The diagnostic model for early detection of gestational diabetes mellitus and gestational diabetic nephropathy

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Funding information
Chongqing Clinical Key Specialty Construction Project

Abstract
Background: Gestational diabetes mellitus (GDM) and gestational diabetic nephropathy (GDN) have become an increasingly serious problem worldwide, which can cause a large number of adverse pregnancy consequences for mothers and infants. However, the diagnosis of GDM and GDN remains a challenge due to the lack of optimal biomarkers, and the examination has high requirements for patient compliance. We aimed to establish a simple early diagnostic model for GDM and GDN.

Methods: We recruited 50 healthy pregnant (HP), 99 GDM patients, 99 GDN patients at Daping Hospital. Renal function indicators and blood cell indicators were collected for all patients.

Results: Compared with HP, GDM, and GDN patients exhibited significantly higher urea/creatinine ratio and NEU. The diagnostic model1 based on the combination of urea/creatinine ratio and NEU was built using logistic regression. Based on receiver operating characteristic curve analysis, the area under the curve (AUC) of the diagnostic model was 0.77 (0.7, 0.84) in distinguishing GDM from HP, and the AUC of the diagnostic model was 0.94 (0.9, 0.97) in distinguishing GDN from HP. Meanwhile, the diagnostic model2 based on the combination of β2-mG, PLT, and NEU in GDM and GDN patients was built using logistic regression, and the area under the ROC curve (AUC ROC) was 0.79 (0.73, 0.85), which was larger than the individual biomarker AUC.

Conclusion: Our study demonstrated that the diagnostic model established by the combination of renal function indicators and blood cell indicators could facilitate the differential diagnosis of GDM and GDN patients.

KEYWORDS biomarker, blood cell indicators, diagnostic model, gestational diabetes mellitus, gestational diabetic nephropathy, renal function indicators
1 | INTRODUCTION

Gestational diabetes mellitus (GDM) is an increasingly serious health problem worldwide.\(^1\) It is one of the most common complications of pregnancy, with the implementation of China’s new fertility policy the incidence rate of GDM is increasing. In 2019, an analysis showed that the incidence rate of GDM in China was up to 12.8%–16.7%.\(^2\) GDM frequently causes fetal disorders during pregnancy and causes fetal mortality and morbidity.\(^3\) Gestational diabetic nephropathy (GDN) is the progression of GDM and there is kidney damage at the same time.\(^4\) It was reported that 20% to 40% of diabetic patients in the United States suffer from varying degrees of kidney injury, and diabetic nephropathy is the first cause leading to end-stage renal disease (ESRD).\(^5\) At the same time pregnancy in women with diabetic nephropathy had a higher risk of fetal/neonatal severe complications.\(^6\)

Early detection of GDM and GDN, timely control of blood glucose, and intervention of kidney injury are of great significance to reduce the risk of maternal and infant. At present, the 75 g oral glucose tolerance test (OGTT) is the diagnostic standard proposed by the international diabetes and pregnancy research group (IADPSG). However, the examination has higher requirements for patient compliance.\(^7\) It is also more troublesome to check with requiring multiple blood. In addition, OGTT is carried out between 24–28 weeks of gestation, which makes it challenging to detect GDM at an early stage.\(^8\) Meanwhile, there are many clinical indicators of kidney injury, such as urinary micro-albumin (ALB), \(\beta_2\)-microglobulin (\(\beta_2\)-mG), cystatin C (Cys-C), urea, Crea et al.,\(^9\) in many cases with normal renal injury indicators, kidney damage has occurred.\(^10,11\)

In recent years, in the study of the relationship between inflammatory factors and microcirculation, it is found that inflammatory response plays an important role in the occurrence and development of GDM and GDN.\(^12\) More and more studies have shown that the increase of leukocyte, PLT is associated with GDM and GDN.\(^13-16\) Neutrophils, lymphocytes, and platelet are important observation indexes in blood cells classification. Neutrophil to lymphocyte ratio (NLR) and platelet to lymphocyte ratio (PLR) are new inflammatory indexes. Increased NLR has been reported in inflammatory bowel disease, diabetes mellitus, thyroiditis, and SARS Cov2 infection.\(^17-19\) Similarly, PLR is associated with irritable bowel disease, type 2 diabetes mellitus, and COVID-19 infection.\(^20-22\) NLR and PLR are used to diagnose many diseases.\(^23,24\) However, the combination of renal function and hemocyte indicators for the diagnosis of GDM and GDN is rarely reported.

Early diagnosis of GDM and GDN can improve the survival and health of mothers and infants.\(^25,26\) Therefore, the present study aims to combine renal function index and hemocyte indicators to establish an early diagnostic model of GDM and GDN patients.

2 | METHODS

2.1 | Study design and participants selection

This was a case–control study. This study included patients between 18 and 45 years of age admitted to the obstetric outpatient of Daping Hospital located in Chongqing, China, from January 2019 to January 2021. Ethics approval was obtained from the Ethics Committee of Daping Hospital, Army Medical University. The patient data were collected from clinical tests within 24–28 gestational weeks. In this study, patients were classified into three cohorts, healthy pregnant (HP), patients with GDM, and patients with GDN by three obstetricians based on a retrospective analysis of the patient’s clinical symptoms and examination results. The HP was from healthy volunteers and matched with patients in age and gestational weeks.

2.2 | Inclusion and exclusion criteria

2.2.1 | Inclusion criteria

Inclusion criteria of GDM refer to the international diabetes and pregnancy research group (IADPSG) on the diagnostic criteria and classification strategy for gestational hyperglycemia. All pregnant women were given an oral glucose tolerance test at 75 g, fasting blood glucose 5.1 mmol/L, 1 h blood glucose 10.0 mmol/L, 2 h blood glucose 8.5 mmol/L at 24–28 weeks of gestation. One of the above abnormalities is diagnosed as GDM and included in the GDM group.\(^27\) The GDN group was included the patients who were diagnosed as GDN after 28 gestation weeks according to the consensus on prevention and treatment of diabetes nephropathy (2014 edition). One of the following can be used to diagnose diabetes nephropathy: heavy albuminuria (>0.3 g/24h); diabetes retinopathy with any stage of chronic kidney disease.\(^28,29\)

2.2.2 | Exclusion criteria

Patients who had previous diabetes, acute complications of diabetes, chronic inflammation, obstetric complications, acute cardiovascular and cerebrovascular diseases, autoimmune diseases, malignant tumors, hematological diseases, history of infection, trauma and other stressful conditions, acute and chronic glomerulonephritis, acute and chronic renal failure, and other primary kidney diseases, and so on were excluded.

2.3 | Data collection

Data were collected between 24 and 28 weeks of gestation, including renal function indicators (urea, creatinine (Crea), urea/creatinine
ratio, Cystatin C (Cys-C), β2-microglobulin (β2-mG), estimated glomerular filtration rate (eGFR) and blood cell indicators (platelets count (PLT), lymphocyte count (LYM), neutropenia count (NEU), neutrophil to lymphocyte ratio (NLR), platelet to lymphocyte ratio (PLR)). Renal function indicators were quantified with a UniCel DxC800 Synchron device (Beckman, USA). The blood cell indicators were measured using the Sysmex XE-2100 automatic hematology analyzer (Sysmex, Japan).

**TABLE 1** Demographic and biological data of the population

| Variables                  | Total (n = 247) | HP (n = 50) | GDM (n = 99) | GDN (n = 98) | p value |
|----------------------------|----------------|-------------|--------------|--------------|---------|
| Age (year)                 | 30.69 ± 4.71   | 30.90 ± 4.97| 30.62 ± 4.50 | NS           | NS      |
| Gestational weeks          | 33.5 (30.18, 39.07) | 33 (30.29, 38.86) | 33.57 (30.25, 38.18) | NS | NS      |
| Urea (mmol/L)              | 2.6 (2.14, 3)  | 3.01 (2.42, 3.58) | 2.94 (2.4, 3.78) | 0.003 | 0.006  |
| Crea (μmol/L)              | 42.2 (38.58, 46.95) | 41.6 (37.8, 47.1) | 43.55 (38.95, 47.55) | NS | NS      |
| Urea/Creatinine ratio      | 15.65 (12.62, 17.23) | 17.83 (13.71, 21.67) | 16.9 (13.75, 20.98) | 0.003 | 0.021  |
| Cys-C (mg/L)               | 1.12 (0.88, 1.33) | 1 (0.85, 1.21) | 1.05 (0.87, 1.23) | NS | NS      |
| β2-mG (mg/L)               | 1.46 (1.24, 1.73) | 1.42 (1.19, 1.7) | 1.51 (1.31, 1.82) | 1 | 0.467  |
| eGFR (ml/min/1.73 m3)      | 204.37 (175.72, 224.5) | 204.73 (177.22, 226.13) | 190.59 (171.37, 219.78) | NS | NS      |
| PLT (10^9/L)               | 179.82 ± 48.42 | 177.94 ± 45.07 | 194.34 ± 52.96 | 0.815 | 0.107  |
| LYM (10^9/L)               | 1.39 (1.21, 1.68) | 1.49 (1.27, 1.74) | 1.57 (1.3, 1.93) | 0.502 | 0.029  |
| NEU (10^9/L)               | 5.48 (4.7, 5.78) | 6.12 (5.22, 6.78) | 7.46 (6.63, 8.72) | 0.001 | 0      |
| NLR                        | 3.7 (2.96, 4.27) | 4.05 (3.3, 5) | 4.7 (3.87, 6.17) | 0.184 | 0      |
| PLR                        | 117 (98.91, 154.36) | 117.56 (95.9137.89) | 114.51 (91.98, 150.86) | NS | NS      |

**FIGURE 1** The result of biomarkers in HP, GDM, and GDN. (A) Violin plot showing the results of biomarkers in HP (n = 50), GDM (n = 99), and GDN (n = 98). Horizontal lines indicate the median and interquartile range. *p < 0.05, **p < 0.01, ***p < 0.001, ns, no significance (Mann–Whitney U test)
2.4 | Statistical analysis

Normally distributed variables are given as means±SD, whereas non-normally distributed variables were given as median (interquartile range, IQR). The comparison between continuous variables was performed using a t-test if the continuous value is normal distribution and homogeneity of variance or Mann–Whitney U test if not. For the establishment of the diagnostic model, indicators with a statistical difference were selected and taken as candidates in multivariable logistic regression models. Then, the regression equation (diagnostic model) was obtained. The receiver operating characteristic (ROC) curve was plotted to evaluate the diagnostic performance of various indicators. The area under the ROC curve (AUC) was calculated, and the cutoff values with the largest Youden index (the sum of sensitivity and specificity) were calculated from the ROC curve. The area under the curve (AUC), sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), positive likelihood ratio (PLR), negative likelihood ratio (NLR), and accuracy, as well as the corresponding 95% confidence interval (CI) were calculated. Statistical was analyzed with SPSS 25.0 statistical software, R v.4.1.1 Statistical Software, and GraphPad Prism9.0. In all statistical analyses, a p value <0.05 was considered statistically significant.

3 | RESULTS

3.1 | Participant characteristics

A total of 247 subjects were included in this study. Fifty HP, 99 GDM, and 98 GDN were consecutively enrolled from January 2019 to January 2021 at Daping Hospital. The demographic and biological data of all participants were summarized in (Table 1). There was no significant difference in scale of age and gestational weeks among these three groups. The mean age was around 30 years, and the median gestational weeks is 33 weeks.

3.2 | Biomarkers in HP, GDM, and GDN

We performed biomarkers analysis among HP, GDM, and GDN. It was observed that compared with HP, GDM patients showed significantly higher urea/creatinine ratio, urea, NEU; GDM patients showed significantly higher urea/creatinine ratio, urea, NEU, NLR, LYM; compared with GDM, GDN patients showed significantly higher β2-Mg, PLT, NLR, NEU. And the NEU showed a clear gradual upward trend as GDN > GDM > HP. No significant difference in Crea, Cys-C, eGFR, PLR, among these three groups (Figure 1).

3.3 | Establishing diagnostic Model of GDM and GDN

To investigate the possibility of combining different biomarkers to distinguish the status of GDM and GDN, we performed a Mann–Whitney U test and discovered the potential of the combination of these indexes to distinguish GDM and GDN from HP (Table 1).
Table 3: The performance of number of biomarkers for distinguishing between HP and GDM, HP and GDN

| Number of biomarkers | Sensitivity (95% CI) | Specificity (95% CI) | PPV (95% CI) | NPV (95% CI) | PLR (95% CI) | NLR (95% CI) | Accuracy |
|---------------------|----------------------|----------------------|--------------|--------------|--------------|--------------|----------|
| **GDM**             |                      |                      |              |              |              |              |          |
| ≥1                  | 74.75% (64.84%, 82.7%) | 80% (65.86%, 89.5%) | 88.1% (78.75%, 93.83%) | 61.54% (48.62%, 73.09%) | 3.74 (2.12, 6.58) | 0.32 (0.22, 0.45) | 76.51%   |
| 2                   | 23.23% (15.58%, 33%)  | 100% (91.11%, 100%) | 100% (82.19%, 100%) | 39.68% (31.2%, 48.81%) | –             | 0.77 (0.69, 0.86) | 48.99%   |
| GDN                 |                      |                      |              |              |              |              |          |
| ≥1                  | 97.96% (92.11%, 99.65%) | 80% (65.86%, 89.5%) | 90.57% (82.93%, 95.13%) | 95.24% (82.58%, 99.17%) | 4.9 (2.81, 8.53) | 0.03 (0.01, 0.1) | 91.89%   |
| 2                   | 25.51% (17.48%, 35.49%) | 100% (91.11%, 100%) | 100% (83.42%, 100%) | 40.65% (32%, 49.89%) | –             | 0.74 (0.66, 0.84) | 50.68%   |

Table 4: Diagnostic performance of additive combination of the two markers between GDM and GDN

| Number of biomarkers | n  | GDN       |
|---------------------|----|-----------|
| 0                   | 21 | 2 (9.52%) |
| 1                   | 56 | 11 (19.64%) |
| 2                   | 80 | 47 (58.75%) |
| 3                   | 47 | 38 (80.85%) |

To establish the diagnostic model based on a combination for differentiating GDM and GDN from HP, all variables with statistical significance were used for multivariable logistic regression model (Table S1). The urea/creatinine ratio and NEU are included in model1. The diagnostic model1 was established as follows: model1: \( P = 1 / [1 + e^{-(−7.636 + 0.171 \times \text{urea/creatinine ratio} + 0.883 \times \text{NEU})}] \). \( P \), predictive value; e, natural logarithm.

3.4 | Biomarker combinations for diagnosing GDM, GDN

Using the best cutoffs derived from the ROC analysis for each of the two biomarkers, we assigned each elevated biomarker a score of 1 (no elevated biomarker scored 0). The number of elevated biomarkers was summed to give a cumulative total (range 0–2). With this approach, an increase in the number of elevated biomarkers was associated with a progressive increase in the risk of GDM and GDN (Table 2). When the score ≥1, the sensitivity of diagnosing GDM is 74.75% (64.84%, 82.7%), and specificity is 80% (65.86%, 89.5%); the sensitivity of diagnosing GDN is 97.96% (92.11%, 99.65%), and specificity is 80% (65.86%, 89.5%); when the score = 2, the sensitivity of diagnosing GDM is 23.23% (15.58%, 33%), and specificity is 100% (91.11%, 100%); the sensitivity of diagnosing GDN is 25.51% (17.48%, 35.49%), and specificity is 100% (91.11%, 100%) (Table 3). When the combined markers were used (model1), the area under the ROC curve (AUC ROC) was 0.77 (0.7, 0.84) and 0.94 (0.9, 0.97); these were larger than individual biomarker AUC (Table S2, Figure 2).

3.5 | Establishing diagnostic Model of GDN

As with the above method, we performed a Mann–Whitney U test and discovered the potential of the combination of these indexes to distinguish GDN from GDM (Table 1). To establish the diagnostic model based on a combination for differentiating GDN from GDM, all variables with statistical significance were used for multivariable logistic regression (Table S3). The j2-mG, PLT, and NEU are included in model2. The diagnostic model2 was established as follows: model2: \( P = 1 / [1 + e^{-(−7.636 + 1.216 \times j2\text{-mG} + 0.009 \times \text{PLT} + 0.605 \times \text{NEU})}] \). \( P \), predictive value; e, natural logarithm.

3.6 | Biomarker combinations for diagnosing GDN

Same as above, using the best cutoffs derived from the ROC analysis for each of the three biomarkers, and assigned each elevated biomarker a score of 1 (no elevated biomarker scored 0). The number of elevated biomarkers was summed to give a cumulative total (range 0–3). With this approach, an increase in the number of elevated biomarkers was associated with a progressive increase in the risk of GDN (Table 4). When the score ≥1, the sensitivity of diagnosing GDN is 97.96% (92.11%, 99.65%), and specificity is 12.12% (6.69%, 20.59%); when the score ≥2, the sensitivity of diagnosing GDN is 86.73% (78.03%, 92.47%), and specificity is 57.58% (47.24%, 67.32%); when the score ≥3, the sensitivity of diagnosing GDN is 38.78% (29.26%, 49.18%), and specificity is 90.91% (83.01%, 95.5%) (Table 5). With the increase of scores, sensitivity decreased, and specificity increased.
When the combined markers were used (model2), the area under the ROC curve (AUC ROC) was 0.79 (0.73, 0.85), this was larger than the individual biomarker AUC (Table S4, Figure 2).

4 | DISCUSSION

In this study, we constructed an early diagnostic model of GDM and GDN by combining renal function indicators with blood cell indicators. The diagnostic model 1 is used to distinguish GDM/GDN patients from HC with higher AUC (0.77 (0.7, 0.84)), (0.94 (0.9, 0.97)) compared with the individual immune indicator. The diagnostic model 2 is used to distinguish GDN with higher AUC (0.74 (0.63, 0.84)) compared with the individual immune indicator.

Hyperglycemia during pregnancy is a severe complication in women during pregnancy. At the same time, hyperglycemia during pregnancy is also the main cause of other complications during pregnancy.\(^1\) When GDM is complicated with renal impairment and developed into GDN, the maternal-fetal risks will be further increased. So early diagnosis of GDM and GDN is very important. At present, the diagnosis of GDM relies on the OGTT test with poor compliance. In the meantime, the diagnosis of GDN relies on renal substantial damage and renal function indicators, but when these indicators appear warning, kidney damage has occurred for a period of time. At present, there are no clear and unified indicators for early diagnosis. It has been reported in the literature that neutrophil gelatinase-associated lipocalin (NGAL) and kidney injury molecule-1 (KIM-1) expression are associated with early renal injury; however, the clinical implementation rate of these projects is not high.\(^2\) Combined with clinical work and research paper results, we established an early diagnosis model that combined renal function indicators and blood cell indicators, showing higher sensitivity and specificity compared with single indicators.

An interesting finding is that when urea/creatinine ratio and NEU are combined in the diagnosis of GDM, it is found that when at least one of the two indicators is early warning, the diagnostic accuracy is (76.51%) higher model 1 (66.44%). When urea/creatinine ratio and NEU are combined in the diagnosis of GDN, it is found that when at least one of the two indicators is early warning, the diagnostic accuracy is (91.89%) higher model 1 (86.49%). This discovery shows the special use of joint diagnosis, and it is feasible to diagnose diseases according to the number of marker warnings.

This study is a single-center study, and the sample size is not large enough. To obtain more stable results, multicenter cooperation should be carried out, and more cases are needed.

In conclusion, our findings suggest that the diagnostic model based on the combination of renal function indicators and blood cell indicators may be an adjunctive but useful method in the diagnosis of GDM and GDN.

AUTHOR CONTRIBUTIONS

Huimin Chong and Jingmi Li analyzed the data and wrote the draft; Kejun Zhang and Dan Liao conceived and designed the study, analyzed the data, and finalized the draft; Caigui Chen and Wan Wang collected the samples, analyzed the data, and participated in the drafting. All authors read and approved the final manuscript.

FUNDING INFORMATION

This study was supported by Chongqing Clinical Key Specialty Construction Project, Chongqing, China.

CONFLICT OF INTEREST

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available in the supplementary material of this article.

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**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Chong H, Li J, Chen C, Wang W, Liao D, Zhang K. The diagnostic model for early detection of gestational diabetes mellitus and gestational diabetic nephropathy. *J Clin Lab Anal*. 2022:36:e24627. doi: 10.1002/jcla.24627