SERUM AMYLOID A AS A NOVEL MARKER TO MORBIDLY ADHERENT PLACENTA

By

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ABSTRACT

Background: Placenta accreta is a placenta where the placental villi adhere directly to the myometrium, placenta increta is a placenta where placental villi invade into the myometrium, and a placenta percreta is a placenta where the placental villi invade through the myometrium and into the serosa. About 75% of morbidly adherent placentas are placenta accretas, 18% are placenta incretas, and 7% are placenta percretas.

Objective: To detect the changes of serum amyloid A in women with morbidly adherent placenta, to assess the changes of serum amyloid A in women with morbidly adherent placenta and to assess use serum amyloid A as biological marker beside ultrasound & doppler findings for prenatal detection of morbidly adherent placenta.

Patients and Methods: This study was conducted on women attending Al Azhar University Maternity Hospital, Obstetric clinic or emergency room and admitted to antepartum inpatient high-risk service. The current study had been conducted on 60 pregnant women who were divided into three equal groups as follows: Group I: placenta previa. Group II: placenta accreta or increta and Group III (control group): included normal pregnant women.

Results: Regarding sensitivity and specificity of serum amyloid-A, maternal serum amyloid-A level ≥15.3 (μg/mL) had high specificity& PPV, and moderate sensitivity& NPV in differentiating previa group. Also, level ≥15.3 (μg/mL) had high specificity& PPV and low sensitivity& NPV in differentiating accreta group from control group. In addition to that maternal serum amyloid-A level ≥15.3 (μg/mL) had high specificity& PPV and low sensitivity& NPV in differentiating previa/accreta groups.

Conclusion: Morbidly adherent placenta (MAP) was characterized by failure of the placenta to separate at delivery, with potential for significant perinatal and maternal morbidity and mortality. Detection of level of maternal SAA can be used as an additional tool to detect MAP.

Keywords: Serum Amyloid A, placenta.

INTRODUCTION

Morbidly adherent placenta (MAP), including placenta accreta, increta and percreta, is characterized by failure of the placenta to separate at delivery, with potential for significant perinatal and maternal morbidity and mortality. Placenta accreta occurrence has risen steadily to 1 in 533 between the years 1982 and 2002. This 13-fold increase parallels the increase in cesarean delivery (CD). When prior CD is combined with placenta previa, the risk for accreta increases with each prior CD, from 11 to 40%, 61 and 67% for one, two, three and
While multiple risk factors for morbidly adherent placenta have been described, mechanisms of accreta development remain opaque and pre-delivery prediction is limited. Proposed mechanisms of accreta include excessive trophoblast invasion into the myometrium (Valentini et al., 2017). Deficient decidua enabling placental implantation onto the myometrium, suggested by increased accreta risk with previa alone, and a combination of both (Bartels et al., 2018).

Antenatal diagnosis of placenta accreta, as it can reduce maternal morbidity by enabling a scheduled delivery by a multidisciplinary team in a tertiary care center. Ultrasound identifies many, but not all, women with MAP and without significant benefit from magnetic resonance imaging (Cheung and Chan, 2012).

SAA exerts immune-regulatory effects and key effects on trophoblastic migration, invasion, and differentiation. At low concentrations, SAA regulates trophoblast invasion and metalloprotease activity within the placental micro-environment, both of which are important for placental homeostasis and placental invasion (Sandri et al., 2014).

Regarding Serum amyloid A (SAA), which is a family of apolipoproteins associated with high-density lipoprotein (HDL) in plasma. Different isoforms of SAA are expressed constitutively (constitutive SAAs) at different levels or in response to inflammatory stimuli (acute phase SAAs). These proteins are produced predominantly by the liver. The conservation of these proteins throughout invertebrates and vertebrates suggests that SAAs play a highly essential role in all animals. Acute-phase serum amyloid A proteins (A-SAAs) are secreted during the acute phase of inflammation. Regarding ultrasonography features of placental lacunae, it appears as multiple linear, irregular vascular spaces within the placenta has been found to be predictive of placenta accreta. It has been shown to have the highest sensitivity to detect placenta accreta among the other diagnostic features. In patients who had a previous caesarean delivery and a low-lying anterior placenta, the lower uterine segment has been measured, and it was found that all patients with placenta accreta had a myometrium of less than 1 mm This was shown to be as predictive for morbidly-adherent placenta as placental lacunae (Tikkanen et al., 2011).

The present work aimed to detect the changes of serum amyloid A in women with morbidly adherent placenta and assess the changes of serum amyloid A in women with morbidly adherent placenta.

**PATIENTS AND METHODS**

**Site:** This study was conducted on women attending Al-Azhar University Maternity Hospital, Obstetric clinic or emergency room and admitted to antepartum inpatient high-risk service.

**Study population:** The current study had been conducted on (60) pregnant women who were divided into three groups as follows: **Group I:** including (20) women with placenta previa. **Group II:** including (20) women with placenta accreta or increta. **Group III** (control group): which included (20) normal pregnant women.
Women were recruited according to the following inclusion and exclusion criteria.

**Inclusion criteria:**

Age: ≥ 18 years old. Gestational age ≥ 28 weeks' gestation. Previous cesarean section or hysterotomy. The diagnosis is usually established by ultrasonography and occasionally supplemented by Doppler.

**Exclusion criteria:**


**Ethical issues:**

The protocol of the study was presented for approval by the ethical committee of the Department of Gynecology and Obstetrics of Al-Zhar University Maternity Hospital.

**Consent process:**

The population sample under study was informed about research protocol and informed consents are granted from each participant before inclusion.

**Study design:**

**Type of study:** Prospective cohort observational study.

**Methodology:** A detailed history and thorough physical examination of all participants had been carried out. A complete laboratory investigation was done, i.e. A complete blood picture with platelet count. Liver functions test. Renal function test. Coagulation profile. Fasting two hours blood glucose level.

**Intervention:** Patients upon admission, after diagnosis of placenta previa, accreta or increta. A sample of venous blood had been taken from each patient participating in the study under aseptic conditions.

**Laboratory method:** Blood samples were centrifuged at 2500 g for 15 minutes at 4°C, separated into serum aliquots, and stored at −80°C until used for the SAA assay. Levels of SAA had been be assayed simultaneously for both groups using the same microliter plates provided with the Human SAA solid-phases and which enzyme-linked immune-sorbent assay kit (Bio Source Europe, Nivelles, Belgium), according to the manufacturer’s protocol. The inter-assay and intra-assay coefficients of variation were 7.4% and 6.1%, respectively. The sensitivity had been <0.004 μg/mL.

**Data collection:**

Demographic data, delivery outcomes and laboratory evaluations were recorded and compared in both groups. Data or results which are collected after arrangement in suitable manner by a process known as processing of data may be manual or computerized. These data should be confidentially protected.

**Statistical Analysis:**

Data were collected, revised, coded and entered to the Statistical Package for Social Science (IBM SPSS) version 23. The quantitative data were presented as mean, standard deviations and ranges when their distribution found parametric. The comparison between two independent groups with quantitative data and parametric distribution were done by using Independent t-test; while the comparison between more than two group
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was done by Using One Way ANOVA test. Spearman correlation coefficients were used to assess the correlation between two quantitative parameters in the same group. Receiver operating characteristic curve (ROC) was used in the quantitative form to determine best cut of point, sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and AUC of serum Amyloid A.

The confidence interval was set to 95% and the margin of error accepted was set to 5%. So, p-value was considered significant when P-value > 0.05.

RESULTS

There was a statistically significant difference found between two groups regarding number of CS, while there was no statistically significant difference found between two groups regarding Age, BMI, and GA (Table 1).

Table (1): Comparison between normal & placenta(previa +accrete) groups regarding demographic data

<table>
<thead>
<tr>
<th></th>
<th>Normal group</th>
<th>Placenta (previa+accreta)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>Mean±SD</td>
<td>27.45 ± 2.68</td>
<td>28.63 ± 2.65</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>23 – 33</td>
<td>23 – 33</td>
</tr>
<tr>
<td>BMI</td>
<td>Mean±SD</td>
<td>29.74 ± 1.36</td>
<td>29.47 ± 1.74</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>27.80 – 31.90</td>
<td>26.40 – 33</td>
</tr>
<tr>
<td>GA</td>
<td>Mean±SD</td>
<td>32.15 ± 2.78</td>
<td>32.38 ± 2.50</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>28.00 – 36.00</td>
<td>28.00 – 37</td>
</tr>
<tr>
<td>No. of CS</td>
<td>Mean±SD</td>
<td>1.80 ± 0.70</td>
<td>2.25 ± 0.71</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>1.00 – 3.00</td>
<td>1.00 – 3</td>
</tr>
</tbody>
</table>

*: Independent t-test

There was a statistically significant difference between two groups regarding serum amyloid A level (Table 2).

Table (2): Serum amyloid A in normal group and placenta (previa and accreta)

<table>
<thead>
<tr>
<th>Serum Amyloid A</th>
<th>Normal group</th>
<th>Placenta (previa+ accreta)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean±SD</td>
<td>11.56 ± 2.19</td>
<td>19.86 ± 5.72</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>8.80 – 16.20</td>
<td>9.20 – 30.3</td>
</tr>
</tbody>
</table>

*: Independent t-test

There was a statistically significant difference between the three groups regarding age, while there was no statistical significance difference between them regarding BMI, GA, and number of cesarean section (Table 3).
Table (3): Demographic data between three groups (normal, placenta previa and placenta accreta)

<table>
<thead>
<tr>
<th></th>
<th>Normal group</th>
<th>Placenta previa</th>
<th>Placenta accreta</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>No. = 20</strong></td>
<td>No. = 20</td>
<td>No. = 20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>Mean±SD</td>
<td>27.45 ± 2.68</td>
<td>27.75 ± 2.81</td>
<td>29.50 ± 2.21</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>23 – 33</td>
<td>23 – 33</td>
<td>25 – 33</td>
</tr>
<tr>
<td>BMI</td>
<td>Mean±SD</td>
<td>29.74 ± 1.36</td>
<td>29.52 ± 2.04</td>
<td>29.42 ± 1.45</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>27.8 – 31.9</td>
<td>26.4 – 33</td>
<td>26.5 – 31.5</td>
</tr>
<tr>
<td>GA</td>
<td>Mean±SD</td>
<td>32.15 ± 2.78</td>
<td>32.50 ± 2.56</td>
<td>32.25 ± 2.49</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>28 – 36</td>
<td>28 – 36</td>
<td>28 – 37</td>
</tr>
<tr>
<td>No. of cesarean section</td>
<td>Mean±SD</td>
<td>1.80 ± 0.70</td>
<td>2.15 ± 0.75</td>
<td>2.35 ± 0.67</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>1 – 3</td>
<td>1 – 3</td>
<td>1 – 3</td>
</tr>
</tbody>
</table>

*: One Way ANOVA test

There was no statistically significant difference between normal group and placenta previa group, while placenta accreta group was found with higher age than normal and placenta previa group with p.value 0.015 and 0.036 respectively (Table 4).

Table (4): Post Hoc analysis by LSD

<table>
<thead>
<tr>
<th></th>
<th>Post Hoc analysis by LSD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P1</td>
</tr>
<tr>
<td>Age</td>
<td>0.715</td>
</tr>
</tbody>
</table>

P1: Normal Vs Previa no significance
P2: Normal Vs Accreta significant
P3: Previa Vs Accreta significant

The best cut off point regarding serum amyloid A to differentiate between normal placenta patients and placenta accreta and previa patients was found > 15.4 with sensitivity of 77.5%, specificity of 95% and area under curve (AUC) of 90.7% (Figure 1).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>AUC</th>
<th>Cut of Point</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum Amyloid A</td>
<td>0.907</td>
<td>&gt;15.4</td>
<td>77.5</td>
<td>95.0</td>
<td>96.9</td>
<td>67.9</td>
</tr>
</tbody>
</table>

Figure (1): Receiver operating characteristic curve (ROC) for the serum amyloid A as a predictor for patients with placenta accrete and placenta previa
There was a statistically significant difference between three groups regarding level of serum amyloid A which was high in placenta previa group (Table 5).

Table (5): Level of serum Amyloid A in normal, placenta previa and placenta accreta

<table>
<thead>
<tr>
<th>Serum Amyloid A</th>
<th>Normal group</th>
<th>Placenta previa</th>
<th>Placenta accreta</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. = 20</td>
<td>No. = 20</td>
<td>No. = 20</td>
<td></td>
<td>0.000</td>
</tr>
<tr>
<td>Mean±SD</td>
<td>11.56 ± 2.19</td>
<td>20.60 ± 6.09</td>
<td>19.13 ± 5.37</td>
<td></td>
</tr>
<tr>
<td>Range</td>
<td>8.8 – 16.2</td>
<td>10.5 – 30.3</td>
<td>9.2 – 29.3</td>
<td></td>
</tr>
</tbody>
</table>

*: One Way ANOVA test

The best cut off point regarding serum amyloid A between normal placenta patients and placenta previa patients was found > 16.2 with sensitivity of 75.0%, specificity of 100.0% and area under curve (AUC) of 91.9%. The best cut off point regarding serum amyloid A between normal placenta patients and placenta accrete patients was found > 15.4 with sensitivity of 80.0%, specificity of 95.0% and area under curve (AUC) of 89.6% (Figure 2).

Figure (2): Receiver operating characteristic curve (ROC) for the serum amyloid A as a predictor for patients with placenta accrete and placenta previa from the normal placenta patients

There was no statistically correlation between serum amyloid A and age, BMI, GA and number of cesarean section (Table 6).

Table (6): Correlation between Serum Amyloid A and Demographic data in (All patients, Placenta previa, and Placenta accrete)

<table>
<thead>
<tr>
<th>Serum Amyloid A</th>
<th>All patients</th>
<th>Placenta previa</th>
<th>Placenta accreta</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>r</td>
<td>P-value</td>
<td>r</td>
</tr>
<tr>
<td>Age</td>
<td>0.249</td>
<td>0.121</td>
<td>0.211</td>
</tr>
<tr>
<td>BMI</td>
<td>0.050</td>
<td>0.757</td>
<td>-0.032</td>
</tr>
<tr>
<td>GA</td>
<td>0.205</td>
<td>0.205</td>
<td>0.107</td>
</tr>
<tr>
<td>Hysterotomy (CS)</td>
<td>0.188</td>
<td>0.246</td>
<td>0.190</td>
</tr>
</tbody>
</table>
DISCUSSION

This study aimed to use SAA as a biochemical marker in diagnosis of MAP with ultrasound & Doppler findings & if maternal serum serum amyloid A level served as an addition to ultrasonography in the diagnosis of MAP.

The current study is a prospective cohort study, conducted on (60) pregnant women who were divided into three equal groups as follows (20) women with placenta previa, (20) women with placenta accreta or increta and (20) normal pregnant women at a tertiary center: Al Azhar University Maternity Hospital during the period between February 2019 and October 2019.

As regards SAA assessment as single marker in patients with placenta previa, patients with placenta previa and patients with normal placenta we found that there was a statistically significant difference between three groups regarding level of serum amyloid A which was high in placenta previa group, The best cut off point regarding serum amyloid A to differentiate between normal placenta patients and placenta accreta and previa patients was found > 15.4 with sensitivity of 77.5%, specificity of 95%, PPV 96.9%, NPV 67.9% and area under curve (AUC) of 90.7%.

Which indicates that it had significant high diagnostic performance in differentiating placenta previa from normal with no significant difference between previa and accreta group.

Which disagree with the results of Shith et al (2009) who found that 3D power Doppler was the best single way for the diagnosis of placenta accreta, with a sensitivity of 97% and a specificity of 92%. 3D power Doppler gave the best PPV (76%), followed by gray-scale and color Doppler (47%). 3D power Doppler alone yielded the fewest false positive diagnoses, this difference with our study may be due to ultrasound interpreter.

These results were in agreement to the results of other studies as follows:

Cali et al. (2013) enrolled 187 patients with placenta previa and history of uterine surgery and performed transabdominal and transvaginal ultrasound examination for early detection of MAP with 2D gray-scale ultrasonography, & using transabdominal 3D power Doppler.

Ultrasound findings were reviewed against the final diagnosis made during Cesarean section.

In 41 cases, a hysterectomy was performed following confirmation of a lack of placental detachment. In each of the 41 confirmed cases, at least two of the sonographic criteria of invasion were present; with at most one of the criteria present in patients without MAP (morbidly adherent placenta) by 2D grayscale US, 3D power Doppler confirmed the gray-scale and color Doppler data. On the other hand, in the case of placenta previa without accreta, 3D rendering confirmed that the serosa-bladder wall interface was not affected by vascularization (Cali et al., 2013).

From my point of view, increase in PPV which represents increase in detection of true cases of MAP, may be the most important of all values in our situation.

This opens the field for the use of multi preoperative modalities in detection of
MAP, rather than use of either sonographic or biochemical marker alone.

As operative interventions in MAP need multidisciplinary team, it seems that accurate preoperative diagnosis of morbidly adherent placenta worthy to have a combination of more than one tool, aiming for proper preoperative preparation regarding transferral to tertiary hospital, preoperative ureteric stents insertion, available blood products, skilled obstetricians in caesarian hysterectomy & devascularization techniques.

Increase in detection rate of MAP makes preoperative preparations beneficial & cost effective.

Being cheap, available test, SAA assay when used in combination with 2DUS with 3DPD showed increase in detection rate of MAP.

SAA assay alone reliable, ultrasound examination results are good, combination of both is better, but we are searching for the ideal aiming to decrease maternal morbidity & mortality & to eliminate the fear associated with preoperative consent taken for hysterectomy in patients with no placental invasion.

CONCLUSION

Morbidly adherent placenta (MAP) is characterized by failure of the placenta to separate at delivery, with potential for significant perinatal and maternal morbidity and mortality. Antenatal diagnosis of placenta accreta, as it can reduce maternal morbidity by enabling for a scheduled delivery. Ultrasound identifies many, but not all, women with MAP. Detection of level of maternal SAA can be used as an additional tool to detect MAP. This opens the field for the use of multi pre-operative modalities in detection of MAP, rather than use of either sonographic or biochemical marker alone.

REFERENCES


صل الاميلويد أ كعلامة في تشخيص المشيمة الملتصقة
خالد زكريا الشيخه، عبد المنصف عبدالحميد صديق، أحمد عبد الحميد أبو زيد*، عمرو حسن محمد الشابي

قسم أغراض النساء والتوليد والباثولوجيا الأكلينية*، كلية الطب، جامعة الأزهر

خلفية البحث: تتميز المشيمة الملتصقة بتنوعها بصعوبة فصلها عند الولادة، مع احتمال حدوث نزيف ووفاة في الفترة المحيطة باليولادة. ارتفعت نسبة ظهور المشيمة بشكل مطرد إلى 1 في 532 بين عامي 1982 و2002. هذه الزيادة 13 مرة توازي الزيادة في الولادة القصيرة مع المشيمة المنزاحة. يوجد عوامل خطورة تؤدي إلى ظهور المشيمة الملتصقة وتوقع هذه العوامل قبل حدوث التصاق المشيمي مازال غامض وتحت الدراسة من هذه العوامل هجوم خلايا تروفوبلاست التي يطفئ الرحم وأيضاً ضعف طبقة بطانة الرحم المكونه للمشبهة يؤدي إلى زيادة احتماله التصاق المشيمي بها. لذلك تشخيص حالات التصاق المشيمي قبل الولادة يؤدي إلى تقليل عدد الوفيات اللتي تحدث بسبب هذا المرض. ويمكن تشخيص هذه الحالات عن طريق السونار ولكن ليس كل الحالات يمكن تشخيصها بالموجات فوق الصوتية فقط. يوجد هناك بروتين يسمى مصل الاميلويد أ هذا البروتين يمكن عن طريق تشخيص حالات المشيمي الملتصقة لانه عند حدوث اختراق لطبقة بطانة الرحم تزويد نسبه بالدم. يعتبر مصل الاميلويد أ واحد من عائلة من البروتينات الشحمية المرتبطة بالبروتين الدهني عالي الكثافة (HDL).

الهدف من البحث: الكشف عن التغييرات في الاميلويد أ المصل في النساء مع المشيمة الملتصقة وتقييم التغييرات في مصل الاميلويد أ في النساء مع المشيمة المنزاحة (المتقدمة) جانب نتائج الموجات فوق الصوتية والدوبير للكشف السابق للولادة للمشيمة الملتصقة.
المرضى وطريق البحث: أجريت الدراسة الحالية على (10) امرأة حامل قسمت إلى ثلاث مجموعات متساوية حيث يلي: المجموعة الأولى بها مشيمة منزاحة (متقدمة) والمجموعة الثانية: بها مشيمة ملتزمة و المجموعة الثالثة بها مشيمة عادية.

نتائج البحث: حدث انخفاض مصل الاميلويد A مع المشيمة المنزاحة (المقدمة) انخفاض ذو دلالات إحصائية عن المشيمة الملتزمة مما يدل على أنه يمكن استخدامه كوسيلة في التشخيص بجانب طريق التصوير فوق الصوتي.

الاستنتاج: الكشف عن مستوى مصل الاميلويد أ بالدم يمكن استخدامه كأداة إضافية للكشف عن مرض المشيمه الملتزمة.