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Group Penalized Logistic Regressions Predict Ovarian Cancer *

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Abstract

Purpose: Ovarian cancer ranks first among gynecological cancers in terms of the mortality rate. Accurately diagnosing ovarian benign tumors and malignant tumors is of immense important. The goal of this paper is to combine group LASSO/SCAD/MCP penalized logistic regression with machine learning procedure to further improve the prediction accuracy to ovarian tumors prediction problem.

Methods: We combine group LASSO/SCAD/MCP penalty with logistic regression, and propose group LASSO/SCAD/MCP penalized logistic regression to predict ovarian cancer. Firstly, we select 349 ovarian cancer patients and divide them into two sets: one is the training set for learning, and the other is the testing set for checking, and choose 46 explanatory variables to divide into 11 different groups. Secondly, we apply the training set and group coordinate descent algorithm to obtain group LASSO/SCAD/MCP estimator, and apply the testing set to compute confusion matrix, accuracy, sensitivity and specificity. Finally, we compare the prediction performances for the five classifiers: group LASSO/SCAD/MCP penalized logistic regression, artificial neural network (ANN) and support vector machine (SVM).

Results: Group LASSO/SCAD/MCP penalized logistic regression finally selects 6/4/1 different groups. The prediction accuracy and AUC for group MCP/SCAD/LASSO penalized logistic regression, SVM and ANN is 93.33%/85.71%/82.26%/74.29%/72.38% and 0.892/0.852/0.823/0.639/0.789, respectively.

Conclusions: Group MCP/SCAD/LASSO penalized logistic regression performs better than SVM and ANN in terms of prediction accuracy and AUC. In particular, group MCP penalized logistic regression predicts the best. Therefore, we propose group MCP penalized logistic regression to predict ovarian tumors.

Keywords: Ovarian benign tumors and malignant tumors; Group coordinate descent algorithm; Group LASSO/SCAD/MCP estimator

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1 Introduction

Ovarian cancer is a malignant tumor growing on the ovary. Its incidence rate is lower than that of cervical and endometrial cancer, whereas its mortality rate is higher than the sum of both cervical cancer and endometrial cancer and ranks first among gynecologic cancers. According to the global cancer data released by the World Health Organization International Agency in 2020, there were 314,000 new cases and 207,000 deaths of ovarian cancer in the world including 55,000 new cases and 38,000 deaths in China. Ovarian tumors are usually not difficult to diagnose. But benign and malignant diagnosis is not easy. Correct benign and malignant diagnosis need further auxiliary examinations, such as ultrasonography, cytology, laparoscopy, determination of tumor markers, radiologic diagnosis, etc. Computer tomography (CT) and magnetic resonance imaging (MRI) can clearly show the image of the tumor, and play a vital role in the diagnosis of ovarian tumors, in the observation of residual neoplastic changes at any time, and in the recurrence of the tumor. Positron emission tomography (PET) is also helpful in differentiating benign and malignant tumors and in diagnosing recurrent cancers. PET-CT has the dual functions of both CT and PET, and help better diagnosis. Advanced ovarian cancer is the leading cause of cancer death in women. Especially after chemotherapy, the recurrence rate was still over 70%. High degree of malignancy, high recurrence rate and poor prognosis from advanced ovarian cancer has become some prominent factors affecting the survival of ovarian cancer patients. Therefore, it is crucial to accurately diagnose ovarian benign tumors and malignant tumors.

There are many studies related to ovarian cancer, including the influencing factors analysis, screening methods, tumor markers, treatment and prognosis. Evaluated the value of tumor markers and clinical characteristics in making a differential diagnosis between MCT and squamous cell carcinoma arising from MCT, demonstrated that there were significant differences in age, tumor size, and levels of squamous cell carcinoma antigen (SCC), CA125, and CEA, as well as a significant difference in the CA19-9 level between MCT and squamous cell carcinoma arising from MCT, found that (1) age and tumor size are important factors in making a differential diagnosis and the optimal cutoff values for age and tumor size were, respectively, 45 years and 99 mm, (2) CEA was the best screening marker for squamous cell carcinoma arising from MCT, whereas age and tumor size were better markers than CA125 or CA19-9, and (3) SCC and CEA levels should be measured in patients age 45 years or older who have an MCT-like ovarian tumor larger than 99 mm in greatest dimension (Kikkawa et al. 1998). Using the Kaplan Meier method to study the prognostic impact of reproductive factors on survival rate after diagnosis of ovarian cancer, and found that high lifetime ovulatory cycles (LOC) and early age at menarche were associated with decreased survival after ovarian cancer (Robbins et al. 2009). Díaz-Padilla et al. summarized the clinical relevance of dynamic changes in CA-125 levels during the primary treatment of EOC and its potential influence both in the patient management and in the design of clinical trials in the adjuvant setting (Díaz-Padilla et al. 2012). Considering considered the contributions of the tumor markers CA125 and human epididymis protein 4 (HE4) as well as the risk ovarian malignancy algorithm (ROMA) and risk malignancy index (RMI) values, evaluated
their utility for establishing this system for patient referrals, and showed that there were no differences in accuracy between CA125, HE4, ROMA, and RMI for differentiating between types of ovarian masses (Anton et al. 2012). A meta-analysis was conducted to evaluate the diagnostic value of CA125, HE4 and ROMA in the diagnosis of ovarian cancer: systematically searched the PubMed and ScienceDirect databases and identified 32 studies that evaluated the role of CA125, HE4 and ROMA in diagnosing OCa, and observed that CA125 had a low specificity in premenopause and its diagnostic value was significantly lower than that postmenopausal, HE4 has higher specificity in premenopausal populations and can effectively reduce the misdiagnosis rate. Therefore, HE4 is suitable for diagnosing OC in premenopausal population, whereas CA125 and Roma are more suitable for the diagnosis of OC in postmenopausal population (Wang et al. 2014). Discussed current trends in diagnostic approaches, updated potential several panels of cancer biomarkers for early detection of ovarian cancer, reported that CA125 in combinations with two or more biomarkers have outperformed single biomarker assays for early detection of the disease, found that CA-125 with CA 19-9, EGFR, G-CSF, Eotaxin, IL-2R, vVCAM, MIF improved the sensitivity with 98.2% and specificity of 98.7% in early stage detection of ovarian cancer, and demonstrated a panel of biomarkers signature as the potential tool for prototype development in future and other advanced approaches for early diagnosis of ovarian cancer to avoid false-diagnosis and excessive cost (Muinao, Boruah & Pal 2019). Machine learning can be used to predict ovarian cancer, combined ovarian malignant tumor riskalgorithm (ROMA) with logistic regression model to accurately classify benign ovarian tumors (BOT) and Ovarian cancer (OC), and demonstrated that the machine learning approach had good potential in predictive modeling for the complex diseases, among others (Lu et al. 2020).

Logistic regression classifier is often used in disease diagnosis and finance forecasting. For high dimension classification problem, group penalized logistic regression classifier can improve the classification prediction performance by introducing group penalties to logistic regression. For example, studied group coordinate descent algorithms for nonconvex penalized regression including group SACD/MCP penalized logistic regression, founded that the estimated parameters converged to a global minimum when the sample size was larger than the dimension of the covariates, and converged to a local minimum otherwise, and found the group selection results of the MCP based and SCAD based GCD algorithms are better than the results selected by the group Lasso in terms of residual sum of squares and correct selection percentage (Wei & Zhu 2012). A block coordinate descent algorithm was used to fit group penalized multireponse and multinomial LASSO models, and verified the effectiveness of the algorithm (Simon, Friedman & Hasti 2013). The classification performance from the multinomial sparse group LASSO is studied based on a coordinate gradient descent algorithm, and showed that the multinomial group LASSO was significantly superior to the multinomial LASSO in terms of classification error rate and feature selection (Vincent & Hansen 2014). The semantic and phonological verbal fluency functional magnetic resonance imaging (fMRI) data was used to make a probabilistic diagnosis of depression, compared the performances of group lasso (gLASSO) and sparse group LASSO (sgLASSO) with standard LASSO(sLASSO), SVM and random forest, demonstrated that gLASSO and sgLASSO outperformed sLASSO in classification robustness, identification
of relevant brain regions and probability prediction were better than commonly used SVM and random forest (Sashimi et al. 2015). A credit scoring model was constructed based on group LASSO logistic regression, established three group LASSO models by AIC, BIC and cross-validation, and showed that the group LASSO method is superior to the backward elimination method in terms of interpretability and predictive accuracy (Chen & Xiang 2017). A multi-task sparsity group LASSO(MT-SGL) framework was used to deal with the loss function of generalized linear model to predict whether subjects are normal, mild cognitive impairment, or Alzheimer’s disease (Liu et al. 2017). Ghosal et al. proposed a new variable selection method in function linear concurrent regression, extended penalized variable selection methods (such as group LASSO, group SCAD and group MCP), and showed that the proposed method with group SCAD/MCP could pick out the relevant variables with high accuracy and had minuscule false positive and false negative rate even when data were observed sparsely, are contaminated with noise and the error process is highly non-stationary (Ghosal et al. 2020). In this paper, we combine group LASSO/SCAD/MCP with logistic regression, and propose group LASSO/SCAD/MCP penalized logistic regression classifier to investigate the benign and malignant ovarian cancer. Firstly, we select the 46 predictor variables like blood routine, general chemical detection, tumor markers and basic information etc., and divide the selected 349 ovarian cancer patients into the two sets: the training set for learning and the testing set for predicting. We develop the group coordinate descent algorithm and the training samples to obtain group LASSO/SCAD/MCP estimator, and apply the testing samples to establish two-class confusion matrix, prediction accuracy, sensitivity and specificity, draw the ROC curve and apply the area under ROC curve (AUC) to assess the prediction performance. Finally, we compare group LASSO/SCAD/MCP penalized logistic regressions with SVM and ANN, and found that the prediction accuracy and AUC for group MCP/SCAD/LASSO penalized logistic regression/SVM/ANN is 93.33%/85.71%/82.26%/74.29%/72.38% and 0.892/0.852/0.823/0.639/0.789, respectively. So group MCP penalized logistic regressions performs the best.

The rest is arranged as follows: Section 2 specifies data source, 49 features and their group processing. Section 3 constructs three group penalized methods. Section 4 reports model estimators and the prediction performances for the five methods. Section 5 is conclusion.

2 Data and Features

2.1 Data source

Here we choose the 349 ovarian cancer patients composed of 178 benign ovarian tumor and 171 ovarian cancer from the Third Affiliated Hospital of Suzhou University from July 2011 to July 2018 selected from the kaggle website (https://www.kaggle.com/saurabhshahane/predict-ovarian-cancer). The data set is divide into two parts: a training set composed of 70% ovarian cancer patients and a testing set composed of 30% ovarian cancer patients, see Table 1.
Table 1. The specific ovarian cancer patients

|                    | Benign ovarian tumor \((Y=0)\) | Ovarian cancer \((Y=1)\) | Total  
|--------------------|---------------------------------|--------------------------|-------
| Training set       | 98                              | 146                      | \(n_1=244\) |
| Test set           | 80                              | 25                       | \(n_2=105\) |
| Total              | 178                             | 171                      | \(n=349\) |

The chosen data set include 49 predictor variables: 19 blood routine tests, 22 general chemical tests, 6 tumor markers, age and menopause listed in Table 2. All patients experienced case diagnosis after operation, and none of them received preoperative radiotherapy and chemotherapy, and the histological type of diagnosis was classified according to the criteria of the World Health Organization.

Table 2. The 49 predictor variables and the response variable

| Notation | Variables | Definition                        | Value range          |
|----------|-----------|-----------------------------------|----------------------|
| X_1      | MPV       | Mean platelet volume              | 7.4 \sim 12.5(fL)    |
| X_2      | PLT       | Platelet count                    | 125 \sim 350(10^9/L) |
| X_3      | PDW       | Platelet distribution width       | 15.5 \sim 18.1(%)    |
| X_4      | PCT       | Thromboctocrit                    | 0.114 \sim 0.282(L/L)|
| X_5      | BASO#     | Basophil cell count               | 0 \sim 0.06(10^9/L)  |
| X_6      | BASO%     | Basophil cell ratio               | 0 \sim 1(%)          |
| X_7      | EO#       | Eosinophil count                  | 0.02 \sim 0.52(10^9/L)|
| X_8      | EO%       | Eosinophil ratio                  | 0.02 \sim 0.52(%)    |
| X_9      | NEU       | Neutrophil ratio                  | 40 \sim 75(%)        |
| X_10     | LYM#      | Lymphocyte count                  | 1.1 \sim 3.2(10^9/L) |
| X_11     | LYM%      | Lymphocyte ratio                  | 20 \sim 50(%)        |
| X_12     | MONO#     | Mononuclear cell count            | 0.1 \sim 0.6(10^9/L) |
| X_13     | MONO%     | Monocyte ratio                    | 3 \sim 10(%)         |
| X_14     | MCV       | Mean corpuscular volume           | 82 \sim 100(FL)      |
| X_15     | MCH       | Mean corpuscular hemoglobin       | 27 \sim 34(Pg)       |
| X_16     | RDW       | Red blood cell distribution width  | 10.6 \sim 15.5(%)    |
| X_17     | HGB       | Hemoglobin                        | 110 \sim 150(g/L)    |
| X_18     | RBC       | Red blood cell count              | 3.5 \sim 5.5(10^12/L)|
| X_19     | HCT       | Hematocrit                        | 0.35 \sim 0.45(L/L)  |
| X_20     | Mg        | Magnesium                         | 0.73 \sim 1.3(mmol/L) |
| X_21     | PHOS      | Phosphorus                        | 0.7 \sim 1.62(mmol/L) |
| X_22     | Ca        | Calcium                           | 1.12 \sim 1.32(mmol/L)|
| X_23     | Na        | Natrium                           | 137 \sim 147(mmol/L) |
| X_24     | K         | Kalium                            | 3.5 \sim 5.3(mmol/L) |
| Notation | Variables | Definition                                      | Value range            |
|---------|-----------|-------------------------------------------------|------------------------|
| $X_{25}$ | CL        | Chlorine                                        | $99 \sim 110$ (mmol/L) |
| $X_{26}$ | ALB       | Albumin                                         | $35 \sim 55$ (g/L)    |
| $X_{27}$ | TP        | Total protein                                   | $60 \sim 82$ (g/L)    |
| $X_{28}$ | GLO       | Globulin                                        | $20 \sim 40$ (g/L)    |
| $X_{29}$ | TBIL      | Total bilirubin                                 | $4 \sim 19$ ($\mu$mol/L) |
| $X_{30}$ | DBIL      | Direct bilirubin                                | $1.5 \sim 7$ ($\mu$mol/L) |
| $X_{31}$ | IBIL      | Indirect bilirubin                              | $2 \sim 15$ ($\mu$mol/L) |
| $X_{32}$ | GGT       | Gama glutamyltransferase                        | $3 \sim 73$ (U/L)     |
| $X_{33}$ | ALP       | Alkaline phosphatase                            | $25 \sim 130$ (U/L)   |
| $X_{34}$ | AST       | Aspartate aminotransferase                      | $6 \sim 40$ (U/L)     |
| $X_{35}$ | ALT       | Alanine aminotransferase                        | $1 \sim 45$ (U/L)     |
| $X_{36}$ | CA125     | Carbohydrate antigen 125                        | $0 \sim 35$ (U/mL)    |
| $X_{37}$ | CA19-9    | Carbohydrate antigen 19-9                       | $0 \sim 37$ (U/mL)    |
| $X_{38}$ | CA72-4    | Carbohydrate antigen 72-4                       | $0 \sim 7$ (U/mL)     |
| $X_{39}$ | AFP       | Alpha-fetoprotein                               | $0 \sim 7$ (ng/mL)    |
| $X_{40}$ | HE4       | Human epididymis protein 4                       | $0 \sim 140$ (pmol/L) |
| $X_{41}$ | CEA       | Carcinoembryonic antigen                        | $0 \sim 5$ (ng/mL)    |
| $X_{42}$ | CREA      | Creatinine                                      | $44 \sim 144$ ($\mu$mol/L) |
| $X_{43}$ | UA        | Urie acid                                       | $90 \sim 450$ (pmol/L) |
| $X_{44}$ | BUN       | Blood urea nitrogen                              | $1.7 \sim 8.3$ (mmol/L) |
| $X_{45}$ | AG        | Anion gap                                       | $8 \sim 30$ (mmol/L)  |
| $X_{46}$ | CO2CP     | Carban dioxide-combining power                  | $18 \sim 30$ (mmol/L) |
| $X_{47}$ | GLU       | Glucose                                         | $3.9 \sim 6.1$ (mmol/L) |
| $X_{48}$ | Age       | Age                                             | $15 \sim 83$          |
| $X_{49}$ | Menopause | Menopause                                       | Premenopausal=0,      |
|         |           |                                                 | Postmenopausal=1      |
| $Y$     | TYPE      | Tumor type                                      | BOT=0,OC=1           |

### 2.2 11 variable groups

In the raw data, total protein is the sum of albumin and globulin, total bilirubin is the sum of direct bilirubin and indirect bilirubin. Therefore, total protein and total bilirubin are deleted from 49 predictor variables. We observe that CA72-4 had 69% missing values. For convenience, we remove this variable. Therefore, total protein ($X_{27}$), total bilirubin ($X_{29}$) and carbohydrate antigen 72-4 ($X_{38}$) are removed, and the remaining 46 predictor variables with small missing values are filled with the mean value (the means of the missing variables is calculate separately.
According to check item, we divide the remaining 46 predictor variables into 11 different variable groups, where blood routine test usually included relevant factors of platelet, white blood cell and red blood cell, liver function and renal function test are the basic tests of physiological functions of liver and kidney, and chemical, acid-base balance and blood sugar are used to measure whether the human body is in balanced state, and tumor markers are used to detect ovarian cancer. In the following Table 3 specifies the 11 variable groups.

| Group       | Check item | Variables | Description                                                                 |
|-------------|------------|-----------|-----------------------------------------------------------------------------|
| Group 1     | Platelet   | $X_1, X_2, X_3, X_4$ | The main function of platelets is to accelerate coagulation, promote hemostasis and repair damaged blood vessels. |
| Group 2     | White blood cell | $X_5, X_6, X_7, X_8, X_9, X_{10}, X_{11}, X_{12}, X_{13}$ | White blood cells can phagocytose foreign materials to produce antibodies, and heal body damage, resist pathogen invasion and disease immunity. |
| Group 3     | Red blood cell | $X_{14}, X_{15}, X_{16}, X_{17}, X_{18}, X_{19}$ | The main work of red blood cells is to transport oxygen and carbon dioxide which can enhance phagocytosis and immune adhesion. |
| Group 4     | Chemical element | $X_{20}, X_{21}, X_{22}, X_{23}, X_{24}, X_{25}$ | Ions are used to measure human body electrolytes. The imbalance of the number of cations and anions will cause electrolyte disorders, which will lead different body damages. |
| Group 5     | Liver function | $X_{26}, X_{28}, X_{30}, X_{31}, X_{32}, X_{33}, X_{34}, X_{35}$ | Liver function examination generally includes protein metabolism function, bilirubin and bile acid metabolism function and serum enzyme indexes. |
Table 3. 11 variable groups

| Group   | Check item         | Variables          | Description                                                                                                                                 |
|---------|--------------------|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| Group 6 | Tumor marker       | $X_{36}, X_{37}, X_{39}, X_{40}, X_{41}$ | Tumor markers can be used for early detection, screening and differential diagnosis of tumors and can also be used for patient efficacy detection, recurrence and prognosis judgment. |
| Group 7 | Renal function     | $X_{42}, X_{43}, X_{44}$ | The main function of kidney is to secrete and excrete urine and toxins, regulate body fluids volume and water, and maintain the balance of body’s internal environment. |
| Group 8 | Acid-base balance  | $X_{45}, X_{46}$ | The pH value of normal people’s blood is always maintained at a certain level. Once the acid-base balance is disturbed, acidosis or alkalosis will occur. |
| Group 9 | Blood sugar        | $X_{47}$           | The glucose in the blood is called blood sugar. The production and utilization of blood sugar are in a state of dynamic balance to maintain the needs of various organs and tissues in the body. |
| Group 10| Age                | $X_{48}$           | Ovarian cancer has a certain relationship with age. The most common age group for ovarian cancer is middle-aged and elderly women, but many young women may also suffer from ovarian cancer. |
| Group 11| Menopause          | $X_{49}$           | Early and late amenorrhea have a certain impact on women’s physical health. The later the amenorrhea, the greater the risk of ovarian cancer. |

3 The three group penalized methods

3.1 Group LASSO penalized logistic regression

Tibshirani firstly introduced $L_1$ function to linear model, and proposed LASSO (Least Absolute Shrink and Selection Operator) penalized linear model for variable selection (Tibshirani
introduce group logistic regression, group LASSO penalized linear regression
and group LASSO penalized logistic regression are proposed. For example, Yuan & Lin (2006)
proposed group LASSO penalized linear regression and constructed its log likelihood function

\[ Q(\beta; \lambda) = \frac{1}{2} \left\| Y - \sum_{j=1}^{J} X_{(j)} \beta_j \right\|^2 + \lambda \sum_{j=1}^{J} \sqrt{d_j} \| \beta_j \|_2, \]  

(1)

where \( \beta = (\beta_1, \ldots, \beta_J) \) with the \( j \)-th group coefficient vector

\[ \beta_j = (\beta_{d_j-1+1}, \ldots, \beta_{d_j-1+d_j})^T \]  

(2)

is the whole coefficient vector, and \( d_j = \dim(\beta_j) \) is the length of the \( j \)-th group. The \( j \)-th
group LASSO estimator is given by

\[ \hat{\beta}_j = \left( 1 - \frac{\lambda \sqrt{d_j}}{\| S_j \|} \right) S_j; j = 1, \ldots, J, \]

(3)

where \( S_j = X_{(j)}^T (Y - \beta_{-j}) \) and \( \beta_{-j} = (\beta_{(1)}^T, \ldots, \beta_{(j-1)}^T, 0, \beta_{(j+1)}^T, \ldots, \beta_J^T) \). Meier, van de Geer
& Bühlmann (2008) combined group LASSO penalty with logistic regression and proposed group
LASSO penalized logistic regression (GLASSO) to investigate ovarian cancer. In this paper we
introduce group logistic regression

\[ \log \frac{P_\beta (X_i)}{1 - P_\beta (X_i)} = \beta_0 + \sum_{j=1}^{11} X_{i(j)}^\top \beta_j = \eta_i, i = 1, \ldots, 244, j = 1, \ldots, 11, \]  

(4)

to study the relation between \( Y \) and \( X = (X_{(1)}, \ldots, X_{(11)})^T \), where

\[ P_\beta (X_i) = P (Y = 1 \mid X_i; \beta) = \frac{\exp \left( \beta_0 + \sum_{j=1}^{11} X_{i(j)}^\top \beta_j \right)}{1 + \exp \left( \beta_0 + \sum_{j=1}^{11} X_{i(j)}^\top \beta_j \right)} \]

is the conditional probability of ovarian benign tumors, \( \beta_0 \) is the intercept, \( \beta_j \) is the \( j \)-th group
parameter vector and \( \beta = (\beta_1, \ldots, \beta_{(11)}) \) is the whole unknown parameter vector. Then, the
negative group log likelihood function for group logistic regression is

\[ L(\beta) = -l(\beta) = -\frac{1}{244} \sum_{i=1}^{244} \left\{ Y_i \log P_\beta (X_i) + (1 - Y_i) \log (1 - P_\beta (X_i)) \right\} \]

\[ = -\frac{1}{244} \sum_{i=1}^{244} \left\{ Y_i \left( \beta_0 + \sum_{j=1}^{11} X_{i(j)}^\top \beta_j \right) - \log \left[ 1 + \exp \left( \beta_0 + \sum_{j=1}^{11} X_{i(j)}^\top \beta_j \right) \right] \right\}. \]

(5)

Group LASSO penalized logistic log likelihood is
\[ Q(\beta; \lambda) = L(\beta) + \lambda \sum_{j=1}^{11} \sqrt{d_j} \| \beta_{(j)} \|, \]

where the tuning parameter \( \lambda \geq 0 \) controls the penalty size. Suppose that for a univariate \( Z \), the univariate soft-thresholding operator is

\[ S(Z, \lambda) = \begin{cases} 
Z - \lambda, & \text{if } Z > \lambda, \\
0, & \text{if } |Z| \leq \lambda, \\
Z + \lambda, & \text{if } Z < -\lambda, 
\end{cases} \]

and for a vector-valued argument \( Z \), the multivariate soft-thresholding operator is

\[ S(Z, \lambda) = S(\|Z\|, \lambda) \frac{Z}{\|Z\|}, \]

where \( Z/\|Z\| \) is the unit vector in the direction of \( Z \). In other words, \( S(Z, \lambda) \) acts on the vector \( Z \) by shortening it towards 0, and if the length of \( Z \) is less than \( \lambda \), the vector is shortened all the way to 0. Let \( \eta = \beta_0 + \sum_{j=1}^{11} X_{(j)}^T \beta_{(j)}, \eta_i = \beta_0 + \sum_{j=1}^{11} X_{(j)}^T \beta_{(j)}, v = \sup \eta_i \{ \nabla^2 L_i(\eta) \} \) with \( L_i(\eta) = Y_i \eta_i - \log(1 + e^\eta_i), i = 1, \ldots, 244 \), so that \( v I - \nabla^2 L(\eta) \) is positive semi-definite matrix at all points \( \eta \). Breheny & Huang (2011,2015) proposed group coordinate descent algorithm for group LASSO penalized logistic regression(GLASSO-PLR) as follows:

**Algorithm 1**  
Group coordinate descent algorithm for GLASSO-PLR

1. Let \( \eta = \beta_0 + \sum_{j=1}^{11} X_{(j)}^T \beta_{(j)}, \eta_i = \beta_0 + \sum_{j=1}^{11} X_{(j)}^T \beta_{(j)}, i = 1, \ldots, 244, \)
   \[ P = e^{\eta_i}/(1 + e^{\eta_i}), L_i(\eta) = Y_i \eta_i - \log(1 + e^{\eta_i}), v = \sup \eta_i \{ \nabla^2 L_i(\eta) \} \; ; \]
2. At the \( j \)-th step of \( m + 1 \) iterations, \( j = 1, 2, \ldots, 11 \), carry out the following (A)-(C):
   (A) compute \( \tilde{r} \leftarrow (Y - P)/v \) and \( Z_{(j)} = X_{(j)}^T \tilde{r} + \beta_{(j)} \);
   (B) update \( \hat{\beta}_{(j)}^{\text{GLASSO}}(m+1) \leftarrow S(v Z_{(j)}; \lambda_j)/v \);
   (C) update residual \( \tilde{r}_T \leftarrow \tilde{r} - X_{(j)}^T \left( \hat{\beta}_{(j)}^{\text{GLASSO}}(m+1) - \hat{\beta}_{(j)}^{\text{GLASSO}}(m) \right) \);
3. Update \( m \leftarrow m + 1 \);
4. Repeat 2 and 3 until convergence.

Introduce \( X_{-j} = \left( X_{(1)}, \ldots, X_{(j-1)}, 0^T, X_{(j+1)}, \ldots, X_{(11)} \right), P_i = e^{\eta_i}/(1 + e^{\eta_i}), \)
\[ \beta_{-j} = \left( \beta_{(1)}, \ldots, \beta_{(j-1)}, 0^T, \beta_{(j+1)}, \ldots, \beta_{(11)} \right), W = \text{diag} \{ P_i (1-P_i) \} \]
Compute \( \tilde{Y} = X^T \hat{\beta}_{(j)}^{\text{GLASSO}}(m) + W^{-1} (Y - P) \) and \( Z_{(j)} = X_{(j)}^T \left( \tilde{Y} - X_{-j} \beta_{-j} \right) \). We call the grprep package and obtain the group LASSO estimator

\[ \hat{\beta}_{(j)}^{\text{GLASSO}} = \frac{1}{v} S(v Z_{(j)}; \lambda \sqrt{d_j}) = \frac{1}{v} S(v Z_{(j)}; \lambda \sqrt{d_j}) \frac{Z_{(j)}}{\| Z_{(j)} \|}, j = 1, \ldots, 11. \]
3.2 Group SCAD penalized logistic regression

Fan & Li (2001) proposed the following SCAD penalty

\[ P_{SCAD}(\beta; \lambda, \gamma) = \begin{cases} 
\lambda \beta, & \text{if } \beta \leq \lambda, \\
\frac{2\lambda \gamma - (\beta^2 + \lambda^2)}{2(\gamma - 1)}, & \text{if } \lambda < \beta \leq \gamma \lambda, \\
\frac{\lambda^2(\gamma^2 - 1)}{2(\gamma - 1)}, & \text{if } \beta > \gamma \lambda.
\end{cases} \tag{10} \]

where \( \lambda > 0 \) and \( \gamma > 2 \). Its first derivative with respect to the parameter vector \( \beta \) is

\[ P'_{SCAD}(\beta; \lambda, \gamma) = \lambda \left\{ I(\beta \leq \lambda) + \frac{(\gamma \lambda - \beta)^+}{(\gamma - 1)\lambda} I(\beta > \lambda) \right\} = \begin{cases} 
\lambda, & \text{if } \beta \leq \lambda, \\
\frac{(\gamma \lambda - \beta)}{\gamma - 1}, & \text{if } \lambda < \beta \leq \gamma \lambda, \\
0, & \text{if } \beta > \gamma \lambda.
\end{cases} \tag{11} \]

Group penalized log-likelihood for group SCAD penalized logistic regression (GSCAD-PLR) is

\[ Q(\beta; \lambda, \gamma) = L(\beta) + \sum_{j=1}^{11} P_{SCAD}(\|Z(j)\|; \lambda \sqrt{d_j}, \gamma). \tag{12} \]

Similar to Algorithm 1, we apply the group coordinate descent algorithm for GSCAD-PLR and obtain the \( j \)-th group SCAD estimator

\[ \hat{\beta}^{GSCAD}_{(j)} = \begin{cases} 
\frac{1}{\sqrt{v}} S(vZ(j), \lambda \sqrt{d_j}), & \text{if } \|Z(j)\| \leq 2\lambda \sqrt{d_j}, \\
\frac{\gamma - 1}{\gamma - 2} \cdot \frac{1}{\sqrt{v}} S(vZ(j), \lambda \sqrt{d_j} \gamma), & \text{if } 2\lambda \sqrt{d_j} < \|Z(j)\| \leq \lambda \sqrt{d_j} \gamma, \gamma > 2, \\
Z(j), & \text{if } \|Z(j)\| > \lambda \sqrt{d_j} \gamma.
\end{cases} \tag{13} \]

3.3 Group MCP penalized logistic regression

Zhang (2010) proposed the nonconvex penalized function MCP (minimax convex penalty)

\[ P_{MCP}(\beta; \lambda, \gamma) = \begin{cases} 
\lambda \beta - \frac{\beta^2}{2\gamma}, & \text{if } \beta \leq \gamma \lambda, \\
\frac{2\lambda^2}{2}, & \text{if } \beta > \gamma \lambda.
\end{cases} \tag{14} \]

where \( \lambda > 0 \) and \( \gamma > 1 \). Its first derivative with respect to the parameter vector \( \beta \) is

\[ P'_{MCP}(\beta; \lambda, \gamma) = \lambda \left( 1 - \frac{\beta}{\lambda \gamma} \right) \text{sgn}(\beta) = \begin{cases} 
\lambda - \frac{\beta}{\gamma}, & \text{if } \beta \leq \lambda \gamma, \\
0, & \text{if } \beta > \lambda \gamma.
\end{cases} \tag{15} \]

where \( \text{sgn}(\beta) = -1, 0, 1 \) corresponds to \( \beta < 0, \beta = 0, \beta > 0 \), respectively. The group penalized log-likelihood for group MCP penalized logistic regression (GMCP-PLR) is

\[ Q(\beta; \lambda, \gamma) = L(\beta) + \sum_{j=1}^{11} P_{MCP}(\|Z(j)\|; \lambda \sqrt{d_j}, \gamma). \tag{16} \]
Similar to Algorithm 1, we apply the group coordinate descent algorithm for GMCP-PLR and obtain the $j$-th group MCP estimator

$$\hat{\beta}_{GMCP}^{(j)} = \begin{cases} \frac{\gamma}{\gamma - 1} \cdot \frac{1}{v} S(vZ(j), \sqrt{d_j} \lambda), & \text{if } \|Z(j)\| \leq \sqrt{d_j} \lambda \gamma, \gamma > 1, \\ Z(j), & \text{if } \|Z(j)\| > \sqrt{d_j} \lambda \gamma. \end{cases}$$

(17)

### 3.4 Two-class prediction accuracy evaluation

For two-class problem, a two-class confusion matrix (accuracy, sensitivity, and specificity), a ROC curve and the area under the ROC curve (AUC) are often used as the prediction performance evaluation indexes. Table 4 lists the two-class confusion matrix.

| Prediction Class 1 | True Class 1 | True Class 2 |
|--------------------|--------------|--------------|
| TP(True Positives) | FN(False Negatives) | FP(False Positives) |
| FN(False Negatives) | TN(True Negatives) |

From table 4, one can compute

- **Accuracy** = \frac{TP + TN}{TP + TN + FN + FP},
- **Precision** = \frac{TP}{TP + FP},
- **Sensitivity** = \frac{TP}{TP + FN},
- **Specificity** = \frac{TN}{TN + FP}.

The ROC curve can be drawn by changing (1 - specificity) and sensitivity at different thresholds. (1 - specificity) is x-axis that represents false positive rate (FPR). The smaller the FPR, the lower the false positive rate, and the less the actual negative class in the predicted negative class. Sensitivity is y-axis that represents true positive rate (true positive rate, TPR). The larger the TPR, the higher the hit, the more the actual positive class in the predicted positive class. The larger the AUC, the better the prediction performance. AUC ranges from 0 to 1: AUC ∈ (0, 0.5) reflects the worse prediction performance than random guess; AUC = 0.5 reflects the bad prediction performance like random guess; AUC ∈ (0.5, 0.7) reflects the low prediction performance, whereas for stock prediction, AUC ∈ (0.5, 0.7) reflects the good prediction performance; AUC ∈ (0.7, 0.9) reflects the relative high prediction accuracy; AUC ∈ (0.9, 1) reflects the very high prediction accuracy and AUC=1 reflects the perfect prediction performance.

### 3.5 Path selection

The tuning parameter $\lambda$ controls the size of the penalized strength. The larger $\lambda$, the stronger the penalized degree, the more coefficients are compressed to 0, and the less non-zero parameters are chosen. Therefore, the choice of the tuning parameter $\lambda$ is crucial. Commonly used methods are Akaike information criterion (AIC), Bayesian information criterion (BIC) and cross validation (CV). However, we are interested in obtaining $\hat{\beta}$ not just for a single value of $\lambda$, but for a range of values extending from a maximum value $\lambda_{max}$ for which all penalized
coefficients are 0 down to \( \lambda = 0 \) or to a minimum value \( \lambda_{\text{min}} \) at which the model becomes excessively large or ceases to be identifiable. Here, we consider \( \lambda \) on a grid \( \{ \lambda_0, \ldots, \lambda_{K+1} \} \), Then, we apply ten cross validation to select the optimal \( \lambda \), and obtain \( \hat{\beta} \) through the optimal \( \lambda \). For default \( \gamma \), group SCAD is 4 and group MCP is 3, the algorithm starts at \( \lambda_{\text{max}} \) and proceeding toward \( \lambda_{\text{min}} \). When the objective function is a strictly convex function, the estimated coefficients continuously vary within \( \lambda \in [\lambda_{\text{min}}, \lambda_{\text{max}}] \) and produce a regularized solution path. Algorithm 1 is an iterative algorithm, the maximum \( \lambda_{\text{max}} \) at \( \beta = 0 \) determined as the iterative initial value, \( \lambda_{\text{max}} = \max_j \{ v\|Z(j)\| \} \) for logistic regression, and going from the maximum toward to the minimum, \( \hat{\beta} \) obtained by the previous \( \lambda \) as the initial value for the next one to ensure that the initial value does not break away from solution.

4 Model estimators and prediction performances

4.1 Model estimators

Group LASSO/SCAD/MCP estimators for group penalized logistic regressions are obtained by the grpreg package. Firstly, the training set is orthogonalised within the group, the optimal \( \lambda \) is selected by ten fold cross validation, and apply the optimal \( \lambda \) and default \( \gamma \) and the formula (9),(13)and (17) to compute group LASSO/SCAD/MCP estimators. Then, the test set is used to compute a confusion matrix, accuracy, sensitivity, specificity and draw ROC curves so that one can compare the prediction accuracy. Fig.1 shows the coefficient path diagrams selected by group LASSO/SCAD/MCP penalty, where the abscissa is the \( \lambda \) value and the ordinate is the coefficient estimators \( \hat{\beta} \).

![Coefficient Path Diagrams](image)

**Fig. 1.** The coefficient path diagrams for group LASSO/SCAD/MCP.

The optimal \( \lambda \) selected by ten fold cross validation are listed in Fig.2. Ordinates represent cross validation errors, abscissa represents \( \log(\lambda) \) and the numbers above indicate the number of variables entered into the model at the corresponding \( \lambda \) value. Table 5 lists the optimal \( \lambda \) selected by ten fold cross validation based on ovarian tumors data sets for group LASSO/SCAD/MCP penalty.
Fig. 2. The cross validation errors diagrams for group LASSO/SCAD/MCP.

Table 5. Optimal lambda values for group LASSO/SCAD/MCP

| Group     | LASSO | SCAD | MCP  |
|-----------|-------|------|------|
| λ         | 0.0462| 0.0430| 0.0520 |

From Table 5, we found that the optimal $\lambda$ of group MCP penalized logistic regression is larger than group LASSO/SCAD penalized logistic regressions. Therefore, the penalization intensity of group MCP is greater, the more coefficients compressed to 0, and the fewer variable groups selected. After determining the optimal $\lambda$ selected by ten fold cross validation, we apply the group coordinate descent algorithm to obtain group estimators for group LASSO/SCAD/MCP penalized logistic regressions. The group coefficient estimators selected by group LASSO/SCAD/MCP penalty are listed in Table 6.

Table 6. Group LASSO/SCAD/MCP estimators

| Variable | Group LASSO | Group SCAD | Group MCP | Variable | Group LASSO | Group SCAD | Group MCP |
|----------|-------------|------------|-----------|----------|-------------|------------|-----------|
| Intercept| -1.7058     | -2.6146    | -4.8250   | $\beta_{(4-24)}$ | 0           | 0          | 0         |
| $\beta_{(1-1)}$ | -0.0781  | 0          | 0         | $\beta_{(4-25)}$ | 0           | 0          | 0         |
| $\beta_{(1-2)}$ | -0.0238  | 0          | 0         | $\beta_{(5-26)}$ | -0.0830     | -0.1234    | 0         |
| $\beta_{(1-3)}$ | -0.0213  | 0          | 0         | $\beta_{(5-28)}$ | -0.1639     | -0.1591    | 0         |
| $\beta_{(1-4)}$ | 0.0104   | 0          | 0         | $\beta_{(5-30)}$ | 0.0370      | 0.0395     | 0         |
| $\beta_{(2-5)}$ | 0         | 0          | 0         | $\beta_{(5-31)}$ | 0.1349      | 0.1628     | 0         |
| $\beta_{(2-6)}$ | 0         | 0          | 0         | $\beta_{(5-32)}$ | 0.0061      | 0.0180     | 0         |
| $\beta_{(2-7)}$ | 0         | 0          | 0         | $\beta_{(5-33)}$ | 0.1263      | 0.1025     | 0         |
| $\beta_{(2-8)}$ | 0         | 0          | 0         | $\beta_{(5-34)}$ | -0.1672     | -0.1532    | 0         |
| $\beta_{(2-9)}$ | 0         | 0          | 0         | $\beta_{(5-35)}$ | -0.1691     | -0.1593    | 0         |
| $\beta_{(2-10)}$ | 0        | 0          | 0         | $\beta_{(6-36)}$ | -0.2574     | -0.2659    | -17.3783  |
| $\beta_{(2-11)}$ | 0         | 0          | 0         | $\beta_{(6-37)}$ | 0.0988      | 0.1180     | 1.1620    |
| $\beta_{(2-12)}$ | 0         | 0          | 0         | $\beta_{(6-39)}$ | -0.0903     | -0.1089    | -5.6947   |
| $\beta_{(2-13)}$ | 0         | 0          | 0         | $\beta_{(6-40)}$ | -0.1336     | -0.1411    | -20.3392  |
| $\beta_{(3-14)}$ | 0         | 0          | 0         | $\beta_{(6-41)}$ | -0.0942     | -0.1122    | -2.8399   |
| $\beta_{(3-15)}$ | 0         | 0          | 0         | $\beta_{(7-42)}$ | 0.1868      | 0.3735     | 0         |
| $\beta_{(3-16)}$ | 0         | 0          | 0         | $\beta_{(7-43)}$ | 0.0229      | 0.0388     | 0         |
Table 6. Group LASSO/SCAD/MCP estimators

| Variable | Group LASSO | Group SCAD | Group MCP | Variable | Group LASSO | Group SCAD | Group MCP |
|----------|-------------|------------|-----------|----------|-------------|------------|-----------|
| \(\beta_{(3-17)}\) | 0 | 0 | 0 | \(\beta_{(7-44)}\) | -0.0045 | 0.0003 | 0 |
| \(\beta_{(3-18)}\) | 0 | 0 | 0 | \(\beta_{(8-45)}\) | 0 | 0 | 0 |
| \(\beta_{(3-19)}\) | 0 | 0 | 0 | \(\beta_{(8-46)}\) | 0 | 0 | 0 |
| \(\beta_{(4-20)}\) | 0 | 0 | 0 | \(\beta_{(9-47)}\) | 0 | 0 | 0 |
| \(\beta_{(4-21)}\) | 0 | 0 | 0 | \(\beta_{(10-48)}\) | -2.1425 | -3.3630 | 0 |
| \(\beta_{(4-22)}\) | 0 | 0 | 0 | \(\beta_{(11-49)}\) | -0.0528 | 0 | 0 |
| \(\beta_{(4-23)}\) | 0 | 0 | 0 | | | | |

From Table 6, we can see that group LASSO penalized logistic regression retains the 6 groups composed of 22 non-zero explanatory variables in 6 groups and compresses the other 5 groups composed of 24 explanatory variables to 0. Group SCAD penalized logistic regression retains the 4 groups composed of 17 non-zero explanatory variables. Group MCP penalized logistic regression selected tumor marker group among 11 groups of explanatory variables, including CA125, CA19-9, AFP, HE4 and CEA, and the coefficients of the remaining explanatory variables are compressed to 0, indicating that the 10 variable groups have no significant influence on the discrimination of benign and malignant ovarian tumors. Therefore, group MCP penalized logistic regression can effectively predict the benign and malignant of ovarian tumors. Its probability estimators are as follows:

\[
P(Y_i = 1|X_i) = \frac{e^{-4.8250-17.3783X_{36}+1.1620X_{37}-5.6947X_{39}-20.3392X_{40}-2.8399X_{41}}}{1 + e^{-4.8250-17.3783X_{36}+1.1620X_{37}-5.6947X_{39}-20.3392X_{40}-2.8399X_{41}}},
\]

\[
P(Y_i = 0|X_i) = \frac{1}{1 + e^{-4.8250-17.3783X_{36}+1.1620X_{37}-5.6947X_{39}-20.3392X_{40}-2.8399X_{41}}},
\]

We conclude that the tumor marker group is the critical indicator for the diagnosis of benign and malignant ovarian tumors.

4.2 Prediction performances

In the following we apply group LASSO/SCAD/MCP penalized logistic regressions to predict benign and malignant ovarian tumors. Apply the test set \(\{(X_i, Y_i), i = n_1 + 1, \ldots, n_1 + n_2\}\) with \(n_1 = 244\) and \(n_2 = 105\) to the probability estimators, and estimate the predicted values \(\hat{Y}_i\) according to the following rules:

\[
\text{If } \hat{P}_i > c, \text{ then } \hat{Y}_i = 1, \text{ else } \hat{Y}_i = 0, \quad (18)
\]

where \(c\) is a given threshold. For balanced data, \(c\) is generally taken as 0.5. For unbalanced data, Youden index is widely used to select the optimal threshold (Raghavan, Ashour & Bailey, 2016).
To evaluate the prediction performance, we compare group LASSO/SCAD/MCP penalized logistic regressions with SVM and ANN. The confusion matrixes and the prediction performances from the five methods are listed in Table 7 and Table 8, respectively.

Table 7. Two-class confusion matrix comparisons

|        | 1(OC) | 0(BOT) |
|--------|-------|--------|
| Group LASSO | 21    | 14     |
| 0(BOT)     | 4     | 66     |
| Group SCAD | 21    | 11     |
| 0(BOT)     | 4     | 69     |
| Group MCP  | 22    | 4      |
| 0(BOT)     | 3     | 76     |
| SVM        | 14    | 16     |
| 0(BOT)     | 11    | 64     |
| ANN        | 24    | 28     |
| 0(BOT)     | 1     | 52     |

Table 8. The prediction performance comparisons

| Model       | Accuracy | Precision | Specificity | Sensitivity |
|-------------|----------|-----------|-------------|-------------|
| Group LASSO | 0.8286   | 0.6000    | 0.8250      | 0.8400      |
| Group SCAD  | 0.8571   | 0.6563    | 0.8625      | 0.8400      |
| Group MCP   | 0.9333   | 0.8462    | 0.9500      | 0.8800      |
| SVM         | 0.7429   | 0.4667    | 0.8000      | 0.5600      |
| ANN         | 0.7238   | 0.4615    | 0.6500      | 0.9600      |

According to Table 8, the prediction performance of group penalized logistic regression is better than that of machine learning method. The prediction accuracy, precision and specificity for group MCP penalized logistic regression are 93.33%, 84.62% and 95%, respectively. The prediction accuracy for group SCAD penalized logistic regression is 85.71%. The predictive accuracy for group LASSO penalized logistic regression is 82.86%. The prediction accuracy of both SVM and ANN are 74.29% and 72.38%, respectively. ANN predicts the worst with a prediction accuracy that differed by 20.95% from the group MCP penalized logistic regression, as well as the lowest precision and specificity with 46.15% and 65%, but the sensitivity is 96% that is the highest. The prediction performance for SVM is slightly better than that of ANN with a prediction accuracy of 74.29%, and its prediction sensitivity is 56% that is also the lowest. The ROC curve and the area under the ROC curve (AUC) are often used to evaluate the predictive performance. The larger the AUC, the better the predictive performance. Here we apply pROC package to visualize their ROC curves, see Fig.3.
As shown in Fig. 3, the sensitivity and specificity of several models are consistent with Table 8, the optimal thresholds for group LASSO/SCAD/MCP penalized logistic regressions are 0.649, 0.740 and 0.988, respectively, and the AUC values are 0.823, 0.852, 0.892, respectively. The optimal threshold for SVM is 0.726 and the AUC value is 0.639. The optimal threshold value for ANN is 0.997 and the AUC value is 0.789. We found that the AUC values for group penalized logistic regressions have exceeded 0.8 that reflects very high prediction accuracy. AUC values for both SVM and ANN are below 0.8 that reflects the relatively poor prediction performance. Thus, group MCP penalized logistic regression has the highest AUC value and the best prediction performance in benign and malignant ovarian tumors prediction problem.

5 Conclusion

In this paper we propose group LASSO/SCAD/MCP penalized logistic regressions to predict ovarian tumors. We select 46 explanatory variables and divide them into 11 variable groups, develop the three group penalized methods to select the significant variable groups and found that the group of tumor markers is the key variable group to predict the benign or malignant ovarian tumors, where the optimal tuning parameters are selected by ten folds cross-validation. Group LASSO penalized logistic regression retains 6 variable groups composed of 22 explanatory variables. Group SCAD penalized logistic regression retains 4 variable groups composed of 17 non-zero explanatory variables. Group MCP penalized logistic regression only selects 1 tumor marker group composed of 5 explanatory variables: CA125, CA19-9, AFP, HE4 and CEA. Finally, we compare the proposed group LASSO/SCAD/MCP penalized logistic regression with ANN and SVM, and found that the prediction accuracy and AUC for group MCP/SCAD/LASSO penal-
ized logistic regression/SVM/ANN is 93.33%/85.71%/82.26%/74.29%/72.38% and 0.892/0.852 /0.823/0.639/0.789, respectively. Obviously, the proposed group MCP penalized logistic re-
gression performs the best. Therefore, we propose group MCP penalized logistic regression to predict ovarian cancer.

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Availability of data The data set used in this study are available from https://www.kaggle.com/saurabhshahane/predict-ovarian-cancer.

Declarations

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.

Ethic approval This is an observational study and do not require ethics approval.

Consent to publication All the authors consented to the publication of this article.
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