Differential Diagnostic Value of Bilateral Inferior Petrosal Sinus Sampling (BIPSS) in ACTH-Dependent Cushing Syndrome: a Systematic Review and Meta-Analysis

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Abstract

**Background**: Previous studies have shown inconsistent results about the usefulness of bilateral inferior petrosal sinus sampling (BIPSS) in differential diagnosis of adrenocorticotropic hormone (ACTH)-dependent Cushing syndrome. This meta-analysis evaluated the diagnostic value of BIPSS via the published literature.

**Methods**: This study searched PubMed, Embase, Web of Science, Cochrane library, and Wanfang database for published data on the use of BIPSS in Cushing syndrome differential diagnosis as of October 2019. Sensitivity, specificity, positive likelihood ratio (PLR), negative likelihood ratio (NLR), diagnostic odds ratio (DOR), and receiver operating characteristic (ROC) curves were calculated based on the relevant data.

**Results**: This meta-analysis included a total of 23 studies with 1,617 patients. The calculated sensitivity, specificity, PLR, and NLR were 0.94 (95% confidence interval, CI: 0.91–0.96), 0.89 (95% CI: 0.79–0.95), 8.8 (95% CI: 4.3–17.9), and 0.07 (95% CI: 0.04–0.11), respectively. The pooled DOR and area under the ROC curve were 129 (95% CI: 48–345) and 0.97 (95% CI: 0.95–0.98), respectively.

**Conclusion**: This meta-analysis indicated that BIPSS had high diagnostic value for detecting ACTH in patients with ACTH-dependent Cushing syndrome, and BIPSS should be used as a routine method to identify ACTH-secretion sources.

**Background**

Adrenocorticotropic hormone (ACTH)-dependent Cushing syndrome (CS) is caused by excessive secretion of ACTH by the pituitary or pituitary tumors, causing bilateral adrenal hyperplasia and excessive cortisol secretion with clinical manifestations such as a moon-shaped face, buffalo hump, and hypertension. The majority of ACTH-dependent Cushing syndrome cases are caused by Cushing disease (CD), a condition in which ACTH-secreting tumors are responsible for elevated ACTH levels. Other cases, such as ectopic ACTH syndrome (EAS), have ectopic sources. These have different therapeutic principles and prognoses. Based only on clinical manifestations, detection of cortisol levels and ACTH, high- and low-dose dexamethasone suppression tests, and imaging, these conditions are not completely distinguishable. Studies have shown that non-functional pituitary tumors are common [1–3], suggesting that even if a pituitary tumor is revealed by magnetic resonance imaging (MRI), the tumor is not necessarily the source of the ACTH. Some ACTH-secreting tumors are small in size, and may not be revealed by MRI. Only 50–70% of these tumors are diagnosed [4, 5]. Therefore, negative MRI does not completely exclude ACTH-secreting tumors. In high-dose dexamethasone suppression test (HDDST), most ACTH-secreting tumors are suppressed, while most EASs are unrepressed. However, a small number of patients have unpredicted presentations on HDDST [6, 7]. An HDDST cannot effectively distinguish between ACTH-secreting tumors and EAS. Therefore, the localization rate of ACTH-secreting tumors is very low. In addition to the positive rate of MRI detection mentioned above, the HDDST has approximately 78-81% sensitivity and 67-81% specificity [8,9], while the corticotrophin-releasing hormone (CRH) stimulation test has 76-91% sensitivity and 95% specificity [10, 11]. For these reasons, more effective diagnostic approaches are needed to distinguish the two diseases.

Bilateral inferior petrosal sinus sampling (BIPSS) has been considered to be the gold standard for differential diagnosis of the above two diseases. BIPSS is an interventional method in which a blood sample from the bilateral inferior petrosal sinus and a peripheral blood sample are used to measure ACTH by calculating the lower
sinus/peripheral (IPS/P) ACTH ratio and left and right inferior petrosal sinus (IPS/IPS) ACTH ratio. The IPS/P ACTH ratio is used to distinguish between CD and EAS. In general, an IPS/P ACTH ratio of ≥ 2 before a CRH stimulation test and an IPS/P ACTH ratio of ≥ 3 after the CRH test are criteria for diagnosing CD [6]. These diagnostic criteria are also recommended by other centers [12, 13]. The ratio of ACTH between the left and right IPS is used to determine the location of pituitary microadenomas, with IPS/IPS > 1.4 indicating a tumor located at the side with higher ACTH, and IPS/IPS ≤ 1.4 indicating a tumor locating near the midline [6]. Studies have shown that vasopressin receptor is present on the surface of ACTH-secreting tumors, and administration of vasopressin stimulates the release of ACTH [14]. Application of desmopressin (DDAVP) during BIPSS enhances diagnostic accuracy [15]. Generally speaking, although BIPSS is a mildly invasive examination, it is relatively safe. It has occasional complications, including groin hematoma, cerebral hemorrhage, and vasovagal reactions (VVRs) [12,16,17]. The incidence of groin hematoma is approximately 4%, and the incidences of cerebral hemorrhage and vasovagal reactions (VVRs) are below 1%. Occasional pulmonary embolism is also reported by some researchers. However, meta-analysis of BIPSS is currently unavailable. This study performed a meta-analysis of BIPSS for the differential diagnosis of ACTH-dependent Cushing syndrome and evaluated the differential diagnostic value for this condition.

**Methods**

**Data Sources, Search Strategy, and Selection Criteria**

This study strictly followed the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines [18] and used PubMed, Embase, Web of Science, Cochrane Library, and Wanfang databases to search for studies using BIPSS for the differential diagnosis of ACTH-dependent Cushing syndrome as of October 2019. The following search terms were used: petrosal sinus sampling, bilateral inferior petrosal sinus sampling, Cushing's syndrome, Cushing disease, and ectopic Cushing syndrome. The search strategies in the various databases were as follows: PubMed: ("petrosal sinus sampling" [Mesh]) AND "Cushing's syndrome" [Mesh]); Embase: (Emtree term-expanded = Cushing's syndrome AND Abstract=petrosal sinus sampling); Web of Science: TS = (petrosal sinus sampling AND Cushing’s syndrome); and Cochrane Library and WanFang: keyword = (petrosal sinus sampling AND Cushing’s syndrome). During searching, keywords and free words were used simultaneously. Manual searches were also used, and relevant references included in the extracted papers were also searched.

Literature was searched by two of the authors (Hao Wang, Run Ce-Cai) independently. If there was a disagreement for inclusion or exclusion, another author (Ying Ba) became involved in the resolution following discussion. The inclusion criteria of this meta-analysis were as follows: (1) patients confirmed with Cushing syndrome (CS) and unclear ACTH source; (2) CS caused by ACTH-secreting tumor or EAS confirmed by postoperative pathology or by clinical manifestations, biochemical tests, and surgery; (3) the study provided true positives (TP), false positives (FP), false negatives (FN), and true negatives (TN) or the data for the calculation of TP, FP, FN, and TN. The exclusion criteria were: (1) studies with incomplete data or data which could not be used to calculate the contingency table, (2) non-original studies, (3) repeated studies, (4) animal studies, and (5) studies with less than 20 patients included.

**Data Collection and Quality Assessment**
Two authors (Qian Xing, Ying Ba) read the included papers and extracted relevant data through discussion. In case of disagreement, another author (Hao Wang) was involved in further discussion. Contents of data extraction in the literature included: name of the first author, year of publication, country of the study, study design (prospective and retrospective), the application of CRH or DDAVP stimulation, the application of prolactin (PRL) correction, TP, FP, FN, and TN. The quality of the included studies was evaluated by two of the authors independently using the Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2) [19,20] according to the four aspects as follows: selection of cases, trials to be assessed, gold standard, and flowchart and progress of cases. Each of the assessments contained seven items which were answered as “yes,” “no,” or “uncertain.” An answer of “yes” indicated that the risk offset of the study was low, while the answers of “no” and “uncertainty” indicated high risk offset.

**Statistical Analysis**

This study used Revman 5.3 for quality evaluation and Stata 14.0 statistical software for data analysis. The TP, FP, FN, TN, sensitivity, specificity, positive likelihood ratio (PLR), negative likelihood ratio (NLR), and corresponding 95% confidence intervals (CIs) of each included study were extracted, otherwise, we calculated the integrated sensitivity, specificity, PLR, and NLR using the bivariate random effects model [21]. The receiver operating characteristic (SROC) curve and the area under the ROC curve were calculated using a hierarchical regression model [22]. The Q statistic and I-square were used for heterogeneity tests. P > 0.10 indicated no significant heterogeneity, while P < 0.10 indicated significant heterogeneity for the Q statistic [23, 24]. The fixed effect model was used when the heterogeneity was low (P > 0.05, I-square < 50%), while the random effect model was used when the heterogeneity was high (P < 0.05, I-square > 50%). A meta-regression analysis of the diagnostic odds ratio (DOR) was performed according to the study design, year of publication, country of publication, application of CRH or DDAVP, application of PRL correction, and the number of patients included in the study[25]. Deek’s asymmetry test was used to evaluate whether a publication bias existed[26]. All reported P values were two-sided, and P < 0.05 was considered statistically significant for pooled diagnostic parameters.

**Results**

As shown in Figure 1 which describes the literature searches and the workflow for study inclusion, there were 822 articles in the initial search, but 256 of them were found to be duplicated and were removed from further analysis. In addition, a total of 472 articles included irrelevant research articles, reviews, commentaries, editorials, and letters, which were further removed. Of the remaining 94 articles, those that contained incomplete data, replicated research, no gold standard, incomplete research descriptions, or less than 20 patients were also removed. Thus, a total of 23 studies were included in this meta-analysis [3,6, 12,13, 27–45].

Table 1 shows the characteristics of the included studies which were published in 1991–2019, including 11 studies conducted in Europe, nine studies conducted in the United States or Brazil, and 3 studies conducted in China or India. There were 3 prospective studies and 20 retrospective studies included in this meta-analysis. Figure 2 shows the quality of the included studies.

The summary results for sensitivity and specificity are presented in Figure 3. The pooled sensitivity was 0.94 (95% CI: 0.91–0.96), the specificity was 0.89 (95% CI: 0.79–0.95), the PLR was 8.8 (95% CI: 4.3–17.9), and the NLR was 0.07 (95% CI: 0.04–0.11). The DOR of further integration of BIPSS on ACTH–dependent Cushing syndrome was 129 (95% CI: 48–345; Figure 4). Lastly, the summary area under the ROC curve was 0.97 (95% CI: 0.95–0.98;
Figure 5), and the results of the DOR forest map for heterogeneity testing were $P = 0.00$, I-square= 99.35. A meta-regression analysis was performed based on the study design (prospective or retrospective), year of publication, country of publication, sample size (number of patients enrolled being 21-100, 100-200, >200), patient ethnicity, application of CRH or DDAVP, and application of PRL correction (Figure 6). The results suggested that the research design was the main cause of heterogeneity. Deek's asymmetry test was used to detect the presence of publication bias, and the results indicated a publication bias ($P = 0.01$; Figure 7).

Table 1. Characteristics of the included studies
| Author          | Year | Country        | Design | Stimulation | PRL adjust | Gold Standard | TP  | FP  | FN  | TN  |
|-----------------|------|----------------|--------|-------------|------------|----------------|-----|-----|-----|-----|
| Oldfield EH^6   | 1991 | USA            | pro    | CRH         | No         | Pathology      | 203 | 0   | 0   | 17  |
| Findling JW^27  | 1991 | USA            | pro    | CRH         | No         | Pathology      | 18  | 3   | 2   | 6   |
| Kaltsas GA^28   | 1999 | UK             | retro  | CRH         | No         | Pathology      | 50  | 0   | 19  | 6   |
| Invitti C^29    | 1999 | Italy          | retro  | DDAVP       | No         | Pathology      | 65  | 0   | 11  | 9   |
| Bonelli FS^13   | 2000 | USA            | retro  | CRH         | No         | Pathology      | 71  | 1   | 6   | 9   |
| Wiggam MJ^30    | 2000 | Northen Ireland | retro  | CRH         | No         | Pathology      | 36  | 0   | 8   | 1   |
| Colao A^12      | 2001 | Italy          | retro  | CRH         | No         | Pathology      | 60  | 0   | 8   | 10  |
| Lefournier V^31 | 2003 | France         | retro  | CRH         | No         | Pathology      | 65  | 2   | 4   | 6   |
| Swearingen B^32 | 2004 | USA            | retro  | CRH         | Yes        | Pathology      | 70  | 2   | 9   | 2   |
| Liu C^33        | 2004 | USA            | retro  | CRH         | No         | Pathology      | 39  | 0   | 3   | 9   |
| Kaskarelis LS^3 | 2006 | Greece         | retro  | CRH         | No         | Pathology      | 40  | 3   | 6   | 5   |
| Machado MC^34   | 2006 | Brazil         | retro  | CRH         | No         | Pathology      | 46  | 0   | 1   | 5   |
| Castinetti F^35 | 2007 | France         | retro  | DDAVP       | Yes        | Pathology      | 32  | 0   | 4   | 7   |
| Tsagarakis S^36 | 2007 | Greece         | retro  | CRH         | No         | Pathology      | 46  | 0   | 1   | 7   |
| Shi XH^37       | 2011 | China          | retro  | No          | No         | Pathology      | 58  | 1   | 10  | 4   |
| Mulligan GB^38  | 2011 | USA            | retro  | CRH         | No         | Pathology      | 33  | 1   | 2   | 1   |
| Anderegenge L^39| 2011 | Switzerland    | retro  | CRH         | No         | Pathology      | 19  | 1   | 1   | 2   |
| Sharma ST^40    | 2011 | USA            | retro  | No          | No         | Pathology      | 16  | 1   | 1   | 7   |
| Shetch SA^41    | 2012 | USA            | retro  | CRH         | Yes        | Pathology      | 195 | 5   | 12  | 5   |
| Grant P^42      | 2012 | UK             | retro  | DDAVP       | No         | Pathology      | 72  | 1   | 0   | 10  |
Discussion

This study was the first meta-analysis to evaluate the differential diagnostic value of BIPSS in ACTH-dependent Cushing syndrome. It included a total of 23 studies and 1,617 patients. Our results suggested that the sensitivity and specificity of BIPSS to pituitary or ectopic ACTH were 94% and 89%, respectively, indicating that BIPSS has high value in the differential diagnosis of ACTH-dependent Cushing syndrome. In addition, the DOR value was also high, suggesting that BIPSS could effectively identify the ACTH source. The area under the SROC curve was 0.97, suggesting that the overall diagnostic performance of BIPSS was effective.

BIPSS has a high value in the differential diagnosis of CS cases that have typical CS presentations clinically and biochemically but have an unclear ACTH source. Because BIPSS does not identify the ACTH source from a morphological perspective, but from a functional perspective, this diagnostic approach is accurate, with relatively high sensitivity and specificity. CD accounts for a large proportion of ACTH-dependent Cushing syndrome cases, and BIPSS is particularly suitable for patients with negative MRI results. Furthermore, BIPSS provides an important basis for guiding the surgical treatment of this disease.

In most cases, the ACTH level of CD was lower than EAS. For example, the ACTH level of the ACTH-secreting tumors was 111.35 pg/ml, while the ACTH level of EAS was 277.01 pg/ml [37]. After CRH or DDAVP stimulation, the ACTH level increased significantly. Many researchers believe that the stimulation intensity of DDAVP on ACTH-secreting tumors is weaker than that of CRH. For example, in Jarial’s study, the ACTH (IPS/P) ratio of ACTH-secreting tumors is increased 11.6-fold after DDAVP stimulation. After CRH stimulation, the ratio is increased by 28-fold [44]. In terms of the maximum ACTH level after stimulation, Bonelli’s study showed that the ACTH levels reached 1,062 pg/ml after DDAVP stimulation, and 3,058 pg/ml after CRH stimulation [13]. This leads naturally to the question of why the stimulation intensity of DDAVP is weaker than that of CRH. We believe that CRH directly stimulates ACTH, and stimulation of DDAVP is due to the presence of vasopressin receptor. Thus, the stimulation intensity of DDAVP is weaker than that of CRH.

BIPSS has a high differential diagnostic value for CD and EAS. Application of CRH or DDAVP stimulation enhances the sensitivity and specificity of BIPSS. However, BIPSS should still be combined with other diagnostic methods, such as imaging, HDDST, and the low-dose dexamethasone suppression test for comprehensive diagnosis.

False negative results can occur in BIPSS. These have been reported to be approximately 10% [32], and may be related to operational failure or abnormal venous drainage from the inferior petrosal sinus. BIPSS is not ideal for identifying the diseased side [31, 46], which may be due to the presence of branches joined to the cavernous sinus and frequent contralateral venous return. A previous study used cavernous sinus sampling instead of BIPSS to obtain a good differential diagnosis for CD and EAS [47]. For BIPSS, the success rate is closely related to the
operator's technique and experience, and accurate catheterization is very important. Results of a previous study suggest that PRL for correction improves the success rate of catheterization [48].

An interesting consideration is whether the false positive rate of BIPSS increases among the patients with positive MRI results, which is only discussed in few studies. The study by Kaskarelis et al. showed that 1 out of 23 MRI-positive patients had a BIPSS-false-negative result (4.3%) and 2 out of 55 MRI-negative patients had BIPSS-false-negative results (3.6%) [3], while the majority of BIPSS-false-positive rates in CS patients ranged from 0 to 5%. Thus, the BIPSS-false-positive rate of the MRI-positive patients in Kaskarelis et al.’s study was higher than that of the MRI-negative patients, and also higher than the average of most other studies. Maybe after being stimulated, the MRI positive patients still secreted a little more ACTH. Since few studies were related to this issue, further studies with increased sample sizes are needed for verification.

This meta-analysis provides implications for future studies as follows: PRL can be used as a reference to improve the accuracy of catheterization during BIPSS. CRH or DDAVP stimulation should also be used during BIPSS to improve the sensitivity and specificity.

The strengths of this study were that we followed a standard protocol and used a comprehensive search strategy. Furthermore, the bivariate random effects model and hierarchical summary ROC analyses were used. Finally, meta-regression analysis suggested that the source of heterogeneity was mainly in the experimental design.

However, our meta-analysis also had some limitations. First, some details of patient characteristics were not available, which might affect our appraisal of the diagnostic value of BIPSS. Second, the analysis used summarized data, which restricted us from conducting more detailed analysis. Finally, the publication bias of this meta-analysis was P < 0.05, suggesting the presence of publication bias. The possible reasons for this were that (1) BIPSS had high diagnostic accuracy of TP and TN for determining the ACTH source and likely shows the ideal statistical results in the software, leading to the calculation of publication bias; (2) authors might have submitted studies only with positive results to increase the chance of being published; and (3) this meta-analysis only included studies published in Chinese and English.

This study was the first meta-analysis to evaluate BIPSS’s effects on determining the etiology of ACTH-dependent Cushing syndrome, suggesting that BIPSS had a great differential diagnostic value for the ACTH source. Results of this study require further large-scale prospective studies to validate the differential diagnostic value of BIPSS for ACTH-secretion sources in different patients.

**Conclusion:**

This meta-analysis indicated that BIPSS had a high diagnostic value for patients with ACTH-dependent Cushing syndrome, and as such, BIPSS should be used as a routine method to identify ACTH-secretion sources. CRH or DDAVP stimulation should be used during BIPSS to improve the test’s sensitivity and specificity.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.
Consent for publication

Not applicable

Availability of data and materials

Not applicable. This study is a systematic review and we used primary data, which are already publicly available.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

HW,YB conceived and designed the study and approved the final draft of the manuscript submitted for review and publication; YB, QX and RCC searched databases, data extracted and study selection. HW performed data analysis. HW,YB,QX,RCC wrote the manuscript. All authors read and approved the final manuscript.

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Deek’s plot for BIPSS in the differential diagnosis of ACTH-dependent CS

Supplementary Files

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