Cervical Length Estimation and Cervicovaginal Fluid for Placental $\alpha$-Microglobulin 1 Testing to Screen Women Had Threatened Preterm Labor for Time till Spontaneous Labor

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Abstract

Objectives: Evaluation of diagnostic performance of rapid testing of cervico-vaginal fluid (CVF) for fetal fibronectin (FFN) and placental $\alpha$-microglobulin 1 (PAMG-1) as screening of women presented by threatened preterm labor (TPTL) with intact membranes for duration till getting spontaneous labor (SL). Patients & Methods: 37 women presenting with TPTL underwent CVF samplings before digital examination and then underwent transvaginal ultrasonography for estimation of cervical length (CL). All women received fluid and anxiolytic therapy and if uterine contractions persisted, all had received tocolytic therapy with oral nifedipine or intravenous magnesium sulphate according to requirements. Incidence of SL within <48 hr, 2 - 7 and 7 - 14 days was recorded and related to CL and result of FFN and PAMG-1 tests.

Results: Incidence of SL was 13.5%, 35.2% and 51.3% within 48 hr, 2 - 7 and 7 - 14 days, respectively. Duration till labor after sampling was positively correlated with CL, while was negatively correlated with positive FFN and PAMG-1 tests. Positive FFN test had high specificity, while positive PAMG-1 test had high sensitivity for labor within 7 days. Regression analysis defined short CL and positive PAMG-1 test as significant predictors for short duration till SL. ROC curve analysis defined short cervix and positive PAMG-1 test as significant predictors for labor within 48 hr and within 2 - 7 days respectively and combined negative PAMG-1 test and CL of 20 - 25 mm were significant predictors for labor within 7 - 14 days. Conclusion: PAMG-1 test had high specificity, if positive, for predicting SL and high NPP, if negative, for excluding labor within 7 days, so it can be used as rapid adjuvant to clinical evaluation to help management decision-making. Moreover, PAMG-1 test is recommended screening test for being easy-to-use bedside test, provides
rapid results, can be used after vaginal exam and coitus and does not require a speculum examination or specialized equipment to analyze results.

**Keywords**

Threatened Preterm Labor, Cervico-Vaginal Fluid, Fetal Fibronectin, Placental α-Microglobulin 1, Screening, Spontaneous Labor

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### 1. Introduction

Threatened preterm labor (TPTL) can be defined as the occurrence of regular uterine contractions before completion of 37 gestational weeks (GW) at rate of one contraction per ten minutes and persists for more than 30 minutes without cervical dilatation [1]. TPTL is associated with neonatal morbidity and mortality as well as maternal morbidity induced by antepartum bed rest and unnecessary treatment [2].

Risk stratification in women with preterm contractions would allow targeting of interventions to women who will indeed deliver preterm and avoiding unnecessary treatments, associated with potential complications, in symptomatic women who are unlikely to deliver preterm [3]. Cervical length (CL) measurement in women with TPTL showed modest accuracy for predicting PTB [4] and combined with fetal fibronectin (FFN) testing may allow risk stratification for spontaneous delivery in women still pregnant 7 days after TPTL [5].

Fetal fibronectin is an extracellular matrix glycoprotein which is produced by amniocytes and cytotrophoblasts [6]. FFN acts normally as glue between maternal decidua and amniotic membranes [7]. FFN can be detected in cervico-vaginal fluid (CVF) in all pregnancies before 22 GW, but between 24 and 34 GW indicates a risk of impending PTL [8] when its concentration in CVF is ≥50 ng/ml, but it showed no adequate sensitivity or specificity to predict a delay of labor at term [9]. For prediction of PTL within 7 days, quantitative FFN measurement was not superior to qualitative test with CL, but added more costs [10]. Accuracy of combined serial CL measurements and FFN for predicting PTB in nulliparous patients was reported to be low [11]. Blood-stained swabs of CVF for FFN testing were still effective for predicting PTL, but had higher false positive rates [12].

Placental α-microglobulin-1 (PAMG-1) is a human 34 kd fetal glycoprotein that was first isolated in 1975 from amniotic fluid [13]. PAMG-1 is an endometrial protein produced by uterine glands since 1st trimester of pregnancy [14]. PAMG-1 is present in the amniotic fluid (AF) and CVF of pregnant women. However, its concentration in AF of pregnant women (2000 - 25,000 ng/ml) is several thousand higher than that found in CVF when the fetal membranes are intact (0.05 - 0.2 ng/ml) [15]. PAMG-1 is not present in urine and at low levels in maternal blood, thus reducing the risk of inaccurate results in the presence of other fluids [16].
1. Objectives

Evaluation of diagnostic performance of cervico-vaginal fluid (CVF) for fetal fibronectin (FFN) and placental α-microglobulin 1 (PAMG-1) as rapid screening of women presented by threatened preterm labor (TPTL) with intact membranes for duration till getting spontaneous labor (SL).

1.2. Design

Prospective clinical intervention.

1.3. Setting

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2. Materials & Methods

The current study was conducted since Jan 2017 after approval of the study protocol by the Local Ethical Committee. All women presenting with TPTL which is defined as development of painful regular uterine contractions persisting for >30 minutes at a rate of one contraction/10 minutes at gestational age (GA) of 24 - 35 GW without cervical dilatation [17] were eligible for evaluations.

Exclusion criteria included cervical dilatation ≥ 3 cm, vaginal bleeding, vaginal discharge, uterine over distention secondary to multiple gestations or polyhydramnios or multiple gestations, systemic infection, fever > 38˚C, fetal distress, intrauterine growth restriction, blood pressure ≥ 140/90 and ≤100/60 mmHg, any obstetric contraindications to tocolytic agents, and history of any thromboembolic disorders. All women were clinically evaluated without digital examination or insertion of speculum until cervical secretion swabs were obtained. Then, vaginal examination was performed and speculum was inserted to assure the cervical status if closed or obtained and if opened membranes were bulging at the level of external os or not. Transvaginal ultrasonography was performed, with empty bladder and minimal pressure, the shortest length between the internal and external os was measures with clearest image after 3 measurements, before and after Valsalva’s maneuver [18].

2.1. Vaginal Sample Obtaining and Testing

According to manufacturer instructions, vaginal samples were obtained for qualitative evaluation of the presence of fFN and PAMG-1 in cervical secretions. Sampling was postponed in women had digital examination or intercourse within 24-hr before attending to the hospital. Also, sampling was prohibited in women had vaginal bleeding or infection.

1. Sample I: Before speculum examination, blind vaginal sample (sample-I) was obtained using a sterile flocked swab that was inserted 5 - 7 cm into the vagina for 30 s, removed and then was placed in a vial containing buffer solution and actively rotated in the buffer for 30 s. Then, the swab was removed and a test strip for PAMG-1 (The PartoSure Immunoassay Kit, Parsagen Diagnostics, Inc.,
Boston, USA) was incubated in the buffer solution for 5 min and then was re-
moved to be read. Test is positive on appearance of two red lines at control and
test sites, negative if only one red line appeared at control site and invalid is only
one red line appeared at test site.

2. Sample II: Speculum was inserted and a sterile Dacron swab was rotated in
the posterior fornix of the vagina for 10 s. Then the swab was inserted into a
tube containing buffer solution in which it was thoroughly mixed for 10 - 15 s,
was removed and a test strip for FFN (QuikCheck™ FFN, Hologic©) was incu-
bated in the buffer for 10 min, removed and read. Positive FFN test (two lines)
suggest cervical secretion FFN concentration is >50 ng/mL, or negative (single
line).

2.2. Management

On admission, all women were asked to relax and stay in bed for 1-hr, and ad-
ministered Lactated Ringer’s solution 500 ml and 50 mg pethidine (Meperidine
hydrochloride, Dolantin). If uterine contractions persisted, oral nifedipine
(Adalat capsules, Bayer AG) 20 mg as a loading dose, then every 30 min till ute-
rine contractions stop; for a maximum total of 3 doses. After regular uterine
contractions stopped, a maintenance dose using nifedipine 20 mg was given
orally every 8 hours for 48-hr [19]. Then, uterine contractions were monitored
by continuous electrical external fetal monitoring for 90 min after the loading
dose and then every uterine contractions and fetal heart sounds were monitored
every 6 for 24 hours [20]. During follow-up, maternal blood pressure was re-
corded every 15 min for 2-hr after the loading dose, and every 6-hr for 48-hr with
monitoring for the side effects of nifedipine.

Management outcome was considered successful if no contractions recurred
after inhibition for 12 h by nifedipine and patients were maintained on main-
tenance dose until 34 weeks of gestation. In case of continued or recurred ute-
rine contractions after inhibition for 12-hr, Intravenous magnesium sulphate
was administered as 4 g loading dose of 4 gm followed by 2 gm/hr continuous
infusion that was increased by 1 g/hr hourly until successful tocolysis or failure
of treatment [21].

2.3. Study Outcomes

Study outcomes included the incidence of spontaneous PTB within <48 hr, 2 - 7
and 7 - 14 days and the relation of this incidence to cervical length and result of
FFN and PAMG-1 tests. The diagnostic performance criteria (sensitivity, speci-
ficity, positive and negative predictive values and positive and negative likeli-
hood ratios) of CL and FFN and PAMG-1 tests for prediction of such outcome
were evaluated.

2.4. Statistical Analysis

Obtained data were presented as mean ± SD, numbers and percentages. Possible
relationships were investigated using Pearson linear regression analysis. The diagnostic performance criteria (sensitivity, specificity, positive and negative predictive values and positive and negative likelihood ratios) of CL and FFN and PAMG-1 tests were calculated versus the time of delivery as a true positive event. Regression analysis (Stepwise method) was used for stratification of studied parameters as specific predictors. Sensitivity & specificity of studied parameters as predictors for time till SL were evaluated using the receiver operating characteristic (ROC) curve analysis judged by the area under the curve (AUC) that was compared versus null hypothesis that AUC = 0.5. Statistical analysis was conducted using the IBM SPSS (Version 23, 2015; IBM, South Wacker Drive, Chicago, USA) for Windows statistical package. p value < 0.05 was considered statistically significant.

3. Results

Through duration of study 45 women presented with manifestations of threatened PTB; 8 women were excluded for not fulfilling inclusion criteria and 37 women were enrolled in the study (Figure 1). The study included 37 women in age range of 18 - 37 with a mean age of 28.5 ± 6.4 years and mean BMI of 29.6 ± 5.5 kg/m² with 16 women were obese with BMI > 30 kg/m². All women were multigravida, but 21 women were nullipara. Majority of women had TPTB

![Figure 1. Consort flow sheet.](image-url)
manifestation in range of 24 - 28 GW. Details of enrollment data of studied women are shown in Table 1.

All enrolled women had short cervix as estimated at time of attendance with a mean CL of 18.9 ± 4.6 mm. Six women (16.2%) had CL < 15 mm, 18 women (48.6%) had CL in range of 15 - 20 mm and 13 women (35.2%) had CL in range of 21 - 24 mm. Five women (13.5%) had labor within 48-hr, 13 women (35.2%) had labor within a range of 2 - 7 days and 19 women (51.3%) had labor within a range of 7 - 14 days and <14 days. Regarding cervical secretion testing, 10 samples (27%) were positive for FFN and 19 samples (51.3%) were positive for PAMG-1, but only 7 samples (18.9%) were positive for both tests.

Duration till labor after sampling showed positive significant correlation (r = 0.484, p = 0.002) with CL, while showed negative significant correlation (−0.404, p = 0.013) with positive PAMG-1 test. On the other hand, duration till labor after sampling showed negative non-significant (r = −0.247, p = 0.141) correlation with positive FFN test.

Evaluation of diagnostic performance criteria of studied tests detected low sensitivity of positive FFN test, while showed high specificity and +LR for prediction of labor during 48 hours or in range of 2 - 7 days after sampling. On

| Variables                  | Findings |
|----------------------------|----------|
| **Maternal age (years)**   | ≤20, n (%) | 1 (2.7%) |
| 21 - 25, n (%)             | 8 (21.6%) |
| 26 - 30, n (%)             | 16 (43.3%) |
| 31 - 35, n (%)             | 7 (18.9%) |
| >35, n (%)                 | 5 (13.5%) |
| Mean (±SD)                 | 28.5 ± 6.4 |
| **Body mass index (kg/m²)**| <25, n (%) | 3 (8.2%) |
| 25 - 30, n (%)             | 18 (48.6%) |
| >30, n (%)                 | 16 (43.2%) |
| Mean (±SD)                 | 29.6 ± 5.5 |
| **Gravidity**              | G1, n (%) | 15 (40.5%) |
| G2, n (%)                  | 12 (32.5%) |
| G3, n (%)                  | 10 (27%)  |
| **Parity**                 | Nullipara, n (%) | 21 (56.8%) |
| Para-1, n (%)              | 11 (29.7%) |
| Para-2, n (%)              | 5 (13.5%)  |
| **Gestational age at inclusion (weeks)** | 24, n (%) | 7 (18.9%) |
| >24 - 28, n (%)            | 27 (73%)  |
| >28, n (%)                 | 3 (8.1%)  |
| Mean (±SD)                 | 26.1 ± 4.6 |
contrary, positive PAMG-1 test showed high sensitivity for detection of women who will get labor within 48-hr or in range of 2 - 7 days after sampling. Moreover, negative PAMG-1 test can exclude the possibility of getting labor within 48-hr or within a range of 2 - 7 days after sampling with high NPV (89.47% & 77.78%, respectively). Positive sample for both FFN and PAMG-1 can define women at high risk of labor within 48-hr or within a range of 2 - 7 days after sampling with high specificity (84.38% and 87.5%, respectively) and +LR (2.56 & 2.46, respectively). Unfortunately, positive FFN and/or PAMG-1 tests had equivocal diagnostic performance criteria for prediction of labor after 7 days (Table 2).

Regression analysis of short CL, positive FFN test and positive PAMG-1 test for prediction of short duration between sampling time and getting labor defined short CL ($\beta = 0.441$) and positive PAMG-1 test ($\beta = 0.351$) as significant ($p = 0.003$ & 0.016, respectively) predictors.

ROC curve analysis of the predictability of short CL, and result of fFN and PAMG-1 tests for risk of labor within <48-hrs defined short cervix as significant sensitive predictor with AUC = 0.116 ($p = 0.006$) (Figure 2), while negative PAMG-1 test was a significant specific predictor for postponing labor beyond >

Table 2. Diagnostic performance criteria of FFN and PAMG-1 testing of samples of cervical secretion of studied women presented by TPTL.

|                  | Sensitivity % | Specificity % | +LR    | -LR    | PPV%  | NPV%  | Accuracy % |
|------------------|---------------|---------------|--------|--------|-------|-------|------------|
| <48-hr           | Both FFN & PAMG-1 | Value        | 40     | 84.38  | 2.56  | 0.71  | 28.57      | 90          | 78.38       |
|                  | CI            | 5.27 - 85.34 | 67.21 - 94.72 | 0.67 - 9.80 | 0.34 - 1.48 | 9.46 - 60.48 | 81.25 - 94.92 | 61.79 - 90.17 |
|                  | FFN test      | Value        | 40     | 75     | 1.60  | 0.80  | 20         | 88.89       | 70.27       |
|                  | CI            | 5.27 - 85.34 | 56.6 - 88.54 | 0.47 - 5.47 | 0.38 - 1.68 | 6.81 - 46.1 | 79.19 - 94.39 | 53.02 - 84.13 |
|                  | PAMG-1 test   | Value        | 60     | 53.12  | 1.28  | 0.75  | 16.67      | 89.47       | 54.05       |
|                  | CI            | 14.66 - 94.74 | 34.74 - 70.91 | 0.57 - 2.86 | 0.25 - 2.31 | 8.21 - 30.91 | 73.46 - 96.31 | 36.92 - 70.51 |
| >48-hr - < 7 days| Both FFN & PAMG-1 | Value        | 30.77  | 87.5   | 2.46  | 0.79  | 57.14      | 70          | 67.57       |
|                  | CI            | 9.09 - 61.43 | 67.64 - 97.34 | 0.65 - 9.36 | 0.53 - 1.17 | 25.95 - 83.53 | 61.17 - 77.56 | 50.21 - 81.99 |
|                  | FFN test      | Value        | 38.46  | 79.17  | 1.85  | 0.78  | 50         | 70.37       | 64.86       |
|                  | CI            | 13.86 - 68.52 | 57.85 - 92.87 | 0.65 - 5.22 | 0.48 - 1.25 | 26.12 - 73.88 | 59.60 - 79.27 | 47.46 - 79.79 |
|                  | PAMG-1 test   | Value        | 69.23  | 58.33  | 1.66  | 0.53  | 47.37      | 77.78       | 62.16       |
|                  | CI            | 38.57 - 90.91 | 36.64 - 77.89 | 0.92 - 3.02 | 0.22 - 1.28 | 33.15 - 62.03 | 59.15 - 89.43 | 44.76 - 77.54 |
| >7 - < 14 days   | Both FFN & PAMG-1 | Value        | 5.26   | 66.67  | 0.16  | 1.42  | 14.29      | 40          | 35.14       |
|                  | CI            | 0.13 - 26.03 | 40.99 - 86.66 | 0.02 - 0.19 | 1.01 - 2.00 | 2.17 - 55.59 | 32.11 - 48.45 | 20.21 - 52.54 |
|                  | FFN test      | Value        | 16.67  | 63.16  | 0.45  | 1.32  | 30         | 44.44       | 40.54       |
|                  | CI            | 3.58 - 41.42 | 38.36 - 83.71 | 0.14 - 1.49 | 0.88 - 1.97 | 11.54 - 48.46 | 34.89 - 54.43 | 24.75 - 57.9 |
|                  | PAMG-1 test   | Value        | 16.67  | 63.16  | 0.45  | 1.32  | 30         | 44.44       | 40.54       |
|                  | CI            | 3.58 - 41.42 | 38.36 - 83.71 | 0.14 - 1.49 | 0.88 - 1.97 | 11.54 - 48.46 | 34.89 - 54.43 | 24.75 - 57.9 |

CI: Confidence interval; LR: Likelihood ratio; PPV: Positive predictive value; NPV: Negative predictive value; fFN: fetal fibronectin; PAMG-1: Placental α-microglobulin-1; p < 0.05 indicates significant difference versus the null hypothesis that AUC = 0.5.
48-hr but to <7 days with AUC = 0.715 (p = 0.033) (Figure 3). On the other hand, negative PAMG-1 test as significant (AUC = 0.297, p = 0.035) sensitive and long cervix are significant (AUC = 0.722, p = 0.021) specific predictors for good response to tocolytics and postponing of labor to occur within a range of 7-14 days (Table 3, Figure 4).

Figure 2. ROC curve analysis of studied variables for prediction of labor within 48 hours after sampling.

Figure 3. ROC curve analysis of studied variables for prediction of labor within >48-hrs - <7 days after sampling.
Figure 4. ROC curve analysis of studied variables for prediction of labor within >7 - <14 days after sampling.

Table 3. ROC analysis of enrolment cervical length and results of fFN and PAMG-1 tests of cervical secretion of studied women presented by TPTL.

| Time of labor          | Variables     | AUC  | SE   | p    | 95% CI     |
|------------------------|---------------|------|------|------|------------|
| Within < 48 hr         | Cervical length | 0.116| 0.069| 0.006| 0.20 - 0.251|
|                         | Positive FFN test | 0.575| 0.144| 0.594| 0.292 - 0.858|
|                         | Positive PAMG-1 test | 0.666| 0.123| 0.239| 0.425 - 0.907|
| Within a range of 2 - 7 days | Cervical length | 0.454| 0.102| 0.645| 0.254 - 0.653|
|                         | Positive FFN test | 0.588| 0.101| 0.382| 0.390 - 0.787|
|                         | Positive PAMG-1 test | 0.715| 0.087| 0.033| 0.544 - 0.886|
| Within a range of 7 - 14 days | Cervical length | 0.722| 0.085| 0.021| 0.555 - 0.889|
|                         | Negative FFN test | 0.385| 0.094| 0.231| 0.201 - 0.569|
|                         | Negative PAMG-1 test | 0.297| 0.088| 0.035| 0.125 - 0.469|

AUC: Area under curve; SE: Standard error; CI: Confidence interval; FFN: fetal fibronectin; PAMG-1: Placental α-microglobulin-1; p < 0.05 indicates significant difference versus the null hypothesis that AUC = 0.5.

4. Discussion

Duration till delivery of studied women who presented by manifestations of TPTL was positively correlated with cervical length (CL), while was negatively correlated with positive FFN and PAMG-1 tests of cervical secretion sample. Positive FFN test had high specificity and +LR, while positive PAMG-1 test had high sensitivity for detecting women who will get labor within 7 days after sampling among studied women presented by manifestations of TPTL. However,
negative PAMG-1 test can exclude the possibility of getting labor within 7 days with high NPV. These findings indicate the possibility of reliance on these rapid tests for prediction of outcome of women presented by TPTL and so can help management decision-making regarding hospital admission with its related costs and consumption of hospital resources.

Moreover, Regression analysis defined short CL and positive PAMG-1 test as significant predictors for short duration between sampling time and getting labor. Differentially, ROC curve analysis defined short cervix as significant predictor for labor within 48-hr, while positive PAMG-1 test was a significant specific predictor for postponing labor within a range of 2 - 7 days and combined negative PAMG-1 test and CL in range of 20 - 25 mm were significant predictors for delivery between 7 and 14 days, while positive FFN test was non-significant predictor for delivery within the range of 7 - 14 days.

These data go in hand with Nikolova et al. [22] who reported that in patients presenting with symptoms of PTL with intact membranes and minimal cervical dilatation (≤3 cm) positive PAMG-1 test indicated spontaneous PTB within 7 days with a high degree of accuracy, but negative result indicated unlikelihood of spontaneous PTL within 14 days and in a comparative study versus FFN and CL, Nikolova et al. [23] found PAMG-1 detection is the best predictor of imminent spontaneous delivery within 7 days with sensitivity, specificity, PPV and NPV of 80%, 95%, 76% and 96%, respectively. Also, Werlen et al. [24] confirmed the excellent specificity and PPV of PAMG-1 test and documented that dependence on PAMG-1 test of can allow reduction of the rate of unnecessary hospitalization. Thereafter, Ehsanipoor et al. [25] reported that the presence of PAMG-1 in cervical secretion of women presented by TPTL is associated with an increased likelihood of delivery within 7 days with sensitivity and specificity rates of 54.6% and 82.7%, and PPV and NPV of 31.6% and 92.5%, respectively. Moreover, Zork et al. [26] found the PPV of FFN for delivery within 7 and 14 days in both singletons and twins with a CL <10 or 11 - 25 mm was low (10% - 25%).

In line with the obtained results concerning the diagnostic performance of PAMG-1 test, Di Fabrizio et al. [27] reported that PAMG-1 test had high efficacy to identify women at risk of imminent PTL within 7 days of testing, and its high NPV can prevent unnecessary admission and therapies and Melchor et al. [28] found the PPV of PAMG-1 was significantly higher than FFN or phosphorylated insulin-like growth factor-binding protein-1 in predicting spontaneous PTB within 7 days of testing in women with TPTL. Recently, Lotfi et al. [29] found PAMG-1 test was statistically superior to standard clinical assessment alone, with respect to specificity in risk assessment of imminent spontaneous PTL in patients with symptoms of TPTL and negative PAMG-1 test result can reduce up to 91% of unnecessary admissions for women presenting with TPTL.

The obtained results concerning the diagnostic performance of PAMG-1 test and review of literature spot light on its efficacy to be used as the sole rapid bedside test for predicting outcome of women had manifestations of TPTL. In
support of the efficacy of PAMG-1 test, Ravi et al. [30] compared the qualitative FFN test at 50 ng/ml threshold to PAMG-1 test for assessing risk of imminent spontaneous PTB in women with symptoms of PTL and found PAMG-1 test is a better predictor of spontaneous delivery within 7 d while maintaining a very high NPV. Also, Melchor et al. [31] found positive PAMG-1 test was >4 times more reliable than positive FFN test in predicting imminent spontaneous PTL and is a more reliable biomarker which is associated with fewer false-positive results that could lead to a reduction in unnecessary admissions, interventions and use of hospital resources.

5. Conclusion

The PAMG-1 test had high specificity, if positive, for predicting and high NPP, if negative, for excluding labor within 7 days in women with TPTB manifestations, so it can be used as a rapid adjuvant to clinical evaluation to help management decision-making. Moreover, the use of PAMG-1 test is recommended for its advantages for being an easy-to-use bedside test, provides rapid results, can be used after vaginal exam and coitus and does not require a speculum examination or specialized equipment to analyze results.

Limitations

One of the limitations of the current study was the small sample size, but this was due to the low incidence of TPTB and conduction of the study in a single institute. Another limitation was the absence of control group to estimate the levels of these parameters in women with normally progressing women.

Recommendation

Wider scale multicenter comparative studies are mandatory to establish the obtained results and suggestions.

Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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