, 4, 5). As the performance of urinary LH surge detection kits has been established, more recent studies have focused on understanding the underlying population variability to improve ovulation prediction, including (a) developing optimal methods and algorithms for detecting the surge and (b) establishing guidance on the cycle-length tailored testing protocol in the menstrual cycle to increase the predictive value of the fertility monitoring methods (6, 7).

There is considerable interpersonal variability in hormone curves and menstrual cycle lengths. Statistical techniques have worked to improve the detection of ovulation (8). Published predictive methods classify 3 broad categories based on how the baseline LH is estimated—fixed days, peak LH day, and estimated LH surge—and define LH surge as a sustained rise in LH concentration above the baseline LH. The primary difference between the methods is how the LH baseline assessment is established (6). The fixed days method requires no previous cycle information, whereas the other 2 methods (peak LH day and estimated LH surge) need complete cycle data and can only be evaluated retrospectively.

Menstrual Cycle Monitoring was a prospective study of normally menstruating women between the ages of 18 and 40 (n = 40) that examined the interindividual variation of urine and serum reproductive hormones and their relationship to ovulation, as confirmed by ultrasound measurements. The investigation found that, among the study participants, the menstrual cycle length ranged from 22 to 37 days, the length of the luteal phase ranged from 3 to 15.5 days, and the day of ovulation ranged from day 8 to day 26 (9). More recent large-scale studies further confirmed the wide variations in the cycle length and date of ovulation at the population level (10, 11). The Menstrual Cycle Monitoring study also found that the LH surge preceded ovulation in all participants by a mean of 0.81 days (9). In a separate report of the same study, the urinary ranges of LH, E3G, and follicle-stimulating hormone reference to the actual ovulation day (determined by ultrasonography) were established. Daily urine samples of a complete cycle were collected and batch analyzed; LH concentrations were evaluated side by side using AutoDELFIA platform and an inhouse developed assay. AutoDELFIA assay detects intact LH, free β LH and the dominant urinary metabolite LH β core fragment, while the in-house assay only recognizes intact LH. The study showed that timing of the urinary LH peak was assay-dependent and did not always precede ovulation. The profiling using the AutoDELFIA assay gave relatively variable patterns of LH profile and also recorded the LH peak about 1-day lag behind the corresponding peak using the intact LH-specific assay. However, both assays worked equitably well in defining the day of the LH surge (12).

Leiva et al. recently conducted a study using ROC analysis to determine the optimal urinary LH threshold for the LH peak method in predicting ovulation within 24 h of LH surge. The best performance was at a threshold of 25 mIU/mL, which yielded a
sensitivity of 54%, specificity of 97%, positive predictive value (PPV) of 50%, and negative predictive value (NPV) of 98%. Therefore, the timing of the LH peak alone should not be used for predicting or determining ovulation status. A complimentary biomarker may be used concurrently to improve specificity and predictive value (7).

Each of these studies adds to our understanding of the individual variability in menstrual cycle physiology. More studies are needed to further clarify the mechanistic basis of this variability and address the complex interplay between female reproductive hormones and other important factors known to affect the female menstrual cycle such as body mass index, race, age, and environmental pollutants.

**Does the use of ovulation predicting kits (OPKs) measuring urine LH and E3G for predicting ovulation in women not treated in a fertility clinic improve outcomes (i.e., increase conception rates, decrease the number of clinic visits/the numbers of unwanted pregnancies) compared to not using prediction tests?**

Studies have shown that the use of OPKs may improve the likelihood of conception among healthy fertile women seeking pregnancy.

Infertility refers to the inability to conceive within 1 year of unprotected intercourse. Infertility can occur in both men and women. Worldwide infertility is estimated at 15% to 25% of women at reproductive age, and there is an upward trend of infertility prevalence in the last 2 decades, especially in developing countries (13, 14). In resource-limited settings, the options of medically assisted reproduction are either unaffordable or inaccessible. The use of OPKs to identify the optimal timing for intercourse in the fertile window is convenient and economical, with the potential of increasing the conception rates or, conversely, for preventing unwanted pregnancies.

Home-use urinary LH testing kits, targeting women who were not under fertility treatment, were developed and became commercially available in the 1980s, and have now become a commonly used tool for evaluating the LH surge to time intercourse in developed countries (15). However, it is still not affordable and may be entirely inaccessible to women in resource-constrained/resource-limited settings. Our literature search revealed the need for professional, nonindustry-sponsored studies to establish product performance independent of the manufacturers. Several recent publications examining the clinical impacts of home-use urinary LH testing as an ovulation predictor for women who are not under fertility treatment are summarized next.

**Impact of OPK use on the conception rate.** The Clearblue fertility monitor, which measures both LH and E3G to signal the onset and the closure of the fertility window, respectively, is the most studied device among the current commercially available urinary LH POCT. In a prospective randomized clinical trial (RCT) with 653 healthy participants, the study results showed that the cumulative pregnancy rate (for 2 cycles) was significantly higher in the test group using Clearblue Easy Fertility Monitor compared with the control group (22.7% vs 14.4%) (16). The improved version, Bluetooth-compatible Connected Ovulation Test System, was evaluated similarly in a more recent study involving 844 volunteers (17). The study results revealed the conception rates of the test vs the control group were 25.4% vs 14.7% (P < 0.001) after 1 cycle and 36.2% vs 26.8% (P = 0.026) after 2 cycles.

In a systematic review conducted to inform WHO guidelines on self-care interventions, Yeh et al. performed a meta-analysis that included 4 studies (3 RCTs and 1 observational study conducted between 1996 and 2013) with a total of 1487 participants from high-income countries. The analysis showed a higher self-reported pregnancy rate in those using OPKs in all 3 randomized control trials. The result from the pooled data analysis found that the ratio of pregnancy among participants with OPK use compared to pregnancy without OPK use (i.e., the relative risk) is 1.36 (95% CI 1.07–1.73) (18).

In addition, a recent study supported the adoption of online tracking systems and fertility monitoring apps as a simple, economical, and effective way to help couples achieve pregnancy. In a 24-cycle prospective effectiveness study (n = 256), the fertile window was identified with Clearblue fertility monitor or cervical mucus monitoring, and an online tool was deployed to primary care clinic for participants to record and monitor their observations and use the data to time intercourse. The study results showed that the pregnancy rate reached 100% at 24 cycles for those women using the hormonal (LH and E3G) fertility monitor (19). There is, however, significant variability among the apps available (20).

**Impact of OPK use on pregnancy avoidance.** Fertility awareness is a collective term describing several different contraceptive methods based on the collection and guided interpretation of various personal fertility signs/symptoms (including BBT, cervical fluid, menstrual cycle length, and urinary assays for reproductive hormones) to predict or identify the fertile window of the menstrual cycle, so an individual can use the information to reduce the probability of pregnancy. All fertility awareness methods meet the WHO criteria of modern methods with “a sound basis in reproductive biology, a precise protocol for correct use, and the evidence of efficacy under various conditions based on appropriately designed studies” (21).

The Marquette method involves users both observing their cervical fluid and measuring urinary hormone levels (both LH and E3G) with the recommended home-use POCT; the Clearblue Fertility Monitor. The effectiveness of this method was supported by a 12-month retrospective study with 204 couples, who were taught to correctly follow the protocol of the method for avoiding pregnancy. The study results showed that the 12-month pregnancy
rate of a “correct use” group was 0.6% (i.e., 99.4% effective) and of a “typical use” group was 10.6% (i.e., 89.4% effective) per 100 users (22). Subsequently, 2 comparative efficacy studies targeting perimenopausal (n = 160) and breastfeeding women (n = 816) suggest this method can be effective for pregnancy prevention in older and nursing women, especially with the adoption of online tracking/monitoring systems (23, 24).

Compared to other more effective contraceptive methods (contraceptive pill and physical barriers), the adoption of fertility awareness methods in the United States is low, but the demand has increased 3-fold from 2008 to 2015 as the result of the wider use of the app-or tech-based fertility trackers as contraception (21, 25, 26). More education, training, and research are needed to properly use hormonal fertility monitors as a means to empower women with awareness and good decision-making for their own reproductive well-being.

Impact of OPK use on other health outcomes. Home-based hormonal fertility monitors also impact other outcome measures, including the general acceptance and satisfaction of end-users to improve fertility awareness and knowledge, and enabling women to know their bodies better. The study findings also supported that the use of OPKs did not cause additional stress/anxiety in general, especially among those who do become pregnant (18).

The wide adoption of online tracking technology may further promote OPK use as an inexpensive and effective tool of fertility management to benefit women living in low-resource settings. However, more research is required to investigate the values/preferences and social harms/adverse events using a diverse set of OPKs other than the Clearblue monitor system, especially in resource-limited settings.

In a study using the data collected from a North American prospective internet-based cohort (n = 8363), it was found that the use of cycle apps to monitor and correlate multiple fertility indicators was associated with a 12% to 20% increase of conception rate after adjusting for possible confounding factors (27). In another study, analysis of big data (total 75 981 cycles) collected from 32 595 users of a connected ovulation monitor confirmed the existence of wide individual variations of cycle length and ovulation dates, and the data were used to derive the population range and a probability table of ovulation per cycle day (11). Meanwhile, the market for fertility apps is rapidly growing, and these apps are gaining wider acceptance for identifying female fertility windows. However, regulatory bodies, academia, and industry should collaboratively develop a standard metric evaluation system to objectively assess the performance and usefulness of these apps based on their intended uses and to inform consumers by periodically reporting the evaluations so they may selectively use only the evidence-based effective products (28).

Does the use of urine LH tests for predicting ovulation in women undergoing fertility treatment improve outcomes (i.e., increase conception rates, decrease the number of clinic visits/the number of fertility treatment cycles) compared to not using prediction tests?

Urinary LH POCT demonstrated a comparable performance among other ovulation monitoring methods for timing intrauterine insemination and confirming sufficient ovulation induction before oocyte retrieval during in vitro fertilization. Limited data supported urinary LH as a cost-effective measure. However, the actual cost-saving may vary depending on the other factors such as population demographics, geographic locations, health resource allocations, and insurance coverage; therefore, the cost-savings need to be empirically determined.

Assisted reproductive technologies that include fertility treatments handling both a woman’s egg and a man’s sperm are viable options for achieving a successful pregnancy (29).

OPK use for timing of insemination in intrauterine insemination (IUI). The timing of IUI is critical to increasing the chance of fertilization and subsequent pregnancy. The use of an at-home ovulation prediction kit (OPK) is one of the available methods to predict the optimal timing of IUI. In a meta-analysis including 2279 infertile couples from 14 RCTs comparing the effectiveness of different synchronization methods for IUI, no differences in pregnancy rate, live birth rates, or adverse events were found among the evaluated methods (i.e., serum or urinary LH detection, ultrasound detection of ovulation, BBT, and human chorionic gonadotropin [hCG]/gonadotropin releasing hormone agonist [GnRHa] administration) (30). A more recent cohort study (n = 232 normoovulatory women) compared the timing of therapeutic donor sperm inseminations using urinary LH vs ultrasound that showed no difference in the outcomes of pregnancy rate, live birth rates, and adverse events between these methods (31).

OPK use for confirming sufficient ovarian stimulation before oocyte retrieval. GnRHa trigger is a recently introduced procedure for ovarian stimulation before egg retrieval. The measurement of serum LH at 12 h after GnRHa trigger has been suggested for confirming the sufficiency of ovarian stimulation. However, the optimal serum LH threshold concentrations after the GnRHa trigger have not yet been established (32). Urinary LH concentrations are highly correlated with serum LH concentration in the fertile window, and urinary LH has been used for timing insemination in IUI. Thus, self-testing of the LH surge (defined as LH ≥ 15 mIU/mL at 12 h after injection) with home-use OPKs followed by communication via cellular phone has been proposed to replace serum LH testing as a simple, safe, and convenient way to confirm the adequacy of GnRHa trigger for oocyte maturation. In a recent multicenter prospective cohort study...
study conducted in Spain, Brazil, and Denmark, urinary LH results were obtained from 359 oocyte donors; 356 participants recorded positive and only 3 had negative urinary LH results, with 1 false positive and 1 false negative as confirmed by ultrasonography and serum LH results. This gave an overall sensitivity of 99.7% (355/356) and specificity of 66.7% (2/3) (33). In fact, the serum LH measurement for the only false-negative case was 18.6 mIU/mL, which is just above the defined positive trigger cutoff but is below the lower detection limit (25 mIU/mL) for the ovulation kit used. Therefore, a few negative LH results from an at-home ovulation monitoring kit would require reflex confirmation by serum testing to rule out false-negative results. A cost analysis (including direct and indirect cost) was also performed, and OPK use was shown to have a significant cost-saving. Serum LH would have cost 14 840€ (approximately US$17 680) while urine LH kits cost only 1855€ (approximately US$2010) (33). More empirical cost–benefit studies are needed to provide definitive evidence before integrating OPK testing as a standard procedure for oocyte retrieval protocol.

So far, many RCTs and a few longitudinal studies on fertility treatments are mostly focusing on the impact of a particular assisted reproductive technologies procedure on the outcome measures of time to pregnancy, the number of clinic visits, and the number of fertility treatment cycles. There is scarce evidence to support or refute that the use of OPK alone can significantly improve these outcomes.

Table 1 summarizes the research design/methodologies, the study size, and the main results of the studies cited in answering the 2 previous questions.

Is the diagnostic accuracy of nonurine ovulation POCT (including marketed and emerging devices) sufficient to predict ovulation when ultrasound is used as a gold standard?

Recently commercialized salivary ferning tests are reusable and easy to use but have subpar reliability and reporting accuracy. The new-generation biosensor-based BBT monitors equipped with algorithms for pattern recognition enhanced their ability to detect ovulation, henceforth may be suitable for family planning. The claimed high accuracy by the device manufacturers needs to be further augmented by postmarket performance studies. In addition, new devices equipped with software algorithms for continuously monitoring skin temperature, pulse rate (PR), and ferning patterns are emerging into the ovulation POC market. Preliminary studies of these emerging devices showed promising results. However, more comprehensive studies are warranted to investigate the actual performance and justify the usability of these emerging devices.

Current commercially available nonurine ovulation tests include BBT monitoring and salivary ferning analysis.

Salivary ferning test. The phenomenon of ferning (or crystallization) results from the cyclical increases of sodium and chloride concentrations in body fluids under the influence of estrogen. The ferning appearance due to the crystallization of NaCl is observable by examining the dried saliva or cervical mucus under a microscope (3).

Two POC devices based on the measurement of salivary electrical resistance were marketed in the 1990s but did not gain traction, likely due to the inferior performance compared to other competing devices in the marketplace. The reported diagnostic accuracy of both devices varied from 52% to 74% (34–36).

Recently, 2 relatively simple and user-friendly salivary ferning test kits were developed and became commercially available; each include a pocket-sized microscope for examining the ferning patterns. Know when Ovulation Monitoring System has a hand-held mini microscope and an accompanying smartphone app for recording the results to predict the fertile window. In a small open-label prospective study, compared with actual ovulation assessed by transvaginal ultrasound, salivary ferning was observed in 29 of 30 ovulatory cycles, while false-positive results occurred twice in 10 anovulatory cycles (37). A subsequent prospective observational study involving 107 healthy women for a total of 114 cycles reported a sensitivity of 88.6% and a specificity of 80% with a PPV of 93.3% and an NPV of 69% compared to the results using ultrasonography (38). Another salivary ferning device, Geratherm® ovu Control, has a mini-microscopic lens with a switchable light source fixed on a lipstick-sized stand. When assessed in women under in vitro fertilization treatment, this device showed a specificity of 78% and a sensitivity of 80% when compared to the rise of estradiol after follicle-stimulating hormone/hCG stimulation. The same study also showed an agreement of 89.4% between the evaluations of the ferning patterns performed by patients and by laboratory staff (39). In another comparison study, 74 healthy women performed Geratherm® ovu Control and a urinary LH test side by side, and the paired results showed a high level of conformity, from the 5th (100%) until the 14th (84%) cycle day, and from the 18th (80%) until the 22nd (96%) cycle day, corresponding to the pre-and postovulatory period; however, the lack of ultrasonography to confirm the ovulation makes the study results hard to interpret (40).

The advantages of the salivary ferning ovulation test are its reusability and relatively low cost, but the impact of physiological condition, disease state, and certain medications on the ferning patterns significantly limits the reliability of its results.

BBT monitors. The pattern of dip–rise–return of BBT reflects the changing level of thermogenic progesterone during the menstrual cycle; BBT reaches nadir approximately 1 day before ovulation, rises 0.5 to 1.0°F after ovulation, plateaus, and then returns to a lower range around the time of menstrual bleeding. BBT monitoring has been, and still is, widely used to estimate the
### TABLE 1. Studies investigating the improvements of clinical outcomes in women using urinary LH POCT to predict ovulation.

<table>
<thead>
<tr>
<th>AUTHORS</th>
<th>STUDY DESIGN</th>
<th>SIZE (n)</th>
<th>METHOD DESCRIPTION</th>
<th>REPORTED OUTCOMES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Robinson et al. (16)</td>
<td>Prospective RCT</td>
<td>653</td>
<td>Compared the self-reported pregnancy rates between test (use fertility monitor) and control.</td>
<td>The cumulative pregnancy rate for 2 cycles: 22.7% in test group vs 14.4% in control group.</td>
</tr>
<tr>
<td>Johnson et al. (17)</td>
<td>Open-label clinical trial</td>
<td>844</td>
<td>A home-based study comparing the self-reported pregnancy rates between test (use app-connected fertility monitor) and control groups.</td>
<td>Conception rates test vs control groups 25.4% vs 14.7% after 1 cycle, 36.2% vs 28.6% after 2 cycles.</td>
</tr>
<tr>
<td>Yeh et al. (18)</td>
<td>Meta-analysis of 4 RCTs</td>
<td>1487</td>
<td>A systematic review comparing women desiring pregnancy who managed their fertility with and without OPKs and reported outcome measures.</td>
<td>Home-based use of OPKs may improve pregnancy rate with no meaningful increase in stress/anxiety and with high user acceptability.</td>
</tr>
<tr>
<td>Bouchard et al. (19)</td>
<td>Prospective effectiveness study</td>
<td>256</td>
<td>Fertile window is identified with Clearblue fertility monitor or cervical mucus monitoring and recorded by online tracking.</td>
<td>Conception rates fertility monitor vs mucus vs combined groups: 80% vs 48%, 83% vs 72%, and 100% vs not available after 6, 12, and 24 cycles, respectively.</td>
</tr>
<tr>
<td>Fehring et al. (22)</td>
<td>Retrospective study</td>
<td>204</td>
<td>Participants from 4 clinics using Marquette fertility awareness method and the unintentional pregnancies were tracked to assess the effectiveness.</td>
<td>The 12-month pregnancy rates were 0.6 and 10.6 per 100 users in correct use group vs typical use group (effectiveness 99.4% vs 89.4%).</td>
</tr>
<tr>
<td>Fehring et al. (23)</td>
<td>Comparison efficacy study</td>
<td>160</td>
<td>The study is conducted in a university based in-person and online family planning service program using methods described in Fehring et al. (22).</td>
<td>The between-group comparison is not statistically meaningful due to the low positive rate.</td>
</tr>
<tr>
<td>Fehring et al. (24)</td>
<td>Comparison efficacy study</td>
<td>816</td>
<td>The same as the methods described in Fehring et al. (22).</td>
<td>The 12-month pregnancy rates were 3% and 14% in correct use group vs typical use group.</td>
</tr>
<tr>
<td>Stanford et al. (27)</td>
<td>Prospective cohort study</td>
<td>8363</td>
<td>Analysis of the tracked self-reporting data to assess the influence of using cycle apps on per cycle probability of conception.</td>
<td>After adjusting for potential confounders, use of cycle apps was associated with increased fecundability of 12%–20% per cycle of attempt.</td>
</tr>
<tr>
<td>Soumpasis et al. (11)</td>
<td>Big data analysis</td>
<td>32 595</td>
<td>Compare users’ perceived cycle characteristics with actual cycle characteristics using anonymized cloud data collected from women trying to conceive.</td>
<td>There exist wide variations in cycle length and the ovulation day. The info was analyzed to provide population ranges and probability table for better timing of the fertile window.</td>
</tr>
<tr>
<td>Cantineau et al. (30)</td>
<td>Meta-analysis of 14 RCTs</td>
<td>2279</td>
<td>A systematic review to evaluate the effectiveness of different synchronization methods for IUI in subfertile couples.</td>
<td>More research is needed to determine whether any difference exists in safety and effectiveness among methods used for synchronizing ovulation and insemination.</td>
</tr>
<tr>
<td>El Hachem et al. (31)</td>
<td>Prospective cohort study</td>
<td>232</td>
<td>Compare the cumulative live birth rates between those using urinary LH strips to detect LH surge vs those using ultrasound monitoring to time sperm insemination.</td>
<td>Urinary LH strip vs ultrasonography groups live birth rates −12.4% vs 9.2% and conception rates −19.9% vs 13.3%.</td>
</tr>
<tr>
<td>Cozzolino et al. (33)</td>
<td>Prospective cohort study</td>
<td>356</td>
<td>Urine LH testing was performed at home 12 h after the GnRHa trigger to replace serum LH for confirming LH surge.</td>
<td>OPK use resulted in an overall sensitivity of 99.7% (355/356) and specificity of 66.7% (2/3). There was a significant cost-saving when urinary LH kits ($2010) were used to replace serum LH ($17 680).</td>
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time of ovulation retrospectively. Modern digital thermometers and the available online services/smartphone apps make the recording of results more convenient. Innovative biosensor-based devices were introduced into the market to facilitate the pattern recognition of BBT and thus improve the predictability of ovulation.

OvuSense and OvuRing are vaginal biosensors. An open-label clinical study (n = 148) showed that OvuRing has 99% accuracy for ovulation date detection and an 89% accuracy rate in predicting the ovulation date when referenced to ultrasound assessment, while OvuSense claimed to have equivalent performance (41, 42).

Rather than its apparent rise during ovulation, BBT monitors rely on a dip of BBT to predict the onset of ovulation; thus, it is less accurate in predicting than in detecting ovulation retrospectively (43). Besides, BBT may fluctuate with the changes in climate, room temperature, alcohol use, medications, many physiological conditions, and stress.

Emerging wearable biosensors. Various forms of wearable sensors, such as arm bands, bracelets, and earbuds, are designed to measure the nocturnal skin temperature; some of them are still at the stage of pilot or proof-of-concept studies, while a few have emerged into market and offer a new way for monitoring ovulation in the menstrual cycle.

The Oura ring is a commercially available wearable sleep and activity tracker with a temperature sensor capable of detecting the dip in body temperature prior to ovulation. In a clinical study, several algorithms for tracking the start of menstruation and ovulation were developed and tested involving 22 women for whom ovulations were predicted by recording and analyzing the changes of intravaginal temperature using proprietary software. The best-performing algorithm was shown to have the sensitivity of 83.3% and with a similar PPV (44). A wrist-worn armband was shown to detect a sustained 3-day BBT shift pattern in 357/437 cycles (82%) in an observational study with 136 healthy nonpregnant participants (45). Another noninvasive wearable device is a thermometer that consists of an earpiece, which measures the ear canal temperature every 5 min during night sleep hours, and a base station that transmits the data to a smartphone application for analysis. Results from a feasibility study of 34 users yielded detection accuracy with a sensitivity of 92.31% when compared with data obtained from an ovulation test kit (46).

The changing levels of estrogen and progesterone are known to affect the cardiovascular system, and studies have demonstrated that PR significantly increases during the fertile window compared to the menstrual phase (47). Ava bracelet (Ava AG) detects the significant, concurrent phase-based shifts in wrist skin temperature, heart rate, and respiratory rate, followed by analyzing the data collected by the bracelet using an accompanying machine-learning algorithm. A prospective longitudinal study involving 237 conception-seeking Swiss women showed the Ava bracelet detected the fertile windows with 90% accuracy when compared with the ovulation date predicted by the Clear Blue Ovulation Monitor (48). In another prospective observational clinical trial with 91 healthy nonpregnant women, the PRs were measured by a photoplethysmography-based wrist sensor for fertility monitoring. A significant increase of PR (2.1 beats per minute, P < 0.01) during the fertile window was observed compared to the PR in menstrual phase (49).

So far, none of the emerging devices have published randomized prospective clinical studies using the gold standard ultrasonography to confirm pregnancy results. Demonstration of correlation between the use of the wearable BBT devices and the improved pregnancy outcome data is required to support the efficacy of these devices as reliably accurate ovulation predictors. Privacy of BBT monitor data transmission have similar concerns to other shared electronic health information. Nonurine ovulation tests provide convenience and home testing convenience for the device users to predict the fertile window either for increasing the possibility of conception or to avoid pregnancy.

**Point-of-Care Pregnancy (Beta-Human Chorionic Gonadotropin) Testing**

**When should POCT hCG testing be considered in place of laboratory hCG analysis?**

POCT should be considered in clinical situations where rapid diagnosis of pregnancy is needed for treatment decisions and laboratory analysis cannot meet the required turnaround time (TAT). This would include situations where patients present in the acute setting with unstable vital signs and symptoms, raising concern for ruptured ectopic pregnancy. A positive urine or whole blood (WB) bedside POCT hCG test combined with an ultrasound suggesting free fluid in the abdomen would be helpful when considering early operative intervention.

β-hCG is a sialoglycoprotein that is initially secreted by the trophoblastic cells of the placenta shortly after implantation of the fertilized ovum into the uterine wall (50, 51). The rapid rise in hCG serum levels after conception allows for the use of hCG as an early biomarker of pregnancy.

POCT has advantages and limitations when compared to laboratory tests (Table 2). POCT urine hCG tests take 3 to 5 min to develop a positive result and 3 to 15 min for a negative result. WB POCT hCG takes 10 to 14 min, while the central laboratory has analytical times of 9 to 25 min. Laboratory TAT is longer than this because it is a combination of analytical time and processing time, including time for the sample to clot and time for centrifugation. This is the inherent advantage of POCT, which utilizes native urine or WB samples that do not require processing. Faster stat laboratory hCG methods for plasma/serum samples are available on some laboratory analyzers (ranging between 9 and 10 min).
These TATs are comparable to the WB POCT hCG analysis times without considering the added processing time required for laboratory samples. In a recent prospective observational study (n = 265), the use of WB to replace urine as a specimen for a rapid hCG bedside pregnancy test resulted in a mean time reduction of 21 min in the emergency department (ED; 95% CI 16–25 min, P < 0.001). There was a 99.6% concordance between WB and urine test results (52). However, WB is not an approved specimen for all rapid hCG test kits and off-label use as a modified method with a WB specimen is not recommended (53).

One advantage of POCT is its portability. POCT kits can be carried to patient rooms in a clinic, by visiting nurses to a patient’s home, and even deployed in military field hospitals and other community settings. This is contrasted with the dedicated space required for laboratory instrumentation. However, some POCT devices and readers require access to a power source and may have downloaders that require fixed space on a countertop.

Laboratory methods are now mostly automated, with walk-away stat capabilities. POCT kits without readers are manual, visually interpreted tests that require timing and manual recording of results in the patient’s medical record. While some POCT devices and readers have electronic interface capabilities, manual documentation risks transcriptional errors. Manual documentation of electronic download of control data is needed to review for quality control shifts over time with both POCT and laboratory methods. POCT devices and readers require minimal maintenance beyond cleaning and disinfection while laboratory instrumentation requires ongoing preventive maintenance and repairs. Urine POCT hCG methods, however, only provide qualitative results with cut-off concentrations for positivity in the 20 to 25 mIU/mL range depending on the manufacturer, some with even higher cutoffs. There are a few WB hCG POCT methods that report quantitative results with positive cutoffs of 5 to 10 mIU/mL, although these methods have limited upper reportable ranges of 1250 to 2000 mIU/mL and POCT WB samples cannot be diluted to extend the range. Laboratory methods, on the other hand, report quantitative results and have wide analytical measurement ranges that can be extended by

<table>
<thead>
<tr>
<th>TABLE 2. Features of POCT and laboratory testing hCG.</th>
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<tbody>
<tr>
<td><strong>POCT hCG</strong></td>
</tr>
<tr>
<td>Fast turnaround of test results</td>
</tr>
<tr>
<td>Unprocessed samples (no centrifugation)</td>
</tr>
<tr>
<td>Native urine and WB samples</td>
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<tr>
<td>Portable</td>
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<tr>
<td>Manual, visually interpreted tests</td>
</tr>
<tr>
<td>Manual result reporting for visual tests (devices and readers have interfaces available)</td>
</tr>
<tr>
<td>Minimal maintenance</td>
</tr>
<tr>
<td>Qualitative results predominantly (few blood quantitative hCG methods)</td>
</tr>
<tr>
<td>No dilution (limited range for quantitative tests)</td>
</tr>
<tr>
<td>POCT does not match laboratory results</td>
</tr>
<tr>
<td>CLIA waived (urine)—blood is moderate complexity</td>
</tr>
<tr>
<td>Ease-of-use (minimal training)</td>
</tr>
<tr>
<td>Room temperature storage (limited refrigeration)</td>
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<tr>
<td>Unit-use tests (implement in low volume areas)</td>
</tr>
<tr>
<td>Analyze one test at a time</td>
</tr>
<tr>
<td>Higher cost per test (individual packaging)</td>
</tr>
</tbody>
</table>

Each row distinguishes a different feature of the test. The feature with an advantage is bolded in each row.
sample dilution to higher reportable results. Clinicians can follow patient trends over time when laboratory results are analyzed by the same method. POCT methods, however, do not necessarily match laboratory results, and variability between different POCT methods may not agree because the methods are different. These steps in the core laboratory give more confidence and reliability in the ensuing results.

Urine POCT hCG is considered waived testing under the CLIA law. CLIA-waived testing has minimum documentation requirements. Staff only need to follow manufacturer’s directions, pay a biennial CLIA certificate fee, and agree to be inspected. This allows the test to be conducted in a wide variety of settings under physician or nursing supervision without laboratory involvement. Laboratory hCG and WB POCT hCG are CLIA moderate complexity, which requires method validation, operator training and competency, daily quality control, participation in a proficiency testing program, biennial inspection, and other quality requirements.

Many variables should be evaluated when considering POCT hCG testing. Operators can perform the tests reliably with minimal training, while laboratory instrumentation requires technical staff with laboratory experience to operate, troubleshoot, and maintain the equipment. POCT kits can be stored at room temperature, depending on the manufacturer, while laboratory reagents and controls require refrigeration and freezers. POCT can be utilized in areas with low test volumes because the tests are unit-use, single-packaged kits. This is an advantage for those settings with occasional testing requirements but can become a limitation as the demand for testing grows. The POCT operator must perform one test at a time, which can slow turnaround of results if multiple samples need to be tested at the same time, unless the optical reader can process multiple tests at the same time. Laboratory analyzers, on the other hand, have higher sample throughput and can analyze multiple tests simultaneously. Laboratory analyzers utilize bulk reagents that can produce more tests at lower cost compared to POCT that are singly packaged. This appears to raise the unit cost of POCT tests.

Thus, POCT hCG has a number of features that allow for rapid test results in a variety of clinical settings. However, urine POCT is limited to qualitative results, and the available quantitative WB POCT has limited reportable ranges.

In what clinical scenarios should hCG POCT be considered? Is WB hCG POCT preferred over urine hCG POCT in patient care?

Compared to laboratory-performed hCG testing, routine urinary hCG bedside testing with POCT devices may reduce the test TAT. hCG POCT should be considered for the rapid determination of pregnancy status in conjunction with ultrasound, symptoms, and medical history. However, location of pregnancy, intrauterine vs ectopic, cannot be determined from POCT alone. The wait time for urine collection often offsets the gain in the testing TAT reduction. WB hCG POCT is more invasive than urine but is not affected by urine dilution or dependent upon the ability to spontaneously void. Thus, WB hCG may be useful when the patient is unwilling to provide a urine sample or unable to urinate due to trauma, an acute presentation, or painful condition making urine collection problematic. Weighing the costs associated with potential harms due to an undiagnosed pregnancy into consideration, accurate perioperative and ED hCG POCT can be advantageous.

Females of childbearing age frequently present to EDs, urgent care centers, physician offices, clinics, and other care providers with symptoms of pregnancy or other clinical conditions that may be pregnancy related, such as abdominal pain, vaginal bleeding, syncope, or shock. Qualitative and quantitative analysis of hCG in urine or blood (WB, serum, and plasma) is used for the early detection of pregnancy (54–56). Qualitative (yes/no) results are useful for detecting pregnancy prior to exposing the patient to radiation, teratogenic medications, and surgical procedures that can harm the fetus. Clinicians need to determine the pregnancy status of their patients to make timely treatment decisions, for teratogenic medications are contraindicated and diagnostic imaging that exposes a developing fetus to radiation should be avoided during pregnancy. However, the menstrual history alone is not reliable; for this reason, a rapid hCG test at the POC may be deployed (57).

Improving patient flow can shorten ED length of stay, and shorter length of stay is associated with higher patient satisfaction. One way to reduce ED length of stay is the adoption of POCT, enabling rapid delivery of test results through immediate bedside testing, to reduce test TAT and potentially lead to shorter time to decision-making, a critical operational benchmark in urgent care settings (58). The significant delay in awaiting the urine collection often compromises the time-saving advantage of bedside urinary POC hCG testing. Blood hCG should be considered when the patient is unable to urinate. Collection of blood samples is more invasive than urine samples, but phlebotomy may be preferred over patient catheterization when delays in testing and treatment/procedures are pending the collection of a urine sample. Spontaneous voiding can be complicated by clinical states where the patient cannot urinate (dehydration, pelvic trauma, and unstable clinical presentation), and blood hCG may be required in these situations. Few published studies report the actual TAT improvement of POC hCG testing in ED settings. One prospective study (n = 498) found that compared to laboratory-performed hCG testing, the adoption of urinary POC hCG testing in the ED significantly reduced the time to initial report and time to availability on the chart, with mean differences of 25 min.
A discriminatory zone based on hCG alone cannot exclude an abnormal pregnancy by hCG measurements alone (62, 63, 64). The discriminatory threshold is the hCG level that distinguishes patients with intrauterine pregnancies in whom a potentially viable intrauterine pregnancy with methotrexate administration or surgery. Thus, interpretation of hCG results in the evaluation of normal and abnormal pregnancy should be done in conjunction with clinical and sonographic findings to arrive at the correct diagnosis (71).

Other sources of hCG, such as an hCG injection, can result in detectable levels of hCG in blood or urine, which can cause diagnostic confusion in patients being screened for pregnancy (72). Chronic renal failure (due to decreased clearance) (73) and passive transfer of hCG to plasma recipients from unknowingly pregnant blood donors (74) have also been reported as sources of elevated hCG in nonpregnant women. There have been rare cases of individuals with familial hCG syndrome—an idiopathic elevation of hCG that has been shown to be inherited in an autosomal manner. The genetic basis of this rare syndrome has not been elucidated but is characterized by persistent low hCG levels in multiple family members with no other underlying etiology (75).

Do quantitative hCG results enhance test utility compared to qualitative hCG at the point-of-care?

Quantitative hCG results at the POC are only available from a few WB methods. WB POCT hCG is of limited clinical value in serial testing and trending hCG results early after conception due to the narrow reportable range of POCT methods and the rapid rise of hCG levels after implantation. If hCG trends are being monitored, POCT results must be followed by the same analytic methodology. hCG results will vary between methods due to differences in anti-bodies and test affinity to various forms of hCG.

Urine hCG tests have reported 99% accuracy for the detection of pregnancy as early as the same day of a missed menstrual period with published cutoffs of 20 to 50 mIU/mL. However, the actual performance is lower due to variability among tests to recognize giving methotrexate. Demonstration of the normal doubling of hCG levels over 48 h suggests a diagnosis of fetal viability, but it does not rule out ectopic pregnancy, and a rate of rising hCG concentration that fails to reach a 50% increase over initial hCG level or plateaus within 2 days suggests a failing or ectopic pregnancy. Falling levels confirm nonviability of the fetus but do not rule out ectopic pregnancy. Ectopic pregnancy may resolve spontaneously through tubal abortion. The risk of tubal rupture is similar across a wide range of hCG values (10–190 000 mIU/mL) (69, 70). Quantitative hCG levels are used not only to diagnose ectopic pregnancies but to guide treatment, as a quantitative hCG level > 5000 mIU/mL is considered a relative contraindication to use of methotrexate for treatment of ectopic pregnancies due to its higher risk of failure in these patients. Once methotrexate is used, the quantitative hCG needs to be measured serially to ensure the pregnancy is resolving and guide further treatment either through continued methotrexate administration or surgery. Thus, interpretation of hCG results in the evaluation of normal and abnormal pregnancy should be done in conjunction with clinical and sonographic findings to arrive at the correct diagnosis (71).

Other sources of hCG, such as an hCG injection, can result in detectable levels of hCG in blood or urine, which can cause diagnostic confusion in patients being screened for pregnancy (72). Chronic renal failure (due to decreased clearance) (73) and passive transfer of hCG to plasma recipients from unknowingly pregnant blood donors (74) have also been reported as sources of elevated hCG in nonpregnant women. There have been rare cases of individuals with familial hCG syndrome—an idiopathic elevation of hCG that has been shown to be inherited in an autosomal manner. The genetic basis of this rare syndrome has not been elucidated but is characterized by persistent low hCG levels in multiple family members with no other underlying etiology (75).
the different forms of hCG present in urine (76). Urine dilution is also a concern, and all urine hCG tests provide qualitative results.

Blood hCG concentrations at or around the time of the missed menstrual period (the fourth completed week since the last menstrual period) report a median serum concentration ranging from 205 mIU/mL (3–7340 mIU/mL) (77) to 560 mIU/mL (6–19 950 mIU/mL) (78). Cutoff concentrations discriminating premenopausal, perimenopausal, postmenopausal, early normal pregnancy, pregnancy in the second or third trimester, or gestational trophoblastic neoplasia may vary depending on the clinical category and manufacturer assay. Quantitative WB POCT hCG methods with an upper reportable range of 1500 to 2000 mIU/mL would be of limited value when trending hCG levels over time, depending on the patient’s initial hCG level, since blood hCG concentration rises rapidly in the first several days of pregnancy.

Early pregnancy loss or spontaneous abortion occurs in about 22% of clinically unrecognized pregnancies and 31% of pregnancies overall (79, 80). Because of the high sensitivity of hCG assays, women with serum/plasma hCG results above the upper reference limit during the initial days after conception may generate negative results in subsequent samples due to natural termination of pregnancy.

Most hCG assays utilize calibrators that are traceable to the WHO’s standard. However, the use of quantitative results to follow patients over time must use the same methodology, since POCT and laboratory hCG immunoassays may generate variable results due to differences in antibody affinity for various forms and fragments of hCG in circulation (76, 81). Contradictory results between POCT and quantitative serum laboratory testing should be closely investigated for the cause of the discrepancies to mitigate possible clinical ramifications. In summary, quantitative WB hCG is of limited clinical value due to the narrow reportable range of POCT methods and the variability of results between different methods, so quantitative POCT has not been shown to be superior to qualitative POCT hCG methods.

What is the stability of hCG in urine?

The stability of hCG in urine at different temperatures has showed significant variation.

Reis et al. and some earlier studies have reported urinary hCG is stable at temperatures of 2°C to 8°C for up to 48 h and temperatures around −20°C for longer periods, extending to over 3 months (82). For POC urinary hCG testing, long-term storage of urine specimens is rarely needed, and it was reported that hCG immunoreactivity in urine was not significantly affected when stored below 10°C for up to 5 days (83). However, the urine samples should not stand at ambient temperature for an extended period before testing to prevent the deterioration of urinary hCG.

On the other hand, quality control materials and population study specimens may need long-term storage. Lempiäinen et al. found variability in urinary hCG stability for samples stored at −20°C from 3 to 10 months, and the magnitude of hCG loss was correlated with high sample urea while adding 5% to 10% glycerol or storing at −80°C mitigated the activity loss. It is noteworthy that the stability of different isoforms of urinary hCG vary and may account for the variations of observed hCG stability (84).

When should measurement of urine specific gravity be considered?

Specific gravity measurement should be considered on urine samples when measuring hCG to determine viable pregnancy.

False-negative urine hCG results can occur due to dilute urine, very early pregnancy stage, or very high hCG concentrations (hook effect). A dilute urine specimen (specific gravity, 1.007) may not contain a high enough hCG concentration to produce a positive result despite the presence of a viable, intrauterine pregnancy. To mitigate the risk of a false-negative urine hCG result, manufacturers recommend analyzing the urine-specific gravity when conducting qualitative or quantitative urine hCG tests. For dilute urine samples, another sample should be collected, ideally the first morning void, or the urine hCG results should be confirmed using a blood sample (85–87).

Does proteinuria and other interferents affect urine pregnancy tests?

There are a few case reports in the literature of false-positive urine pregnancy tests as a result of proteinuria/other interferents.

Heterophile antibodies and other interferences are the primary causes for false-positive or false-negative urine hCG tests. However, urine tests for hCG may not give appropriate results based on the alkalinity or acidity of the urine specimen following medication/drug use or if there are very high levels of hCG such as in trophoblastic diseases (88, 89). In septic shock, transient passage of interferents into the urine have been shown to result in falsely elevated urine hCG (90). False-positive urine pregnancy tests have been seen in patients with nephrotic range proteinuria, systemic lupus erythematosus, ovarian failure, elevated gonadotrophins, and tuboovarian abscess. In addition, a false-positive urine pregnancy test has been reported in a 28-year-old woman with a history of tubal ligation who had a delayed diagnosis of obstructive pyelonephritis due to renal calculus. Marzinke et al. reported a false-positive urine pregnancy test in a patient with membranoproliferative glomerulonephritis type I (91–93). Proteinuria and rheumatoid factor have been shown to
produce false-positive urine hCG (93).

There are anecdotal reports of false-positive urine hCG results in patients with a urinary tract infection (UTI), in addition to warnings on POC hCG kit package inserts and inaccurate statements on health websites (94, 95). However, there are scant data in the scientific literature that support such claims. Mitchell et al. performed a study using 95 urine samples testing positive on urine culture with Gram-positive bacteria (95). Only 5 of the 95 samples tested positive for hCG. Of these, 2 were from pregnant women and 3 were from cancer patients. The authors concluded that UTIs do not cause false-positive results. This study had some limitations in that data were collected from a single site, a single test kit was used, and the study was not designed to evaluate the potential for false-negative results. Nevertheless, the results seem to be compelling and point toward the inability of bacterial isolates to produce false-positive results with these assay kits.

Overall, there are limited published studies on the role of interference by UTIs. The evidence does not suggest that UTIs can cause false-positive results on urine-based hCG POC assays. One should be cautious when laboratory findings are inconsistent with the clinical scenario and exercise prudence by utilizing different methods/assays as test platforms and/or other sample types (e.g., serum) for confirmation of ambiguous results.

**Are qualitative POC hCG devices susceptible to false-negative results due to antigen excess?**

False-negative qualitative POC hCG results may occur when testing urine specimens containing elevated concentrations of intact hCG or hCG variants.

Urine hCG devices use a sandwich immuno-assay format consisting of an immobilized capture antibody and a soluble, labeled detector antibody. In the presence of hCG, antibody-hCG-antibody “sandwich” accumulates at the test line, forming a visible band. Excess labeled antibody flows past the test line and accumulates at the control line, confirming validity of the test result. When hCG is absent, labeled antibody accumulates at the control line only, indicating a negative result. Any substance that prevents formation of the test line when hCG is present, creates a test line when hCG is absent, or prevents the formation of a control line may lead to misclassification of pregnancy status, delayed treatment, and possible adverse patient outcomes.

Qualitative POC hCG devices are designed to accommodate intact hCG concentrations typically observed in normal pregnancy (peak around 10–12 weeks with levels between 25 700–288 000 mIU/ mL). The hook effect is an immunologic phenomenon whereby the effectiveness of antibodies to form immune complexes can be impaired when concentrations of an antigen are very high. Device manufacturers often evaluate this threshold and indicate in the package insert the highest intact hCG concentration confirmed to generate positive results. Two commercially available devices indicate resistance to the hook effect up to 500 000 mIU/ mL, although it is important to note that this threshold varies by manufacturer and a careful reading of the package insert is recommended.

Several types of tumors produce an elevated level of the hCG, the high-level free β-subunit is especially related to the aggressiveness of the malignancies. hCG level can reach > 3 000 000 mIU/mL in patients with the complete hydatidiform mole (advanced molar pregnancy) or disseminated gestational choriocarcinoma. The amounts of hCG in the specimens of these patients greatly exceed and thus saturate the antibodies present in POC hCG devices and cause false-negative results due to the hook effect. While gestational choriocarcinoma is rare, hydatidiform mole occurs in approximately 1 per 1000 pregnancies in the United States. Choriocarcinoma can evolve very rapidly and result in death if left without treatment. Because the levels of elevation of hCG in hCG-secreting tumors vary widely, hCG testing should not be used solely in the initial assessment of the suspected cases. In fact, the false-negative urine/serum hCG results caused by the hook effect is a well-known problem that complicates the diagnosis of these 2 types of gestational trophoblastic diseases, as evidenced by the sporadic case reports in the literature (96–98). Compared with intact hCG or the free β-subunit of hCG in pregnancy, gestational trophoblastic neoplasms, and germ cell tumors, the low predictive value of the free alpha-subunit of hCG excludes its clinical use in these settings.

False-negative results may also occur when variant forms of hCG are present at sufficiently high concentrations to saturate the hCG antibodies in the test device. Excess hCG variant prevents the device antibodies from binding to intact hCG present in the sample, even though the intact hCG concentration may be well below the device’s indicated upper limit of detection. The variant hook effect is most commonly caused by hCG beta core fragment (hCGβcf), a product of proteolytic digestion of the hCGβ subunit formed during renal excretion and present only in urine. While the variant hook effect due to hCGβcf may be observed at any point in pregnancy, it is most likely when urine hCGβcf concentrations are highest, typically coinciding with the peak of plasma hCG concentrations observed between weeks 8 and 12 (99). hCGβcf is present only in urine and has not been demonstrated to interfere with hCG measurement in serum, plasma, or WB specimens. Although urine hCGβcf concentrations are typically highest between 10 and 16 weeks’ gestation, excretion patterns are variable, and false-negative results due to excess hCGβcf may be encountered at any point in pregnancy (100, 101). Commonly used qualitative POC hCG devices vary in their susceptibility to hCGβcf interference as some devices generated strong positive results, others generated weak positive results, and some generated clear negative results when used to evaluate a series of urine specimens with known intact hCG and hCGβcf concentrations (102). Some devices have been
reformulated, resulting in improved performance in specimens containing high concentrations of hCGβcf (103, 104).

Recommendations to reduce the hCG hook effect:

- Maximal intact hCG concentrations listed in the device package insert can be verified by clinical laboratory personnel.
- If a false-negative result due to the hook effect is suspected in the previously discussed clinical conditions, the urine specimen should be diluted using an hCG-free material and retested. A positive result upon dilution confirms the hook effect. Alternatively, a serum hCG test can be conducted.
- When clinically appropriate, quantitative hCG measurement should be performed in serum or plasma specimens to avoid hCGβcf interference. If urine testing is required, a device with minimal susceptibility to hCGβcf variant effect should be used.

Does excess biotin intake affect hCG measurement on POCT devices?

If repeat testing on a patient sample yields an invalid result or a result that is inconsistent with clinical history or previous laboratory testing, interference from biotin supplementation should be considered. Recommended next steps include quantitative serum/plasma hCG measurement or performance of a qualitative urine hCG test that is not subject to interference by biotin supplementation. After pausing biotin intake for 24 h, a second urine sample can be collected in nonemergent cases.

Biotin has recently become a focus of concern as an immunoassay interferent due to its increased popularity as a vitamin supplement, with several case reports and research studies demonstrating that excess biotin in patient samples can interfere with assays reliant on streptavidin-biotin interaction (105, 106). A recently published AACC guidance document on biotin interference lists immunoassays impacted by biotin; hCG can be falsely decreased in the presence of excess biotin on 2 commonly used immunoassay instruments (106).

Approximately 50% of biotin is secreted in urine unchanged. In general, 20% of ingested biotin dose is excreted in urine within 4 h. The rate of biotin excretion varies between 3 to 40 h at low doses and 7.8 to 18.8 h at high dose (107, 108). Therefore, qualitative urine hCG testing devices are subject to biotin interference in individuals taking dietary biotin supplements such as B-complex vitamins; coenzyme R; dietary supplements for hair; skin, or nail growth; multivitamins; prenatal vitamins; vitamin B7 supplements; and vitamin H. However, most over-the-counter multivitamins do not contain sufficient biotin to interfere with urine hCG devices when the manufacturer-recommended dosing is followed. However, repeated ingestion of multivitamin doses that exceed manufacturer recommendations or use of the recommended dose of biotin-specific supplements (typically ≥5 mg per pill) can cause high urine biotin concentrations that may interfere with test performance (109).

In one study, the authors investigated potential interference of biotin in qualitative POCT for hCG both in vitro and in vivo and showed that excess biotin produced invalid results on the QuickVue urine hCG device by preventing streptavidin-biotin interaction at the control line. However, other devices (Alere 20, Alere 25, Icon 20, Osom, QuPID, and SureVue) were not affected (110). Failure to form a control line will generate an invalid test result, potentially delaying care and prompting additional unnecessary testing.

If biotin interference is suspected, testing personnel should contact the ordering clinician to inquire about biotin use. Unfortunately, when asked to list current medications, many patients do not include over-the-counter supplements, and clinicians may not be aware of their patients’ biotin supplementation. Regardless of the information provided by the clinical team regarding biotin supplementation, laboratorians should suggest quantitative plasma or serum hCG measurement using an analytical platform free from biotin interference (105). In nonemergent situations, the patient may be instructed to discontinue biotin supplementation for 24 to 48 h, after which a second urine specimen may be collected. However, a longer period of discontinuation may be required in patients on an extremely high dose whose specimens will be tested by a test method that is particularly sensitive to biotin interference. As biotin is water-soluble and is rapidly removed from circulation, serum and urine concentrations should fall below the threshold required to interfere with test performance within 24 h of discontinuing biotin supplementation (111). As an alternative strategy, repeat testing of the original urine specimen may be performed using a qualitative hCG device that is not subject to biotin interference.

Point-of-Care Premature Rupture of Membranes Testing

What tests are available to predict prelabor rupture of membranes (PROM)?

Conventionally, testing for PROM includes observed pooling of amniotic fluid in vagina/posterior fornix, pH testing of the fluid (pH of vaginal fluid 3.8 to 4.5 vs amniotic fluid 7.1–7.3), and microscopic examination of dried vaginal fluid (ferning pattern is suggestive of PROM). More recently, multiple commercial test kits are also available to aid the diagnosis of PROM. These include Actim, Amnisure and ROM Plus in the United States and/or Canada. Fetal fibronectin should not be used for PROM diagnosis. Digital examination should be avoided to reduce risk of infection.

According to ACOG (112), PROM is defined as membrane ruptures prior to the onset of labor. PROM that occurs before 37 weeks of gestation is considered preterm PROM. It complicates approximately 2% to 3% of pregnancies in the United States. At
term > 37 weeks of gestation), PROM occurs in approximately 8% of pregnancies in the United States and is often followed by prompt delivery. PROM testing signals rupture of the membranes and clinical management depends largely on gestational age of the fetus. The most significant complication to the fetus relates to prematurity. As to the mother, IUI risk increases with duration of membrane rupture. Therefore, the ability to accurately diagnose PROM is crucial in clinical management of mother and fetus.

As previously noted, vaginal pooling, pH testing, and ferning are often used to confirm the diagnosis of PROM (112). Commercially available test kits (rapid lateral flow immunoassay) such as Actim, Amnisure, ROM Plus, and Amnioquick Duo are also available (113–115). Amnisure uses a biomarker called placental-α-microglobulin-1. Actim and ROM Plus use a biomarker called placental protein 12/insulin-like growth factor binding protein-1, which are the same protein (116). These proteins are present in substantially higher concentration in the amniotic fluid than in cervico-vaginal secretions with intact fetal membrane or maternal serum, making them ideal markers to detect ROM (117, 118). Unlike Actim and Amnisure test kits, the ROM Plus test kit and Amnioquick Duo also use a second biomarker, α-fetoprotein. The α-fetoprotein level increases in amniotic fluid as gestational age advances but decreases during third trimester.

When should PROM testing be performed?

Most instances of PROM can be diagnosed based on physical examination, clinical presentation, and patient history. PROM testing using commercial kits alone is not recommended without clinical signs of ROM, such as leakage of amniotic fluid from the cervical opening (119). As with other tests, the test result must be interpreted in the context of the patient’s clinical presentation. Unfortunately, there have been reports of misuse resulting in death and health complications for fetus and/or mother (120).

PROM management varies depending on gestational age and fetal presentation (112). PROM is typically of minimal risk compared to preterm PROM. Risks associated with preterm PROM are significantly higher with gestational age < 34 weeks (112). Currently, there is insufficient literature for the committee to recommend whether testing for PROM is clinically beneficial for ≥37 weeks pregnancy. In the United States, not all the commercial test kits are approved for PROM testing for any gestational age by the Food and Drug Administration. For example, Actim PROM is only approved for use with gestational age > 29 weeks, whereas Amnisure and ROM Plus can be used for all gestational age. However, there is little published literature on these tests’ accuracy for gestational age < 20 weeks. Healthcare providers need to be aware of such limitations to assure regulatory compliance and proven test performance for use with all stages of pregnancy.

In 2018, the Food and Drug Administration released alerts to providers about risks associated with improper use of PROM tests (120). Briefly, diagnosis of PROM and subsequent management of patients should not be made solely based on test results—overreliance on test results can cause patient harm. As with other tests, false positives or false negatives can occur. Testing in the absence of clinical signs of PROM can be particularly misleading. In women at term, positive Amnisure test results (121, 122) had been reported without clinical sign of ROM, and the positivity rate is higher with patients in active labor. While this is not considered PROM because these women were in labor, the studies suggest potential leakage of amniotic fluid or small amniotic protein as the delivery date approaches. Therefore, PROM testing should only be performed if clinically indicated and must be interpreted in conjunction with clinical assessments to diagnose PROM.

How does the performance of commercial test kits (e.g., Actim, Amnisure, ROM Plus) compare to traditional testing (pH, pooling, ferning test)?

Multiple studies have shown that performance of commercial test kits (e.g., Actim, Amnisure, ROM Plus) is equal to or surpasses the performance of the standard clinical assessment (SCA), which includes traditional testing methods (vaginal pooling, pH testing, and ferning). The committee recommends the use of commercial test kits for aiding in the diagnosis of PROM in women with suggestive symptoms.

Multiple prospective, observational studies have compared the performance of rapid commercial test kits to (a) the performance of SCA and (b) clinical outcomes (48 h follow-up and chart review) and are summarized in Table 3. In most of these studies, final PROM diagnosis was adjudicated by a physician who was blinded to the immunoassay test results. Most studies did not define the exact criteria of PROM, so the final diagnosis likely varied depending on the physician. However, some of the following criteria were taken into consideration in many studies: the SCA (some or all 3), clinical signs of fetal distress, latency period (varied between 48 h to 7 days), chorioamnionitis, etc. Esplin et al. (123) determined a sensitivity, specificity, PPV, and NPV of 91.7%, 97%, 94.8%, and 95.1% for the ROM Plus; 93.4%, 95.0%, 91.9%, and 96.0% for the Amnisure; and 95.0%, 98.5%, 87.4%, and 97.1% by SCA compared to the primary outcome, respectively. The performance characteristics of the 2 kits was statistically equivalent, and there was no significant difference when compared to the SCA methods.

Albayrak et al. (124) also determined that there was no significant difference in the performance of Actim, Amnisure, and the SCA methods compared to the final clinical diagnosis as determined by medical records after delivery. The sensitivities, specificities, PPV, NPV, and accuracy are, respectively, as follows: Actim commercial test kit, 89.8%, 97.5%, 97.5%, 89.5%, and
### TABLE 3. Summary of studies that assess the diagnostic performance of PROM tests.

<table>
<thead>
<tr>
<th>CONFIRMED CLINICAL DIAGNOSTIC CRITERIA OF PROM</th>
<th>n</th>
<th>TESTS</th>
<th>Sens(^a) (%)</th>
<th>Spec(^b) (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
<th>ACCU(^c) (%)</th>
<th>REFERENCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>48-hour follow up medical records review by an experienced physician who was blinded to the immunoassay test results</td>
<td>323</td>
<td>Amnisure</td>
<td>93.4</td>
<td>95.0</td>
<td>91.9</td>
<td>96.0</td>
<td></td>
<td>Esplin et al. (123)</td>
</tr>
<tr>
<td></td>
<td>322</td>
<td>ROM PLUS</td>
<td>91.7</td>
<td>97.0</td>
<td>94.8</td>
<td>95.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>323</td>
<td>SCA (all 3 present: ferning, pH, and pooling)</td>
<td>95.0</td>
<td>98.5</td>
<td>87.4</td>
<td>97.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visualization of vaginal amniotic fluid and/or persistence of oligohydramnios</td>
<td>75</td>
<td>Actim</td>
<td>100</td>
<td>96.7</td>
<td>97.8</td>
<td>100</td>
<td>98.7</td>
<td>Galletta et al. (129)</td>
</tr>
<tr>
<td></td>
<td>82</td>
<td>Amnisure</td>
<td>90.2</td>
<td>100</td>
<td>100</td>
<td>86.1</td>
<td>93.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>63</td>
<td>pH test</td>
<td>85.7</td>
<td>95.2</td>
<td>97.3</td>
<td>76.9</td>
<td>88.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>99</td>
<td>Amnisure</td>
<td>93.2</td>
<td>100</td>
<td>100</td>
<td>94.9</td>
<td></td>
<td>Eleje et al. (170)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AmnioQuick Duo</td>
<td>97.3</td>
<td>100</td>
<td>100</td>
<td>98.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Two of the following criteria: delivery within 48 hours to 7 days, chorioamnionitis, membrane rupture before delivery, adverse perinatal outcomes associated with prolonged PROM</td>
<td>220</td>
<td>Amnisure</td>
<td>95.5</td>
<td>89.1</td>
<td>89.7</td>
<td>95.1</td>
<td>92.3</td>
<td>Abdelazim et al. (128)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AmnioQuick Duo</td>
<td>93.6</td>
<td>86.4</td>
<td>87.3</td>
<td>93.1</td>
<td>90.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>pH test</td>
<td>72.7</td>
<td>80.9</td>
<td>79.2</td>
<td>74.8</td>
<td>76.8</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>76.4</td>
<td>83.6</td>
<td>82.4</td>
<td>78.0</td>
<td>80.0</td>
<td></td>
</tr>
<tr>
<td>Review of medical records after delivery by a physician blinded to the immunoassay test results</td>
<td>211</td>
<td>Amnisure</td>
<td>95.7</td>
<td>100</td>
<td>100</td>
<td>75</td>
<td></td>
<td>Ng et al. (125)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SCA (2 of 3 present: ferning, pH, or pooling)</td>
<td>78.1</td>
<td>100</td>
<td>100</td>
<td>36.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ferning</td>
<td>77.5</td>
<td>100</td>
<td>100</td>
<td>25.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>pH test</td>
<td>62.6</td>
<td>100</td>
<td>100</td>
<td>36.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>pooling</td>
<td>86.6</td>
<td>100</td>
<td>100</td>
<td>49.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Review of medical records after delivery. ROM was considered when the membranes are absent during vaginal examination or a positive pad chart was obtained.</td>
<td>285</td>
<td>ROM PLUS</td>
<td>99</td>
<td>91</td>
<td>95</td>
<td>99</td>
<td>96.5</td>
<td>Thomasino et al. (127)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SCA (visualization of fluid or 2 of 3 present: ferning, pH, or pooling)</td>
<td>85</td>
<td>98</td>
<td>99</td>
<td>77</td>
<td>89.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ferning</td>
<td>99</td>
<td>72</td>
<td>80</td>
<td>99</td>
<td>86.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>pH test</td>
<td>93</td>
<td>83</td>
<td>90</td>
<td>88</td>
<td>89.5</td>
<td></td>
</tr>
<tr>
<td>Review of medical records after delivery by a physician blinded to the immunoassay test results</td>
<td>167</td>
<td>Actim</td>
<td>89.8</td>
<td>97.5</td>
<td>97.5</td>
<td>89.5</td>
<td>93</td>
<td>Albayrak et al. (124)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Amnisure</td>
<td>94.3</td>
<td>97.5</td>
<td>97.6</td>
<td>93.9</td>
<td>95</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>SCA (visualization of fluid or 2 of 3 present: ferning, pH, or pooling)</td>
<td>88.6</td>
<td>94.9</td>
<td>95.1</td>
<td>88.2</td>
<td>91</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)Sensitivity.
\(^b\)Specificity.
\(^c\)Accuracy.
What might affect the performance of PROM tests?

Extended membrane rupture and minimal residual fluid can cause false-negative results. The presence of semen, cervical mucus, and hepatitis C can cause false-positive ferning results. Blood, semen, urine, bacterial vaginitis, cervicitis, trichomonas, and antiseptic agents have also been reported to cause false-positive pH/nitrazine results (125, 130, 131). Blood also affects the performance of different commercially available PROM tests to various degrees (132, 133).

Ramsauer et al. (132) investigated the impact of vaginal bleeding on the diagnostic accuracy of the Actim and AmniSure tests in patients with unknown membrane status. Cases that were identified as contaminated with blood had an increased occurrence of false positives and nonevaluable results (2% in AmniSure and 11% in Actim).

Bushman et al. (133) performed an in vitro study using WB samples spiked with varying concentrations of amniotic fluid proteins to evaluate the effect of blood contamination on the performance characteristics of three commercially available test kits, Actim, AmniSure, and ROM Plus. Blood contamination had no effect on the specificity or PPV value of each assay, which remained 100%. In this study, blood contamination increased the number of false negatives in all 3 assays, resulting in a reduction of sensitivity, NPV, and overall accuracy (Actim: 56.7%, 69.8%, and 78.3%; AmniSure: 61.0%, 72.3%, and 80.7%; ROM Plus: 96.7%, 96.6%, and 97.9%, respectively). ROM plus was able to detect amniotic proteins significantly better than the AmniSure or Actim PROM in this study. These discrepancies of the Ramsauer et al. (132) and the Bushman et al. (133) studies may be due to different cutoffs (analytical sensitivities) of commercial kits for placental-α-microglobulin-1/ insulin-like growth factor binding protein-1/placental protein 12 or α-fetoprotein levels (Table 4) and the different experimental design (real patients vs in vitro spiking samples) utilized.

Is the use of PROM test cost-effective?

There are currently limited studies in evaluating the cost-effectiveness of PROM testing. Given that prevalence of PROM is approximately 2% to 3% preterm and 8% of all term pregnancies in the United States (112), concern for PROM is not uncommon for pregnant women. Timely diagnosis is key in ensuring the health of mother and fetus. Accurate rule-out of PROM may reduce unnecessary emergent visits to the hospital/clinic or patient transfer to a higher level care facility and therefore reduce healthcare cost (134, 135).

According to Ferro et al. (134), at a large hospital in New Jersey, out of 1250 unscheduled visits, 68 had a primary concern of suspected PROM. Of these, 58 were discharged with PROM rule-out. Therefore, if there is a safe and accurate way for pregnant women to determine the presence of PROM, the emergent visits could be avoided. Although there is no at-home device for PROM in the United States currently, an at-home pH indicator has been trialed to differentiate amniotic fluid leakage (136, 137). However, the use of the device without proper clinical evaluation can result in false-negative diagnosis and presents a significant safety concern.

Echebiri et al. (135) use computer modeling and showed that the use of PROM testing can rule out 39% to 45% of the suspected cases. In resource-limited settings, this can lead to fewer patient transfers to facilities with comprehensive management capabilities for a higher level of care. According to Echebiri et al. (135), a single patient transfer costs $800 to $8800 in the United States. This is significantly higher than the Medicare reimbursement cost per test (<US$100). Therefore, it presents a significant cost-saving, as well as improved patient convenience.

Is the use of PROM test cost-effective?
Point-of-Care Fetal Scalp Lactate/Cord Blood Gas Testing

What is the clinical utility of fetal scalp sampling for pH or lactic acid during high-risk deliveries?

Fetal blood scalp (FBS) lactate values are being used to predict fetal acidosis and to indicate the need for intervention. Although an FBS sample taken within 1 h prior to birth correlates well with umbilical arterial or venous lactate, pH, and base excess, currently available studies indicate that FBS testing has minimal utility in prevention of metabolic acidosis.

Generally, FBS lactate testing is recommended over FBS pH testing for the management of intrapartum pregnancy with nonreassuring fetal heart tracing (138, 139). An exception was the National Institute for Health and Care Excellence (NICE) guideline where both FBS lactate and fetal scalp pH were presented as screening options following an abnormal fetal heart rate, with no preference given to one test over the other (140).

Hypoxia in the fetus induces changes in fetal scalp lactate and pH (141). FBS has been found to be unreliable due to multiple variables affecting sample integrity, including alteration of results by the alkaline amniotic fluid (142) and acidic meconium containing bile acid (143). Therefore, the recent NICE guidelines on intrapartum care has concluded that FBS increases the caesarean section rate and operative vaginal delivery rates (140).

Does fetal scalp pH or lactic acid monitoring reduce the rate of caesarian delivery?

There is insufficient evidence to conclude that use of FBS will reduce the rate of caesarian delivery.

One prospective RCT compared auscultation with cardiotocography (CTG) and CTG plus FBS of 695 high-risk deliveries where 232 were monitored by auscultation alone, 233 by CTG alone, and 230 by CTG and FBS. When comparing CTG with CTG plus FBS, there was a reduction of CS (18% vs 11%) in the CTG plus FBS (151, 152). Data from the systematic review of CTG + FBS trials and observational data from Australian hospitals support a conservative estimate of up to a 40% relative risk reduction in caesarean sections when FBS is added to CTG monitoring (153). In contrast, a recent Cochrane review concluded that use of FBS as an adjunct to CTG increased instrumented deliveries and reduced fetal acidosis but did not impact any other fetal outcomes (154).

Can fetal scalp pH and fetal scalp lactic acid be used interchangeably to predict fetal acidosis and hypoxia during complicated deliveries?

Fetal scalp lactate is used more than fetal scalp pH for fetal acidosis due to higher success rate and low volume of sample required.

Two RCTs (151, 152) compared some of the effectiveness and risks of fetal scalp sampling for lactate and pH measurement to assess fetal well-being, after a nonreassuring CTG trace during 3348 deliveries. No statistically significant difference between the 2 groups was found for neonatal encephalopathy, death, low Apgar score at 5 min, admission to neonatal intensive care unit, metabolic acidosis, or mode of birth. Because lactate testing...
What is the advantage of measuring fetal scalp lactate using a small POCT lactate analyzer over measuring pH?

A small volume, approximately 1 μL of blood, is required to measure fetal scalp lactate whereas 40–90 μL of sample is required to measure blood pH. POCT devices for measurement of lactate reduce TAT to a significant level.

For analysis of lactate by POCT strip analyzers, the volume of blood required is only 1 μL, while for pH, either measured on a POCT analyzer or on a blood gas analyzer, much more sample is needed (40–90 μL, depending on the device or cassette used). The use of POCT lactate measurements thus represents an attractive option to obtain quick TAT with a small volume of blood. The disadvantage, of course, is that lactate alone is measured, while a full blood gas including pH and lactate gives lot more data and confidence in data interpretation but needs a larger volume of blood and more training and competency for staff performing the test. The CLSI recommends a storage time of ≤15 min for WB lactate at room temperature (157).

What should be done for the validation of small POCT lactate analyzers?

For validation of fetal lactate devices, precision analysis and method comparison between POCT devices and laboratory-based blood gas analyzer or plasma lactate are recommended. The total allowable error tolerance should be much less than the biological variability of lactate in the normal range, as studies have shown that biological variability in patients with elevated lactate levels is lower.

Precision analysis may be performed using the CLSI: EP5A Complex Precision Protocol. In addition to the usual validation criteria, desirable precision when testing patients and evaluating them for lactic acidosis should be tighter, and total allowable error should be half of normal (158). Comparison studies should not be done on 2 portable devices but rather between the POC device and a central lab blood gas analyzer or plasma lactate method. The lack of a reference method for lactate analysis is one problem in the validation of a POC device to perform fetal scalp lactate. One investigator used the average of the Roche and Vitros plasma lactate as an internal reference method, as the Roche and Vitros methods were originally calibrated against different comparator methods (159). Another approach would be to use a blood gas analyzer as the reference method, since many blood gas analyzers have calibration schemes that return similar results across the range of lactate values normally encountered. When comparing blood gas analyzers to commercial plasma lactate assays, the blood gas analyzers exhibit a negative bias at higher lactate values (159, 160).

What is the clinical utility and cost-effectiveness of universal (vs selective) umbilical cord blood gas analysis?

Paired umbilical cord blood gas testing (Arterial and Venous) is recommended either immediately after delivery or from a clamped section of the umbilical cord within 2 to 3 min of delivery to provide appropriate care to newborns at birth, quality management, and training purposes. There is currently insufficient evidence to suggest that universal (every birth) collection of paired blood gas samples is more effective than collection in situations where the infant has low Apgar scores or poor outcome is suspected.

Umbilical artery blood gas and acid-base analysis provides objective information about fetal metabolic condition, specifically the presence of hypoxic (respiratory) or metabolic acidosis, at the time of birth. Guidelines from ACOG (150) and NICE (140) recommend selective testing of umbilical cord blood gas shortly after delivery. NICE guidelines recommend that if the baby is in “poor condition,” then the umbilical cord should be double clamped and paired (arterial and venous) samples should be drawn and sent for blood gas analysis. ACOG guidelines previously recommended that physicians should attempt to obtain paired umbilical cord blood gas samples in circumstances of cesarean delivery for fetal compromise, low 5-min Apgar score, severe growth restriction, non reassuring fetal heart tracing, maternal thyroid disease, intrapartum fever, or multifetal gestation (these guidelines have since been withdrawn) (140). In contrast, guidelines from the Society of Obstetricians and Gynecologists of Canada (161) recommend universal (all births) testing of paired umbilical cord blood gas samples drawn either immediately after delivery or from a clamped section of the umbilical cord within 2 to 3 min of delivery. Reasons given for universal sampling include that the information may help provide appropriate care to the newborn at birth and having blood gas data from all births can assist in quality assurance and quality improvement activities (161). Other advantages cited for universal umbilical cord blood gas analysis include staff becoming more proficient at collection, processing, and testing of samples so accurate results will exist when really needed, and umbilical blood gas results will always be available for interpretation when there are adverse outcomes to delivery (medicolegal argument) (162).

requires less specimen volume than pH, fetal scalp sampling success rate was estimated to be 98.7% and 79.4% for lactate and pH, respectively (139, 155). A reanalysis of the multicenter trial (156) found that the frequency of total operative inter-ventions was similar, but more caesarian deliveries were performed in the lactate group (16.5 vs 12.4%; relative risk 1.33, 95% CI 1.02–1.74).
Does arterial umbilical cord pH predict adverse neurological outcomes?

Low umbilical artery pH at birth is a risk factor for subsequent short- and long-term neurological outcomes but is a poor diagnostic test for predicting intermediate-or long-term complications related to childbirth.

A systematic review and meta-analysis of studies found that low arterial umbilical cord pH (studies used cutoffs between 7.0 and 7.2) had a strong, consistent, and temporal association with neonatal morbidity and mortality (hypoxic ischemic encephalopathy, seizures, intraventricular hemorrhage, or periventricular leukomalacia) and possibly the long-term outcome of cerebral palsy (163). Despite increasing evidence that low umbilical arterial pH is associated with poor neonatal and long-term outcomes, it is important to remember that few infants born with low umbilical arterial pH will develop cerebral palsy (164), and most infants that go on to have neurodevelopmental morbidities as children will be born with normal umbilical arterial pH (165, 166). Thus, low umbilical arterial pH at birth is a risk factor for subsequent short-and long-term neurological outcomes but is a poor diagnostic test for predicting intermediate- or long-term complications related to childbirth.

Does implementation of universal (as opposed to selective) umbilical cord blood gas analysis improve neonatal outcomes in centers implementing this practice?

There is limited evidence that universal cord blood gas testing as opposed to selective testing improves the neonatal outcomes.

A single center study was conducted over a 3-year period after implementing universal cord blood gas testing found that outcomes as measured by percentage of infants with umbilical artery pH <7.1 and number of nursery admissions did not change over the 3-year period, nor did universal testing affect the distribution of Apgar scores at 5 min or percentage of infants requiring resuscitation (167). Another study found that universal umbilical cord blood gas screening could identify infants with encephalopathy that were missed by other screening methods and predicted that universal screening would facilitate the timely initiation of therapeutic hypothermia to prevent secondary brain injury in the setting of hypoxic ischemia (168). We could not find any published evidence that universal testing leads to more reliable blood gas values as measured by number/percentage of samples meeting Westgate criteria for valid samples (based on relationship between arterial and venous pH and pCO2 in paired samples) or percentage of samples rejected by the laboratory or testing personnel.

Is universal umbilical cord blood gas analysis cost-effective?

There is insufficient evidence to conclude that universal umbilical cord blood gas analysis is cost-effective in most hospital practices with decreased nursery admissions.

One study performed at a single medical center (the same medical center observing improved neonatal outcomes associated with universal umbilical cord blood gas analysis) found that universal cord blood gas was cost-effective due to cost-savings associated with decreased nursery admissions resulting from universal cord blood gas testing (169). This large study from Australia observed that the initial cost of universal umbilical cord blood gas analysis was high but that there was significant savings from reduced special care nursery costs. Because findings are limited to one medical center, there is insufficient evidence to conclude that universal umbilical cord blood gas analysis is cost-effective in most hospital practices.

SUMMARY

POCT is gaining popularity in the field of reproductive medicine from predicting ovulation and diagnosing pregnancy to managing premature rupture of membranes and fetal distress at birth. By providing quick results at the site of patient care, POCT can offer the opportunity for faster medical intervention. The AACC Academy formed an expert committee to examine the key clinical questions and published literature surrounding the use of POCT in fertility and reproduction. This guidance was a consensus of expert opinion based on current peer-reviewed literature. These recommendations updated the previous LMPG: Evidence-Based Practice for Point-of-Care Testing and provide recommendations for best practice in the utilization of POCT. A discussion of supporting literature, as well as challenges and limitations of POCT was provided for each recommendation.

Nonstandard Abbreviations

POCT, point-of-care testing; LMPG, Laboratory Medicine Practice Guidelines; ACOG, American College of Obstetrics and Gynecology; LH, luteinizing hormone; POC, point of care; PPV, positive predictive value; NPV, negative predictive value; OPKs, ovulation predicting kits; RCT, randomized clinical trial; IUI, intrauterine insemination; hCG, human chorionic gonadotropin; GnRH, gonadotropin releasing hormone agonist; PR, pulse rate; BBT, basal body temperature; E3G, estrone-3-glucuronide; TAT, turnaround time; ED, emergency department; WB, whole blood; UTI, urinary tract infection; PROM, prelabor rupture of membranes; SCA, standard clinical assessment; FBS, fetal blood scalp; NICE, National Institute for Health and Care Excellence; CTG, cardiotocography.
Author Contributions
All authors confirmed they have contributed to the intellectual content of this paper and have met the following 4 requirements: (a) significant contributions to the conception and design, acquisition of data, or analysis and interpretation of data; (b) drafting or revising the article for intellectual content; (c) final approval of the published article; and (d) agreement to be accountable for all aspects of the article thus ensuring that questions related to the accuracy or integrity of any part of the article are appropriately investigated and resolved.

Authors’ Disclosures or Potential Conflicts of Interest

Role of Sponsor
The funding organizations played no role in the design of study, choice of enrolled patients, review and interpretation of data, preparation of manuscript, or final approval of manuscript.

Acknowledgments
The authors wish to acknowledge the information manufacturers provided in their package inserts, some of which is cited in this guidance document. Neither the authors of this guidance document nor the AACC Academy are endorsing any manufacturer product. The mention of specific products is intended only as an example supporting key points made in the guidance, and the examples mentioned are not representative of all products that are available world-wide. The committee developing this guidance would like to remember Dr. Harvey Kincaid Jr. who contributed to this document but unfortunately passed before the document was completed. We send our condolences to his family and are thankful for all of Harvey’s contributions.

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