BMJ Open Epidemiology of placenta previa accreta: a systematic review and meta-analysis

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ABSTRACT

Objective To estimate the prevalence and incidence of placenta previa complicated by placenta accreta spectrum (PAS) and to examine the different criteria being used for the diagnosis.

Design Systematic review and meta-analysis.

Data sources PubMed, Google Scholar, ClinicalTrials.gov and MEDLINE were searched between August 1982 and September 2018.

Eligibility criteria Studies reporting on placenta previa complicated by PAS diagnosed in a defined obstetric population.

Data extraction and synthesis Two independent reviewers performed the data extraction using a predefined protocol and assessed the risk of bias using the Newcastle-Ottawa scale for observational studies, with difference agreed by consensus. The primary outcomes were overall prevalence of placenta previa, incidence of PAS according to the type of placenta previa and the reported clinical outcomes, including the number of peripartum hysterectomies and direct maternal mortality. The secondary outcomes included the criteria used for the prenatal ultrasound diagnosis of placenta previa and the criteria used to diagnose and grade PAS at birth.

Results A total of 258 articles were reviewed and 13 retrospective and 7 prospective studies were included in the analysis, which reported on 587 women with placenta previa and PAS. The meta-analysis indicated a significant (p<0.001) heterogeneity between study estimates for the prevalence of placenta previa, the prevalence of placenta previa with PAS and the incidence of PAS in the placenta previa cohort. The median prevalence of placenta previa was 0.56% (IQR 0.39–1.24) whereas the median prevalence of placenta previa with PAS was 0.07% (IQR 0.05–0.16). The incidence of PAS in women with a placenta previa was 11.10% (IQR 7.65–17.35).

Conclusions The high heterogeneity in qualitative and diagnostic data between studies emphasises the need to implement standardised protocols for the diagnoses of both placenta previa and PAS, including the type of placenta previa and grade of villous invasiveness.

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INTRODUCTION

Placenta accreta is a pathological condition of placentation associated with a high risk of massive obstetric haemorrhage during delivery. Initially described in 1937 by Irving and Hertig as the abnormal adherence of the placenta to the myometrium due to the partial or complete absence of decidua basalis, it was subsequently redefined by Luke et al as a spectrum of abnormally adherent and invasive placentation disorders. Placenta accreta is now graded according to the depth of the villous penetration into the uterine wall starting with the abnormally adherent placenta or creta, where the villi attach directly to the surface of the myometrium without invading it, and extending to the invasive grades of placenta increta, where the villi penetrate deeply into the myometrium up to the uterine serosa, and placenta percreta, where the invasive villous tissue penetrates through the uterine serosa often entering the surrounding pelvic tissues.

The different grades of the placenta accreta spectrum (PAS) can coexist in the same specimen and can be focal (just a small area of the placental bed) or extensive (including much of the placental bed). Over the last two decades, a growing body of epidemiology research has identified the effect of the rapid increase in caesarean delivery rates on the risks of PAS. The main additional risk factor after a previous caesarean delivery is placenta previa. A large multicentric US cohort study noted that for women presenting...
with placenta previa and prior caesarean delivery, the risk of PAS was 3%, 11%, 40%, 61% and 67% for first, second, third, fourth and fifth or more caesarean deliveries, respectively. A national case–control study using the UK Obstetric Surveillance System found that the incidence of PAS increases from 1.7 per 10000 births overall to 577 per 10000 births in women with both a previous caesarean delivery and placenta previa.8

Both abnormal adherence and invasion of villous tissue into the myometrium result in failure of the placenta to separate spontaneously from the uterine wall at delivery.2–4 When unsuspected at the time of delivery, attempts to manually remove accreta villous tissue typically provoke rapid bleeding from the uteroplacental circulation.5,11 In invasive cases, this can lead to massive obstetric haemorrhage due to the disruption of the deep uterine vascularature of the increta or percreta area.4,9 Not surprisingly, prenatal diagnosis of PAS has been shown to decrease maternal morbidity and mortality, and has thus become essential in improving its management.12,13 Tabsh et al were the first in 1982 to report on the prenatal ultrasound diagnosis of a case of placenta increta.14 A recent systematic review and meta-analysis of prenatal ultrasound diagnosis of placenta previa with PAS in women with a history of caesarean delivery has found that the overall diagnostic accuracy of ultrasound in specialist units is in 90.9%.15 However, in countries with well-established screening programmes for fetal anomalies, over half the cases of PAS are not diagnosed before delivery.3,10,16

Accreta placentation and in particular its invasive forms are impacting maternal health outcomes globally and its prevalence is likely to increase. Women with a history of previous caesarean delivery presenting with placenta previa complicated by PAS in an ongoing pregnancy are now the cohort of obstetric patients with the highest risk of delivery complications,16 however, their epidemiology has not been comprehensively reviewed yet. Health provision for the development of maternity centres with specialist teams, equipment, drugs, blood bank and intensive care infrastructure to safely manage women presenting with placenta previa and PAS requires an accurate evaluation of its epidemiology. The objective of this meta-analysis is to review the epidemiology of women presenting with placenta previa and to examine the different criteria used by the authors of cohort studies to diagnose placenta previa and PAS prenatally and to confirm the diagnosis of PAS at birth.

**MATERIALS AND METHODS**

A systematic review was undertaken of articles providing data on prevalence and incidence of PAS in women presenting with a placenta previa where the populations sampled were defined. PubMed, Google Scholar, ClinicalTrials.gov and MEDLINE were searched for studies published between the first prenatal ultrasound description of placenta accreta in August 1982 by Tabsh et al.8 and September 2018. The overall search strategy was inclusive of MeSH headings for the following terms: ‘placenta accreta, placenta increta, placenta percreta, abnormally invasive placenta, morbidly adherent placenta and major placenta previa’ (search strategy in online supplementary data 1). Title, abstracts and full text were independently assessed by the authors for content, data extraction and analysis. Additional relevant studies were identified from reference lists of reviews and editorials and by handsearching key journals and websites. All search results were combined in a reference database. Duplicates were removed by hand. The search was limited to articles published in English.

Two independent investigators (EJ and LG) selected studies in two stages. The abstracts of all potentially relevant papers were individually examined for suitability. Papers were only ruled out at this stage if they obviously did not meet the inclusion criteria. The remaining were obtained in full text and were independently assessed for content, data extraction and analysis. Disagreements between the two original reviewers were resolved by discussion with the third investigator (JL-R). Articles were excluded if; they were published before August 1982, contained no data on the study population such as the overall pregnancies, births and/or deliveries numbers, were case reports or were overlapping.

Study characteristics were extracted using a pre-designed data extraction protocol including: author institution, year of publication, country of origin, study period, study type (retrospective, single institution, multiple institutions), total number of cases in the study population, type of placenta previa, diagnosis of PAS at birth (search strategy in online supplementary data 2). Outcome measures included the need to perform a peripartum hysterectomy and direct maternal mortality. Prior surgical history was also recorded. The reference standard for differential diagnosis between minor and major placenta previa was recorded based on the placental position inside the uterine cavity on transvaginal ultrasound with relation to the internal cervical os. For the diagnosis of accreta placentation, we referred to the clinical grading based on surgical findings at delivery as previously described17 and to histopathological findings when a caesarean hysterectomy was performed, that is, placental villi directly attached to the myometrium without interposing decidua or invading the uterine wall.

Two independent reviewers (EJ and LG) undertook the quality assessment with difference agreed by consensus. The Newcastle-Ottawa scale for observational studies was used to establish the risk of bias in selection (representativeness of the exposed cohort, ascertainment of exposure and the demonstration that the outcome of interest was not present at the start of the study), comparability (evaluation of the cohorts based on the design or analysis) and outcome assessment.18 These included retrospective versus prospective studies, single versus multiple institutions studies, prenatal ultrasound description of low-lying/placenta previa and PAS, histopathological confirmation of the diagnosis of the PAS and corresponding grade of invasiveness and detailed data on management and maternal outcomes. Studies that
scored four stars for selection, two stars for comparability and three stars for ascertainment of the outcome were regarded to have a low risk of bias. Studies with two or three stars for selection, one for comparability and two for outcome ascertainment were considered to have a medium risk of bias. We deemed any study with a score of one for selection or outcome ascertainment, or zero for any of the three domains, to have a high risk of bias. No study was excluded based on the risk of bias assessment.

Analyses were conducted using STATA software (V.15; StataCorp). Standard Kurtosis analysis indicated that some values were not normally distributed and study specific estimates are therefore presented as median and IQR. A random-effects model was used to combine the studies while incorporating variations among studies unless there were three or less studies contributing to the meta-analysis in which case a fixed-effect model was used. Statistical heterogeneity was assessed with the Cochran’s Q-test and the I² statistic (the proportion of variation in study estimates because of heterogeneity rather than sampling error). Forest plots are presented to graphically summarise the study results and the pooled results. A test for heterogeneity between subgroups (ie, study types) was conducted.

Patients and public involvement

Patients and the public were not involved in the design or planning of the study.

RESULTS

The initial search provided 256 records with cross-referencing providing an additional two studies, making a total of 258 potentially relevant articles. After exclusion of duplicates and the two which were not available (figure 1), 220 remained. On screening the titles and abstracts, a further 162 were excluded as the reported outcomes were not relevant, leaving 58 studies which were obtained for full text review. An additional 38 articles were excluded after full review including letters (n=16), narrative reviews (n=10), commentaries (n=9), conference proceedings (n=2) and duplication of data in another publication (n=1), leaving 20 articles for the final analysis.

There were 13 retrospective 19 20 23 25–27 29–31 33–35 38 and 7 prospective,21 22 24 28 32 36 37 studies including a total of 1 207 296 births and 23 864 cases referred as pregnancies. There were 15 studies from a single study including 19–24 27–30 32–34 37 38 and 5 from multiple institutions25 31 or a geographical region.26 33 36 Overall, 18 studies had low or medium risk of bias (full data in online supplementary data 3).

Table 1 presents the epidemiology data of the 20 studies. These studies included 587 women with placenta previa complicated by PAS out of 6628 cases of placenta previa. The median prevalence of placenta previa in the 20 studies was 0.56% (IQR 0.39–1.24) whereas the median prevalence of placenta previa with PAS was 0.07% (IQR 0.05–0.16). The median incidence of PAS in women with a placenta previa was 11.10% (IQR 7.65–17.35).

All authors except two29 33 reported on the criteria used for the prenatal ultrasound diagnosis of placenta previa. Six studies24 26 30 32 37 39 only included major placenta previa in their cohort as defined as the placenta completely covering or partially covering the internal os of the cervix. The others included both major and minor placenta previa. The definition of minor placenta previa varied with two studies31 36 using the placental edge being <2 cm from the internal os, two studies using <3 cm22 25 32 and one study using <3 cm or <5 cm if associated with abnormal fetal presentation.21 The gestational age at confirmation of the prenatal diagnosis of placenta previa was reported in six studies22–24 28 32 37 and ranged between 20 and 34 weeks and in one study the diagnosis of placenta previa was confirmed at birth when the placenta was found to be inserted in the lower segment.19

The ultrasound diagnostic signs for PAS were reported in six studies24 28 30 32 36 37 with two studies also reporting on the use of MRI.29 38 The clinical criteria used for the diagnosis of PAS at birth were reported by nine studies19 20 23 27 28 30 33 36 37 and included a difficult delivery of the placenta without easy separation uterine wall or requiring a ‘piecemeal removal’ associated with heavy bleeding and excessive bleeding from the placental bed after placental delivery. One author described the presence of invasive villous tissue at delivery27 and one the need to suture the placental bed.23 None of the other authors reported on the gross appearance of the uterus or surgical findings at the time of caesarean delivery. In 12 studies,19 23 24 27–31 33 34 36 37 the prenatal and/or clinical diagnosis was confirmed by histopathological examination with detailed description of the microscopic criterion only reported in six.19 27 28 30 31 37 Detailed histopathological findings on the depth of villous invasiveness were reported in 9 studies24 27–29 31 33 34 36 37 out of the 20 studies (table 2). These included 283 cases of placenta previa accreta graded for 171 (60.4%) as placenta accreta (adherent), 74 (26.2%)
Table 1  Prevalence of placenta previa with placenta accreta spectrum (PAS) per pregnancies or births in the corresponding obstetric population and incidence of PAS per cohorts of placenta previa

| References              | Obstetric population | Prevalence (%) | Incidence (%) |
|-------------------------|----------------------|----------------|---------------|
| Chattopadhyay et al 19  | 41,206 births        | 26 (0.063)     | 26/222 (11.7) |
| Zaki et al 20           | 23,070 births        | 12 (0.052)     | 12/110 (10.9) |
| Ziadeh et al 21         | 18,651 births        | 13 (0.070)     | 13/65 (20.0)  |
| Ghourab 22              | 18,670 births        | 11 (0.059)     | 11/138 (8.0)  |
| Bahar et al 23          | 42,487 births        | 53 (0.125)     | 53/306 (17.3) |
| Hamada et al 24         | 2413 births          | 5 (0.207)      | 5/70 (7.1)    |
| Jang et al 25           | 35,030 births        | 53 (0.151)     | 53/560 (9.5)  |
| Rosenberg et al 26      | 185,476 births       | 23 (0.012)     | 23/779 (3.0)  |
| Kassem and Alzahrani 27 | 29,053 births        | 25 (0.085)     | 25/122 (20.5) |
| Maher et al 28          | 24,661 births        | 42 (0.170)     | 42/577 (7.3)  |
| Alchalabi et al 29      | 16,845 births        | 23 (0.137)     | 23/81 (28.4)  |
| Asicioglu et al 30      | 112,819 births       | 46 (0.401)     | 46/364 (12.6) |
| Sumigama et al 31       | 96,670 births        | 46 (0.408)     | 46/954 (4.8)  |
| Ahmed et al 32          | 3841 births          | 14 (0.365)     | 14/52 (26.9)  |
| Cheng and Lee 33        | 81,497 births        | 39 (0.048)     | 39/921 (4.2)  |
| Cho et al 34            | 11,210 pregnancies   | 39 (0.348)     | 39/442 (8.8)  |
| Kollmann et al 35       | 218,876 births       | 13 (0.006)     | 13/328 (4.0)  |
| Pillioni et al 36       | 108,000 births       | 37 (0.034)     | 37/314 (11.8) |
| Rezk and Shawky 37      | 12,654 pregnancies   | 53 (0.419)     | 53/74 (71.6)  |
| Wortman et al 38        | 14,8031 births       | 14 (0.010)     | 14/157 (8.9)  |

Table 2  Studies presenting detailed histopathological data on the depth of villous invasiveness (PAS grades)

| References              | No of cases analysed/no of cases included in the study | PAS grades |
|-------------------------|-------------------------------------------------------|------------|
|                         |                                                       | PC (%)     | PI (%)     | PP (%)     |
| Hamada et al 24         | 5/5                                                   | 3 (60.0)   | 2 (40.0)   | --         |
| Kassem and Alzahrani 27 | 19/25                                                 | 13 (68.4)  | 5 (26.3)   | 1 (5.3)    |
| Maher et al 28          | 42/42                                                 | 28 (66.6)  | 13 (31.0)  | 1 (2.4)    |
| Alchalabi et al 29      | 23/23                                                 | 15 (65.2)  | 4 (17.4)   | 4 (17.4)   |
| Sumigama et al 31       | 46/46                                                 | 14 (30.4)  | 21 (45.7)  | 11 (23.9)  |
| Cheng and Lee 33        | 39/39                                                 | 36 (92.3)  | --         | 3 (7.7)    |
| Cho et al 34            | 39/39                                                 | 24 (63.3)  | 11 (31.3)  | 4 (13.3)   |
| Pillioni et al 36       | 17/37                                                 | 7 (41.2)   | 4 (23.5)   | 6 (35.3)   |
| Rezk and Shawky 37      | 53/53                                                 | 31 (58.5)  | 14 (26.4)  | 8 (15.1)   |

Total 283/309 171 (60.4) 74 (26.2) 38 (13.4)

as placenta increta and 38 (13.4%) as placenta percreta. These studies included a total of 383,003 pregnancies or births and the prevalence for the different grades of placenta previa accreta was 0.05%, 0.02% and 0.01% for increta, increta and percreta, respectively.

The meta-analysis indicated statistically significant (p<0.001) level of overall heterogeneity between study estimates for the prevalence of placenta previa (figure 2), the prevalence of placenta previa with PAS (figure 3) and the incidence of PAS in the placenta previa cohort (figure 4). There was strong evidence of inconsistency between study types with I² values greater 85%. The difference in heterogeneity between prospective versus retrospective studies was not statistically significantly (p=0.839) different (figure 2) whereas it was significant (p=0.014) for the prevalence of placenta previa accreta (figure 3). Adjusting for type of study (prospective vs retrospective) did not reduce inconsistency between studies. The in-between placenta previa major only versus minor and major placental previa was not significant (p=0.067) for

PAS, placenta accreta spectrum
the incidence of PAS in patient with placenta previa (figure 4).

All authors but two reported on prior surgical history including cesarean section, uterine curettage, and myomectomy. Data on surgical management were available in 14 out of the 20 studies with 314 out of 441 women presenting with a placenta previa complicated by PAS. The median peripartum hysterectomy rate of 69.2% (IQR 50.0–84.0). Data on maternal mortality were available in 13 studies with 5 maternal deaths out of 387 (1.3%) cases of placenta previa with PAS.

Figure 2 Forest plots showing the heterogeneity of prevalence data in prospective and retrospective cohort studies of women presenting with a placenta previa. Only first author’s name is given for each reference. ES, effect size.

Figure 3 Forest plots showing heterogeneity in the prevalence data for prospective and retrospective cohort studies of women diagnosed with placenta previa accreta. Only first author’s name is given for each reference. ES, effect size.

Discussions
This study provides a comprehensive evaluation of the prevalence of placenta previa complicated by PAS and the incidence of PAS in women presenting with a placenta previa. Women with a prior history of cesarean delivery presenting with a low-lying/placenta previa represent more than 90% of the cases of PAS. The
Figure 4 Forest plots showing the heterogeneity in cohort studies reporting incidence data for women diagnosed with placenta previa major and PAS and those with either placenta previa minor or major and PAS. ES, effect size.; PAS, placenta accreta spectrum.

meta-analysis indicates high heterogeneity for both the prenatal diagnosis of placenta previa and for the confirmation of the diagnosis of PAS at delivery. These findings highlight the need to use international standardised clinical protocols for the screening and management of this complex obstetric condition. The current situation limits the capacity building of healthcare providers on improvements in training, implementation of guidelines and changes in clinical practice behaviour.

Defining the position of the placenta inside the uterus was one of the first aims of obstetric ultrasound examination.39 40 Following the development of real-time ultrasound imaging, placental location became an integral part of the mid-pregnancy ultrasound examination.41 Placenta previa was initially described with transabdominal scan as a placenta developing within the lower uterine segment and classified according to the relationship and/or the distance between the lower placental edge and the internal os of the uterine cervix, that is, minor placenta previa when lower edge is inside the lower uterine segment down to the internal os and major placenta previa when the placenta covers the cervix. Minor placenta previa can be further subdivided into low-lying placenta when the placental edge is less than 2 cm from the internal os and marginal placenta previa when it does. Major placenta previa can also be described as partial or complete depending on the amount of placental tissue covering the cervix. The use of transvaginal scanning has allowed for a more precise evaluation of the distance between the placental edge and the internal os42 43 but as demonstrated in our meta-analysis, the reporting of the ultrasound criteria used for the diagnosis of placenta previa has been heterogeneous.

In addition, we found also wide variation in the gestational age at diagnosis. The timing of the confirmation of the diagnosis has a direct impact on epidemiology data as up to 70% of minor placenta previa at 20–23 weeks of gestation will resolve by 32–35 weeks.44 45 An expert panel of the American Institute of Ultrasound in Medicine46 has recently recommended ceasing the use of the terms ‘partial’ and ‘marginal’ and using the term ‘placenta previa’ only when the placenta lies directly over the internal os. The placenta should be reported as ‘low-lying’ when the placental edge is less than 2 cm from the internal os and as normal when the placental edge is more than 2 cm from the internal os. The findings of our meta-analysis highlight the need for the use of such a classification in further studies.

Only 6 of the 20 studies included in the present meta-analysis provided data on the prenatal ultrasound diagnosis of PAS in patients with placenta previa. We included in the systematic review all studies published since the first ultrasound description of PAS by Tabsh et al.14 We found no studies between 1982 and 1993 (table 1), which corresponds to the time when high-resolution grey-scale ultrasound imaging became widely available. Colour Doppler imaging was introduced for the diagnosis of PAS in 1992,47 however, the sensitivity and specificity of grey-scale imaging alone in diagnosing for placenta previa accreta are high when performed by the experience operators.15 These findings indicate that the prenatal diagnosis of PAS can be performed using standard ultrasound equipment. Unlike placenta previa which is routinely screened for at the time of the fetal anomaly scan, PAS is currently not screened for and the data available on
the prenatal diagnosis of the condition come exclusively from specialist centres. In these centres, the diagnostic accuracy of ultrasound imaging is over 90%, but similar to placenta previa, the description of the ultrasound signs used for the diagnosis of PAS has also been highly variable over the last two decades. The European Working Group on Abnormally Invasive Placenta and the Abnormally Invasive Placenta international expert group have recently proposed standardised descriptions of the ultrasound signs used for the prenatal diagnosis and a protocol for the ultrasound assessment of PAS. The use of these protocols in prospective studies should also facilitate the screening of patients at high risk of PAS and in particular those with multiple prior caesarean deliveries presenting with a low-lying or placenta previa.

We found significant heterogeneity in the qualitative definition and diagnosis of PAS at birth among the nine studies that provided a description of the clinical findings. Only one of these studies described the invasive appearance of placental tissue at delivery whereas the others reported a difficult delivery of the placenta without easy separation from the uterine wall or requiring a ‘piecemeal removal’ associated with heavy bleeding as diagnostic of PAS. These clinical criteria were first described by Irving and Hertig in 1937 who did not have invasive cases in their cohort limiting their definition to abnormally adherent placenta and not to placenta increta or percreta. This definition also fails to clearly differentiate between abnormal adherence and placental retention as both present with similar clinical symptoms and aetiology leading to possible overdiagnosis of placenta previa accreta. Similarly, the finding of excessive bleeding from the placental bed after delivery of the placenta is a common complication of non-accreta placenta previa due to the implantation of the placenta in the lower uterine segment which contains less muscular fibres than the upper segment and is often thinner and dehiscent after multiple caesarean deliveries.

Detailed histopathological reports can only be obtained in those patients who have a hysterectomy or a partial myometrial resection and thus in many studies there is not histopathological confirmation of the clinical diagnosis. The main histological diagnostic criteria of accreta placentaion, that is, absence of decidua between the tip of anchoring villi and the superficial myometrium, is found with increasing incidence with advancing gestation in pregnancies with no clinical evidence of PAS. Thus, the combination of clinical criteria that do not differentiate between placenta retention and adherent accretta and the use of non-diagnostic criteria of villous invasiveness may result in the overdiagnosis of the adherent grade of PAS (table 2), in particular in those studies reporting a low rate of caesarean hysterectomy. Overall, this can explain the wide range in the prevalence (0.04%–0.42%) of placenta previa with PAS and incidence (2.9%–71.6%) of PAS in women presenting with placenta previa (figures 3 and 4).

Overall, management strategies and outcomes will vary depending on the accuracy of prenatal diagnosis, local surgical expertise and more recently access to a centre of excellence with multidisciplinary team approach. In cases of high suspicion of PAS during caesarean delivery, 60%–70% of obstetricians-gynaecologists proceed with a peripartum hysterectomy. By contrast with a conservative management approach, radical surgery is often considered to be safer, in particular in cases of invasive placentaion. The association between a placenta previa and a PAS increases the risks of both maternal morbidity and mortality. In the present study, we found that a caesarean hysterectomy was the primary management option in around 70% of the patients presenting with a placenta previa and PAS. The interstudy range was wide with four studies reporting peripartum hysterectomy rates <50%, five had rates between 50% and 99% and four had rates of 100%. This may be due to differences in study protocols, local expertise and the impact of prenatal diagnosis on management strategies but also as suggested by our analysis to differences in the rates of the different grades of PAS and the accuracy of clinical diagnosis at birth and detailed histopathological examination confirming the diagnosis.

The main limitations of this review are the quality of the published data. Thirteen out of 20 studies included in the analysis studies were retrospective and there was wide variation in the use of different ultrasound criteria for the prenatal diagnosis of placenta previa, in the clinical diagnosis of PAS at delivery and in the authors providing detailed histopathology data to confirm the clinical diagnosis. This is hampering the meta-analysis of the clinical outcomes in particular the incidence of major haemorrhage at delivery and the need and amount of blood transfusion but also the choice in management protocols and in particular the use of conservative management procedures. We would not, therefore, recommend the use of the pooled estimates beyond that of a support towards the development of standardised diagnostic protocols.

The prevalence of PAS in the general population of women giving birth varies widely. A systematic review and meta-analysis of the prevalence of placenta praevia has found evidence suggestive of regional variation. As both conditions are often associated with prior caesarean sections, it is likely that national and local caesarean delivery rates, expertise in diagnosing both conditions antenatally and access to perinatal pathologist to confirm the diagnosis of PAS at birth will influence these epidemiology data. There is a need for further prospective multicentre studies with participatory methodologies involving local service providers and facility management to accurately evaluate the consequences of high caesarean sections rates on maternal health within a particular population. Within this context, accurate epidemiological data on PAS disorders are essential in planning screening programmes and in making provision for the development of centres of excellence for the management of this increasingly common complex
obstetric condition. While the concept of core outcome measures within clinical trials is now well recognised and championed, greater efforts are required to disseminate this approach in epidemiological research to facilitate global estimation and recognition of problems emerging on a worldwide scale. Our study supports implementation, in both clinical practice and in reporting data on placenta previa accreta in the medical literature, of standardised protocols for prenatal diagnosis of both placenta previa and PAS, for the clinical diagnosis of PAS at birth and for the histopathological confirmation examination.

Contributors EJ, CB and JL-R contributed equally to the study design. EJ, LG and JL-R collected the data and carried out the qualitative analysis. CB and EJ carried out the quantitative analysis. EJ, JL-R and SLC drafted the manuscript. All authors were involved in the critical discussion and approved this final version for publication. EJ is the guarantor of the study.

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