Introduction

The Placenta Accreta Spectrum disorder (PAS) encompasses previous terms such as morbidly adherent placenta, placental invasion, and abnormally invasive place,[1] and is the most accepted term used in the clinical practice to standardize the terminology and is included in the Federation of Gynecology and Obstetrics (FIGO) consensus guidelines. The most common risk factors for PAS are prior cesarean delivery and placenta previa, with rising incidence in the past years.[2,3] Moreover, the risk of PAS is progressively increased with each Cesarean section (C-section).[4] With a likelihood of 11% of PAS in a patient with a single prior C-section and 61% in women with three prior C-sections.[5,6] Other risk factors include increasing maternal age and history of curettage or other uterine surgery.[3] PAS is associated with maternal morbidity and can lead to intrapartum hemorrhage and is a life-threatening situation for the mother and the fetus,[7,8] and the prenatal diagnosis of PAS is associated with reduction of these complications,[9,10] indicating its significant importance.

Ultrasound is the most widely used method for the diagnosis during the prenatal period given wide availability and relatively low-cost method and has a good accuracy mainly for the anterior uterine wall evaluation. Besides that, ultrasound allows a longitudinal assessment of women at risk for PAS increasing its possibility to reach the final diagnosis.[10] Magnetic Resonance Imaging (MRI) is usually performed only once as a secondary tool in cases in which ultrasound is inconclusive,[10] mainly for the evaluation of the posterior uterine wall,[7,11-15] and improves surgical planning by identifying the location of invasive placentation.[16] These different methods have already been compared how to stratify the surgical risk of women affected by PAS.[26]

Magnetic resonance imaging and previous cesarean section in placenta accrete spectrum disorder: Predictor model

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ABSTRACT

Objective: To evaluate objective criteria of Magnetic Resonance Imaging (MRI) of Placenta Accreta Spectrum disorder (PAS) analyzing interobserver agreement and to derive a model including imaging and clinical variables to predict PAS.

Methods: A retrospective review including patients submitted to MRI with suspicious findings of PAS on ultrasound. Exclusion criteria were lack of pathology or surgical information and missing or poor-quality MRI. Two radiologists analyzed six MRI features, and significant clinical data were also recorded. PAS confirmed on pathology or during intraoperative findings were considered positive for the primary outcome. Variables were tested through logistic regression models.

Results: Final study included 96 patients with a mean age of 33 years and 73.0% of previous C-sections. All MRI features were significantly associated with PAS for both readers. After logistic regression fit, including MRI signs with a moderate or higher interobserver agreement, intraplacental T2 dark band was the most significant radiologic criteria, and ROC analysis resulted in an AUC = 0.782. After including the most relevant clinical data (previous C-section) to the model, the ROC analysis improved to an AUC = 0.893.

Conclusion: Simplified objective criteria on MRI, including intraplacental T2 dark band associated with clinical information of previous C-sections, had the highest accuracy and was used for a predictive model of PAS.
Therefore, the objective of this study was to evaluate the objective criteria of MRI for the diagnosis of PAS, correlating to intraoperative findings and pathology and then creating a model to predict PAS, including imaging and clinical variables.

Materials and methods

Subjects

The Institutional Review Board (IRB) approved this retrospective study and granted a waiver of consent. This is a tertiary hospital, and the obstetric department is a reference for high-risk patients. Patients that are diagnosed by ultrasound with placenta previa and have the suspicion of PAS are routinely sent to MRI. Therefore, the authors included in the study pregnant women who performed an MRI under those circumstances during the period from July 2008 to December 2017. At last, the authors had a total of 110 patients, but 14 patients were not included in the analysis for different reasons, such as an anatomo-pathological study of the placenta not available (n = 2), surgical description not available (n = 3), MRI images not available (n = 2) and low-quality MRI images (n = 7) (Fig. 1).

The following clinical data were also recorded: maternal age, gestational age at MRI, gestational age at delivery, the time between MRI and delivery, ethnic group, parity, previous C-sections, abortions, history of curettage, and other uterine surgeries. PAS confirmed on the pathology of surgical specimen and/or signs of PAS during intraoperative findings were considered as positive for the primary outcome (Table 1).

Image acquisition

All MRI examinations were performed on a 1.5T unit (Signa HD × TM, General Electric Healthcare) with body array coils, including axial, coronal, and sagittal T2-weighted Single Shot Fast Spin-Echo (SS-FSE); axial, coronal, and sagittal balanced steady-state free precession (Fast Imaging Employing Steady-State Acquisition – FIESTA); axial and sagittal fat-suppressed ultra-fast spoiled gradient-echo T1-weighted (LAVA) without intravenous contrast media.

The parameters for SS-FSE images were Repetition Time (TR)/Echo Time (TE), 650–16000 / 50–90 msec; Flip Angles (FA), 90; Field of View (FOV), 120–480 mm; slice thickness, 4.5 mm. For balanced – steady-state free-precession – FIESTA images, they were TR/TE, 3800–6000/ 50–90 msec; FA = 90; FOV, 120–480 mm; slice thickness, 4.5 mm, and for T1 LAVA images, they were: TR/TE, 200–230 / 1.8–11 msec; FA = 80; FOV, 340–380 mm; slice thickness, 4.5 mm.

Table 1

| Clinical data                        | Mean ± SD or n (%) |
|--------------------------------------|--------------------|
| Maternal Age (years)                 | 33.65 ± 5.39       |
| Gestational age at MRI (weeks)       | 32.20 ± 3.79       |
| Gestational age at delivery (weeks)  | 35.70 ± 2.68       |
| Race                                 |                    |
| · White                               | 77 (80.2%)         |
| · Nonwhite                            | 16 (16.7%)         |
| · Uninformed                          | 3 (3.10%)          |
| Parity                               |                    |
| · 1                                   | 11 (11.5%)         |
| · 2                                   | 21 (21.9%)         |
| · 3                                   | 18 (18.8%)         |
| · 4                                   | 22 (22.9%)         |
| ≥5                                   | 24 (25.0%)         |
| Abortions                            |                    |
| · 0                                   | 53 (55.2%)         |
| · 1                                   | 28 (29.2%)         |
| · 2                                   | 10 (10.4%)         |
| · 3                                   | 5 (5.20%)          |
| Previous C-section                   |                    |
| · 0                                   | 27 (28.1%)         |
| · 1                                   | 33 (34.4%)         |
| · 2                                   | 18 (18.8%)         |
| · 3                                   | 15 (15.6%)         |
| · 4                                   | 3 (3.10%)          |
| Other surgeries (myomectomy)         |                    |
| · 0                                   | 93 (96.6%)         |
| · 1                                   | 3 (3.10%)          |
| Curettage                            |                    |
| · 0                                   | 63 (66.6%)         |
| · 1                                   | 23 (24.0%)         |
| · 2                                   | 5 (5.20%)          |
| · 3                                   | 5 (5.20%)          |

SD, Standard Deviation; n, number.

Image interpretation

Two board-certified blinded radiologists with different levels of expertise (2 and 8 years of experience in abdominal radiology) retrospectively analyzed the exams on a Likert scale[1–5] for each sign of PAS.

The MRI signs used for evaluation were:

1- Intraplacental abnormal vascularity: tortuous enlarged flow voids observed on T2 and corresponding high signal on FIESTA, indicating vascular flow[18] (Fig. 2).

- 5: ≥ 6mm
- 4: 5–6 mm
- 3: 3–4 mm
- 2: 1–2 mm
- 1: absent

Fig. 1. Flowchart of the study population. *PACS (Picture Archiving and Communication System).

Fig. 2. 28-year-old woman at 34 weeks gestation with two previous C-sections. (A) Sagittal T2 weighted image shows intraplacental abnormal vascularity with a flow void in the placenta (arrow), and on (B) axial FIESTA there is a high signal on the structure indicating the presence of vascular flow.
Intraplacental T2 dark band: wedge-shaped areas of low signal intensity on T2-weighted images and FIESTA [19] (Figs. 3 and 4).

- 5: > 20 mm
- 4: 15-20 mm
- 3: 10-14 mm
- 2: < 10 mm
- 1: absent

2- Placental bulge: focal bulging of the uterine contour. [14,19,24,27]
- 5: clear loss of myometrial congruency or invasion of adjacent organs
- 4: focal bulges
- 3: disruption of the normal pear shape of the uterus with the lower uterine segment being wider than the fundus
- 2: slight irregularities on uterine contours
- 1: normal contours

3- Placental protrusion sign: placenta pushing inferior to the internal os.[21]
- 5: > 15 mm
- 4: 11-15 mm
- 3: 6-10 mm
- 2: < 5 mm
- 1: absent

4- Myometrial thinning: focal thinning and indistinctness of the myometrium and loss of thin dark uteroplacental interface on T2-weighted images.[18,28,29]

5- Heterogeneous placenta: marked heterogeneous intensity within the placenta (Fig. 5).[18,30-32]

Fig. 3. 34-year-old woman at 36 weeks' gestation and one previous C-section. (A) Sagittal T2 weighted image shows intraplacental T2 dark band (arrow), and on (B) sagittal FIESTA the band remains with the low signal intensity.

Fig. 4. 42-year-old woman at 35 weeks' gestation and one previous C-section. (A) Axial T2 weighted images shows intraplacental T2 dark band (arrow) and placental bulges (dashed arrows).

Fig. 5. 35-year-old woman at 35 weeks' gestation and two previous C-sections. (A) Sagittal T2 weighted image shows a heterogeneous placenta (arrows).

**Statistical analysis**

**Pathological and surgical data**

The surgical description had three outcomes: normal discharge of the placenta, focal attachment (small bleeding during placental discharge but no clinical relevance/no need for blood transfusions or other intervention), and hysterectomy. Only hysterectomy was considered positive for PAS. For pathological analyses, there were five outcomes: normal placenta, focal accretion, accreta, increta and percreta. Normal placenta and focal accretion were considered negative for pathological PAS.

**Clinical and imaging analyses**

Descriptive analyses of the data were performed using absolute and relative frequencies, central tendency, and dispersion measures. The comparison between the groups of PAS and no PAS was made with a Mann-Whitney test for non-parametrical variables and $t$-Student test for parametrical variables.

The interobserver variability of all MRI signs was assessed using the Kappa ($\kappa$) value. A $\kappa$-value of $\leq 0.20$ was interpreted as slight agreement, $0.21-0.40$ as fair agreement, $0.41-0.60$ as moderate agreement, $0.61-0.80$ as substantial agreement, and $\geq 0.81$ as almost perfect agreement.[33]

For the development of the predictor model, both readers reassessed the discordant criteria and reviewed it in consensus. This strategy was made to extract the best possible information from the MRI and build the best predictor model possible. To identify which MRI signs and clinical variables were most significant, analyses through multiple binary logistic regression models were made in other to identify its coefficients, Odds Ratios (OR), and their respective 95% Confidence Intervals (95% CI). The significant variables ($p < 0.005$) were tested on the multiple modeling and those which presented a value of $p < 0.20$ by the Stepwise technique.

Receiver Operating Characteristic (ROC) analysis was used for evaluation of the diagnostic performance using only the most significant MRI sign and using the most significant MRI sign combined with the most significant clinical data for PAS. The areas under the ROC Curves (AUC) were estimated nonparametrically for non-ordinal score assessments.
Analyses of clinical data and association with placental acretism.

| Clinical variables                                      | PAS       | No PAS    | p      |
|---------------------------------------------------------|-----------|-----------|--------|
| Maternal Age (years), mean (SD)                         | 33.7 ± 5.4| 32.9 ± 5.6| 0.486* |
| Gestational age at MRI (weeks), mean (SD)               | 31.5 ± 4.7| 32.6 ± 2.8| 0.194* |
| Gestational age at delivery (weeks), mean (SD)          | 35.2 ± 3.2| 36.1 ± 2.0| 0.114* |
| Time interval between MRI and delivery (weeks), mean (SD)| 3.6 ± 3.4| 3.4 ± 2.4| 0.808* |
| Parity, median (min-max)                                | 2 (0–8)   | 1 (0–5)   | <0.0001 |
| Previous C-Section, median (min-max)                    | 2 (0–4)   | 1 (0–3)   | <0.0001 |
| Abortions, median (min-max)                             | 0 (0–3)   | 0 (0–3)   | 0.464* |
| Curettage, median (min-max)                             | 0 (0–3)   | 0 (0–3)   | 0.061* |

SD, Standard Deviation; min, minimum; max, maximum.

* t-Student test.

b Mann-Whitney test.

Analyses were conducted using statistical software (Statistical Package for the Social Science – SPSS, version 20.0 for Windows).

Results

The final sample of 96 patients was divided into two groups according to anatomopathological study and/or surgical descriptions: patients with PAS (n = 42) and patients without PAS (n = 54) (Table 1).

Clinical variables

For all the clinical variables analyzed, only parity and previous C-section showed significant differences between the PAS and no PAS groups. All other variables: abortions, curettage, maternal age, gestational age at MRI, gestational age at delivery, and time interval between MRI and delivery showed no significant differences between the PAS and no PAS groups (Table 2).

Differences in each MRI sign between PAS and no PAS groups and diagnostic performance

All MRI signs (intraplacental abnormal vascularity, intraplacental T2 dark band, placental bulge, placental protrusion sign, myometrial thinning, and heterogeneous placenta) showed significant differences between the PAS and no PAS groups for both reader A and reader B (Table 3).

The Sensitivity (SEN), Specificity (SPE), Positive Predictive Value (PPV), and Negative Predictive Value (NPV) for each MRI sign are shown in (Table 3).

Interobserver agreement

Concordance coefficient values are shown in (Table 4). Intraplacental T2 dark band had a substantial agreement between reader A and reader B, moderate agreement for abnormal vascularity and heterogeneity, poor agreement for placental bulge and myometrial thinning, and slight agreement for placental protrusion sign.

Predictor model

Initially, in other to create a reproducible predictor model, the authors excluded the MRI signs that showed a slight agreement (κ-value of ≤ 0.20) and fair agreement (κ-value of 0.21–0.40) in the interobserver analysis.

The remaining MRI signs (heterogeneity, abnormal vasculation, and intraplacental T2 dark band) and the most relevant clinical sign

Table 2
Analyses of clinical data and association with placental acretism.

| Clinical variables                                      | PAS       | No PAS    | p      |
|---------------------------------------------------------|-----------|-----------|--------|
| Maternal Age (years), mean (SD)                         | 33.7 ± 5.4| 32.9 ± 5.6| 0.486* |
| Gestational age at MRI (weeks), mean (SD)               | 31.5 ± 4.7| 32.6 ± 2.8| 0.194* |
| Gestational age at delivery (weeks), mean (SD)          | 35.2 ± 3.2| 36.1 ± 2.0| 0.114* |
| Time interval between MRI and delivery (weeks), mean (SD)| 3.6 ± 3.4| 3.4 ± 2.4| 0.808* |
| Parity, median (min-max)                                | 2 (0–8)   | 1 (0–5)   | <0.0001 |
| Previous C-Section, median (min-max)                    | 2 (0–4)   | 1 (0–3)   | <0.0001 |
| Abortions, median (min-max)                             | 0 (0–3)   | 0 (0–3)   | 0.464* |
| Curettage, median (min-max)                             | 0 (0–3)   | 0 (0–3)   | 0.061* |

SD, Standard Deviation; min, minimum; max, maximum.

* t-Student test.

b Mann-Whitney test.

Analyses were conducted using statistical software (Statistical Package for the Social Science – SPSS, version 20.0 for Windows).

Table 3
MRI feature analyses and performance for each radiologist.

| Radiologist A          | PAS median (min–max) | No PAS median (min–max) | p*      | SEN   | SPE   | PPV   | NPV   |
|------------------------|----------------------|-------------------------|---------|-------|-------|-------|-------|
| Intraplacental T2 dark | 5 (1–5)              | 1 (1–5)                 | <0.0001 | 73.8% | 74.1% | 68.9% | 78.4% |
| Abnormal vascularity   | 5 (1–5)              | 2.5 (1–5)               | <0.0001 | 81.0% | 50.0% | 55.7% | 77.1% |
| Placental Bulge        | 4 (1–5)              | 3 (1–5)                 | <0.0001 | 95.2% | 37.0% | 54.1% | 99.9% |
| Heterogeneity          | 4 (1–5)              | 2 (1–5)                 | <0.0001 | 78.6% | 55.6% | 57.9% | 76.9% |
| Myometrial thinning    | 5 (1–5)              | 3 (1–5)                 | <0.0001 | 95.2% | 44.4% | 57.1% | 92.3% |
| Placental protrusion   | 2 (1–5)              | 1 (1–4)                 | <0.0001 | 33.3% | 90.7% | 73.7% | 63.6% |
| Radiologist B          |                      |                         |         |       |       |       |       |
| Intraplacental T2 dark | 4.5 (1–5)            | 1 (1–5)                 | <0.0001 | 76.2% | 75.9% | 71.1% | 80.4% |
| Abnormal vascularity   | 5 (1–5)              | 2 (1–5)                 | <0.0001 | 81.0% | 63.0% | 63.0% | 81.0% |
| Placental Bulge        | 4 (1–5)              | 2 (1–4)                 | <0.0001 | 85.7% | 72.2% | 70.6% | 86.7% |
| Heterogeneity          | 4 (1–5)              | 2 (1–4)                 | <0.0001 | 81.0% | 57.4% | 59.6% | 79.5% |
| Myometrial thinning    | 4 (2–5)              | 3 (1–5)                 | <0.0001 | 83.3% | 72.2% | 70.0% | 84.8% |
| Placental protrusion   | 2 (1–5)              | 1 (1–4)                 | <0.0001 | 45.2% | 96.3% | 90.5% | 69.3% |

Min, minimum; max, maximum; SEN, Sensitivity; SPE, Specificity; PPV, Positive Predictive Value; NPV, Negative Predictive Value.

* Mann-Whitney test.

Concordance coefficient values are shown in (Table 4). Intraplacental T2 dark band had a substantial agreement between reader A and reader B, moderate agreement for abnormal vascularity and heterogeneity, poor agreement for placental bulge and myometrial thinning, and slight agreement for placental protrusion sign.

Predictor model

Initially, in other to create a reproducible predictor model, the authors excluded the MRI signs that showed a slight agreement (κ-value of ≤ 0.20) and fair agreement (κ-value of 0.21–0.40) in the interobserver analysis.

The remaining MRI signs (heterogeneity, abnormal vasculation, and intraplacental T2 dark band) and the most relevant clinical sign

| MRI signs               | Radiologists A and B (Kappa coefficient) |
|------------------------|------------------------------------------|
| Intraplacental T2 dark | 0.749                                    |
| Abnormal vascularity   | 0.547                                    |
| Placental bulge        | 0.332                                    |
| Heterogeneity          | 0.482                                    |
| Myometrial thinning    | 0.363                                    |
| Placental protrusion   | 0.179                                    |

A κ-value of ≤ 0.20 was interpreted as slight agreement, 0.21–0.40 fair agreement, 0.41–0.60 as moderate agreement, 0.61–0.80 as substantial agreement and ≥ 0.81 as almost perfect agreement.
(previous C-section) were analyzed using a multiple binary logistic regression model. The most significant clinical variable in isolation was the presence of a previous C-section with an OR = 3.35 (95% CI 0.68–5.97), and the most significant MRI sign was intraplacental T2 dark band with an OR = 12.67 (95% CI 3.97–40.45) as shown in Table 5.

The result of ROC analyses for the prediction of PAS using only the most significant MRI sign (intraplacental T2 dark band) and using the most significant MRI sign combined with the most significant clinical data (previous C-sections) are shown in Fig. 6. For the prediction using only the most significant MRI sign, the area under the curve is 0.782 (95% CI 0.685–0.878), and for the prediction of the most significant MRI sign combined with previous C-sections, the area under the curve is 0.893 (95% CI 0.829–0.957).

Finally, the authors derived a predictor model for the estimated PAS probability according to the number of previous C-sections for patients that had or not the presence of the intraplacental T2 dark band sign (Fig. 6).

Discussion

The analysis in the present study has confirmed that MRI signs of PAS (intraplacental abnormal vascularity, Intraplacental T2 dark band, placental bulge, placental protrusion sign, myometrial thinning, and heterogeneous placenta) were associated with the presence of PAS for both inexperienced and experienced radiologists as observed in previously published studies.

The authors have also evaluated the interobserver variability of MRI analyses of experienced and less experienced radiologists, knowing that the previous experience of the radiologist improves the diagnostic performance, as shown by Ghezzi et al.[34] The authors found a substantial agreement for the intraplacental T2 dark band and a moderate agreement for abnormal vascularity, similarly to previous reports. [19,21,22,28,35,36] The heterogeneity sign also had moderate agreement even though the authors understand that this finding is very influenced by subjective evaluation as resembled on previous studies, [22,28] and it is known that normal placenta can show some heterogeneity. [37] The placental bulge had a fair agreement, a variability slightly higher than previously reported. [19,22,35] Myometrial thinning also had fair agreement similar to what was reported by Lax et al. [19] and in correlation with the knowledge that normal myometrium can become thin during pregnancy, especially in the third trimester and when the placenta was posterior location. [19,20,27] Placental protrusion sign had a slight agreement and had a low incidence as previously reported by Bourgioti et al. [36] fact that can explain its low agreement and limit its use once it is a rare finding in a rare disease, making it a difficult sign to be studied and correctly interpreted. These findings indicate the need for objective standardization of MRI assessment of PAS disorders that can provide a more reproductive interpretation of the exam [17] as suggested by the consensus of Jha et al. [38] and potentially reduce the effect of the radiologists’ previous experience. [34]

Regarding clinical variables, they can be important risk factors for the development of PAS, mainly in relation to the previous C-section, as previously shown. [4–6] In the present study, radiologists were blinded for clinical information, but the authors understand that the availability of this information to the radiologist at the moment of the exam evaluation is extremely important for the elaboration of his final diagnosis.

The predictive model, including only MRI sign (intraplacental T2 dark band), had great accuracy (AUC = 0.782), which was further improved (AUC = 0.893) when adding previous C-sections. These results demonstrate that this model can be useful for obstetricians to evaluate the estimated probability of PAS on delivery very easily so that they can properly manage it in advance and are not surprised with a potential high-risk procedure.

Table 5
Multiple binary logistic regression models of independently most significant variables.

| Variables                  | Coefficient | SE  | OR adj | 95% IC (inf-sup) | p        |
|----------------------------|-------------|-----|--------|------------------|----------|
| Initial analysis           |             |     |        |                  |          |
| Intercept                  | -3.1        | 0.6 |        |                  |          |
| Heterogeneity              | -1.0        | 1.1 | 0.3    | 0.04–3.4         | 0.878    |
| Abnormal vascularity       | 0.7         | 0.6 | 2.0    | 0.5–7.4          | 0.266    |
| Intraplacental dark T2 band| 2.9         | 1.2 | 19.4   | 1.8–210          | 0.014    |
| Previous C-sections       | 1.2         | 0.3 | 3.4    | 1.8–5.9          | <0.0001  |
| Final analysis             |             |     |        |                  |          |
| Intercept                  | -3.1        | 0.6 |        |                  |          |
| Intraplacental dark T2 band| 2.5         | 0.6 | 12.6   | 3.9–40.4         | <0.0001  |
| Previous C-sections       | 1.2         | 0.3 | 3.4    | 1.8–5.9          | <0.0001  |

SE, Standard Error; adj, adjusted; inf, inferior; Sup, Superior; p <0.05.

Fig. 6. (A) ROC curve analyses for the prediction of PAS including only MRI sign (intraplacental T2 dark band) with an AUC = 0.782 (95% CI 0.685–0.878) and (B) including MRI sign and relevant clinical data (previous C-sections) with an AUC = 0.893 (95% CI 0.829–0.957). (C) Predictor model for the estimated PAS probability versus number of previous C-sections of the patients.
There are some limitations of the present study. To begin with, it was performed in a tertiary hospital with high-risk patients selected from abnormal ultrasound, inducing selection bias. Moreover, radiologists in the study were well trained for this rare condition, given the particularity of the hospital, and may not reflect standard radiology practice. Some of the advantages of the study are its high number of patients included compared with previous studies and the use of a gold standard combining surgical description and pathological analyses. According to the sample, the results must be analyzed carefully. The reduced sample size contributes to reproducibility issues, including false positives and false negatives. However, the model was analyzed considering the number of outcomes to avoid an overfitting model.

Conclusion

Simplified objective criteria on MRI (intraplacental T2 dark band) combined with clinical data (previous C-sections) contributed to the creation of a predictive model, which the authors believe can facilitate and improve the diagnostic accuracy by providing a more objective result for the MRI report. Besides, it allows a higher uniformization of the analyses between radiologists with different levels of expertise and facilitates interobserver agreement in MRI assessment of severity of placenta accreta spectrum disorders. Ultrasound Obstet Gynecol 2020;55(4):467–73.

References

1. Clark HR, Ng TW, Khan A, Happe S, Dashe J, Xi Y, et al. Placenta accreta spectrum: correlation of MRI parameters with pathological and surgical outcomes of high-risk pregnancies. AJR Am J Roentgenol 2020;214(6):1417–23.
2. Higgins MF, Monteith C, Foley M, O'connor P, Canning M, Ehrlich Y. Real increasing incidence of hysterectomy for placenta accreta following previous caesarean section. Eur J Obstet Gynecol Reprod Biol 2013;171(1):54–6.
3. Wu S, Kocherginsky M, Hibbard JU. Abnormal placentation: twenty-year analysis. Am J Obstet Gynecol 2005;192(2):458–61.
4. Iacovelli A, Liberiati M, Khalil A, Timer-Trisch I, Leombroni M, Buca D, et al. Risk factors for abnormally invasive placenta: a systematic review and meta-analysis. J Matern Fetal Neonatal Med 2020;33(3):471–81.
5. Hall AD, Moore TR. Multiple repeat cesareans and the threat of placenta accreta: incidence, diagnosis, management. Clin Perinatol 2011;38(2):285–96.
6. Silver RM, Landon MB, Rouse DJ, Leveno KJ, Spong CY, Thom EA, et al. Maternal morbidity associated with multiple repeat Cesarean Deliveries. Obstet Gynecol 2006;107(6):1226–32.
7. Committee opinion no. 501: Maternal-fetal intervention and fetal care centers. Obstet Gynecol 2011;118(2 Pt 1):405–10.
8. Oyelese Y, Previta Smulian JCPlacenta, Accreta Placenta, Previa Vasa. Obstet Gynecol 2006;108(3 Part 1):694.
9. Buca D, Liberiati M, Cali G, Forlani F, Caisutti G, Flacco ME, et al. Influence of prenatal diagnosis of abnormally invasive placenta on maternal outcome: systematic review and meta-analysis. Ultrasound Obstet Gynecol 2016;52(3):304–9.
10. Familiari A, Liberati M, Lim P, Pagani G, Cali G, Buca D, et al. Diagnostic accuracy of magnetic resonance imaging in detecting the severity of abnormal invasive placenta: a systematic review and meta-analysis. Acta Obstet Gynecol Scand 2018;97(3):507–20.
11. Fuentes E, Alfrevic Z, Bhide AG, Belfort MA, Burton GJ, Collins SI, et al. Placenta praevia and placenta accreta: diagnosis and management: green-top guideline No. 27a. BJOG 2019;126(1):e1–e48.
12. Maher MA, Abdelaziz A, Bazeed MF. Diagnostic accuracy of ultrasound and MRI in the prenatal diagnosis of placenta praevia. Acta Obstet Gynecol Scand 2013;92(9):1017–22.
13. Palacios Jaraquemada JM, Bruno CH. Magnetic resonance imaging in 300 cases of placenta accreta: surgical correlation of new findings. Acta Obstet Gynecol Scand 2005;84(8):716–24.
14. Collins SI, Ashcroft A, Braun T, Calda P, Langhoef-Roos J, Morel O, et al. Proposal for standardized description of abnormally invasive placenta (AIP). Ultrasound Obstet Gynecol 2016;47(3):271–5.
15. Jimaraina E, Bhide A, Kennedy A, Woodward P, Hubinont C, Collins SI, et al. FIGO consensus guidelines on placenta accreta spectrum disorders: Prenatal diagnosis and screening. Int J Gynecol Obstet 2018;140(3):274–80.
16. Masselli G, Brundelli R, Casciani E, Polettiu E, Piccinii MG, Anzecchi M, et al. Magnetic resonance imaging in the evaluation of placental adhesive disorders: correlation with color Doppler ultrasound. Eur Radiol 2008;18(6):1292–9.
17. Finazzo F, Fantonio F, Masselli G, Forlani F, Palacios-Jaraquemada J, Minazzi G, et al. Interobserver agreement in MRI assessment of severity of placenta accreta spectrum disorders. Ultrasound Obstet Gynecol 2020;55(4):467–73.
18. Derman AY, Nickar K, Herbanek S, Zelenko N, Opha S, Flyer M. MRI of placenta accreta: a new imaging perspective. AJR Am J Roentgenol 2011;197(6):1514–21.
19. Lan A, Prince MR, Memmit KW, Schwebach JR, Budrick NE. The value of specific MRI features in the evaluation of suspected placental invasion. Magn Reson Imaging 2007;25(1):87–93.
20. Allen BC, Leyendecker JR. Placental evaluation with magnetic resonance. Radiol Clin North Am 2012;51(6):955–66.
21. Ueno Y, Kitajima K, Wakawami F, Maeda T, Suwana Y, Takahashi S, et al. Novel MRI finding for diagnosis of invasive placenta praevia: evaluation of findings for 65 patients using clinical and histopathological correlations. Eur Radiol 2014;24(8):3811–8.
22. Ueno Y, Maeda T, Tanaka U, Naimura K, Kitajima K, Suwana Y, et al. Evaluation of interobserver variability and diagnostic performance of developed MRI-based radiological scoring system for invasive placenta previa. J Magn Reson Imaging 2016;44(3):573–83.
23. D'Antonio F, Iacovella C, Palacios-Jaraquemada J, Bruno CH, Manzoli L, Bhide A. Pre-natal identification of invasive placenta placentation using magnetic resonance imaging: systematic review and meta-analysis. Ultrasound Obstet Gynecol 2014;44(1):8–16.
24. Teo TH, Law YM, Tay KH, Tan BS, Cheah FK. Use of magnetic resonance imaging in evaluation of placental invasion. Clin Radiol 2009;64(5):511–6.
25. Kilcoyne A, Shenoy-Bhangle AS, Roberts DJ, Soxinda RC, Gervais DA, Lee SL. MRI of placenta accreta, placenta increta, and placenta percreta: pearls and pitfalls. AJR Am J Roentgenol 2017;208(1):214–21.
26. Palacios-Jaraquemada JM, D'Antonio F, Buca D, Fiorillo A, Larraza P. Systematic review on near miss cases of placenta accreta spectrum disorders: correlation with invasion topography, prenatal imaging, and surgical outcome. J Matern Fetal Neonatal Med 2020;33(19):377–84.
27. Baughman WC, Corteville JE, Shah RR. Placenta accreta spectrum: a review of US and MR imaging findings. Radiographics 2008;28(7):1905–16.
28. Bour L, Place V, Bensdavid S, Fargeaudho Y, Portal JJ, Ricbourg A, et al. Suspected invasive placenta evaluation with magnetic resonance imaging. Eur Radiol 2014;24(12):3150–60.
29. Maldjian C, Adam R, Pelosi M, Pelosi M 3rd, Rudelli RD, Maldjian J. MRI appearance of placenta percreta and placenta accreta. Magn Reson Imaging 1999;17(7):965–71.
30. Elsayes KM, Trout AT, Friedkin AM, Liu PS, Bude RO, Platt JF, et al. Imaging of placental invasion. Clin Radiol 2009;64(5):511–6.
31. Masselli G, Guaidi G. MR imaging of the placenta: what a radiologist should know. Abdom Imaging 2013;38(3):573–87.
32. Varghese B, Singh N, George RA, Gilvaz S. Magnetic resonance imaging of placenta accreta. Indian J Radiol Imaging 2013;23(4):379–85.
33. Landis JR, Koch GG. The measurement of observer agreement for categorical data. Biometrics 1977;33(2):159–74.
34. Grezzi CLA, Silva OK, Casagrande AS, Westphalen SS, Salazar CC, Vettorazzi J. Diagnostic performance of radiologists with different levels of experience in the interpretation of MRI of the placenta accreta spectrum disorder. Br J Radiol 2021;94(1128):20210827.
35. Sato T, Mori N, Hasegawa O, Shigihara T, Fujimori K, Taiho K, et al. Placental recess finding for diagnosing placental invasion. Abdom Radiol (NY) 2019;44(2):W41–6.
36. Naderi M, Sabour S. MRI features predictive of invasive placenta with extrauterine spread in high-risk gravid patients: methodologic issues. AJR Am J Roentgenol 2019;212(2):W41.
37. Blaicher W, Bruggner PC, Mittermayer C, Schwindt J, Deutinger J, Bernaschek G, et al. Color Doppler MRI parameters in the evaluation of placental invasion. AJR Am J Roentgenol 2011;196(2):304–10.