A high serum phosphate and calcium-phosphate product is associated with cerebral small vascular disease in patients with stroke; a real world study.

Wenjing Lv  
Qingdao University

Zixuan Wang  
Qingdao University

Junqi Jiang  
Qingdao University

BinBin Deng  
Wenzhou Medical University First Affiliated Hospital  
https://orcid.org/0000-0002-4058-0738

Qingxia Lin  
Wenzhou Medical University

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Cerebral small vessel disease (CSVD) is a very common neurological whole-brain disease in older people, which results from pathologies in cerebral arterioles, capillaries and venules. It causes up to 45% of dementia, 20% of all stroke, and cognitive impairment, depression and gait problems are also frequently seen in CSVD patients (Pantoni, 2010). The diagnosis of CSVD has relied on imaging findings including lacunes, white matter hyperintensities (WMHs), and microbleeds (CMBs) (Shi & Wardlaw, 2016). Epidemiology of CSVD might differ in Asian compared with non-Asia population (Shi & Wardlaw, 2016). CSVD has a robust association with large artery atherosclerosis, and often accompanies by stroke in USA (Del Brutto et al., 2020). It is often ignored when accompanied by stroke among inpatients, since its manifestation is less devastating and progress slower than large vessel occlusion. Few studies investigate the epidemiology of CVSD accompanied by stroke in China. Antiplaatlet and management of traditional risk factors still remain the most important therapeutic and preventive approach, due to limited understanding of CSVD, however, vascular risk factors and large artery disease explain quite small part of the variance of CSVD (Wardlaw et al., 2014). Exploring the risk factor of CSVD is necessary for targeted prevention and treatment of CSVD.

Artery calcication is part of the atherosclerotic process, it is considered as ectopic deposition of bone components and is highly associated with WMHs, lacunes, and CMBs (Chen et al., 2019). Serum phosphate (P), calcium (Ca) and calcium-phosphate product (Ca×P) is calcium and phosphorus metabolism (CPM)-related factors. In extreme cases, such as chronic kidney diseases (CKD), CPM homeostasis disruption can promote ectopic vascular calcification in blood vessel, including elevated P levels and increased Ca×P (Reiss et al., 2018). Lowering dialysate Ca concentration can improve the aortic stiffness and decrease the carotid intima-media thickness (He et al., 2016). However, CPM disorders are not quite common risk factors in stroke, and less attention has been paid to the potential involvement of CPM in CSVD patients who without CPM disorders.

Materials And Methods

This study was approved by the Ethics Committee of the First Affiliated Hospital of Wenzhou Medical University. The experiments were undertaken with the understanding and written consent of each subject, and that the study conforms with World Medical Association Declaration of Helsinki. From 2016.4.1 to 2018.7.31, there were 1102 ischemic stroke patients were admitted to the First Affiliated Hospital of Wenzhou Medical University. 588 of them were included in this study. 140 of whom had no CSVD signs served as ‘no CSVD’ group, 448 of whom had any signs of CSVD in the cranial MR images served as ‘CSVD’ group. 448 of the include cases had lacunes, 212 of them had scattered lacunes, and 236 of them had multiple lacunes. 100 of the included cases had WMHs, and 55 of the included cases had CMBs. Cases with overlapping subtypes were repeated included in each subtype.

Exclusion criteria

Patients lacked of serum Ca and P tests. Patients had any diseases that could influence CPM, including chronic kidney disease with blood creatinine level>186 µmol/L, bone metastasis of malignant tumor, parathyroid disease, pituitary disorder. The demographic, chronic disease, hematologic parameters, and imaging findings of all patients were collected using standardized data records.

Clinical and laboratory assessments

All blood indicators were assessed on the morning of the second day of admission after overnight fasting. Cranial MR examinations were completed within 3 days of admission. Vascular risk factors of cases were collected according to patient description.

Classification criteria

Lacunes are shown as hypointense lesions with hyperintense rim on FLARE and T1, with hyperintense signal on T2. The diameters are 3-15mm. It can be rounded, ovoid or tubular, and can distributed in the white matter, basal ganglia, pons or brainstem. Recent small ischemic lacunes is included, that are best identified on DWI, as hyperintense on DWI:T2 and FLAIR, hypointense on T1, with usual diameter ≤20mm. WMHs are usually symmetrically and bilaterally distributed in the white matter including the pons and brain stem, and also occur in deep grey matter. They appear hyperintense to the normal brain on T2 or FLAIR, and can be patchy or confluent. CMBs are regarded as small round and homogeneous foci of hypointensity on T2-weighted MRI and susceptibility-weighted imaging. CSVDs are with single or combined signs of lacunes, WMHs and CMBs (Shi & Wardlaw, 2016).

Statistical analysis
All baseline data and risk factors of excluded and include cases were statistically analyzed by logistic regression, using the probability as propensity score. Their distribution were viewed in a population pyramid graph. For baseline data and risk factors of included cases, categorical variables were compared with the chi-square test, and continuous variable were compared with the Mann-Whitney U test, p-value<0.05 was considered statistically significant. Logistics regression analysis of univariate was used for identifying risk factors, and factors with p value <0.1 were analyzed in multivariate analysis with logistic regression, with a method of Backward LR. The nomograms were constructed performed with R. After logistic regression analysis and calculation of probabilities in multivariate analysis, we assessed the performance of the model by calculating the AUC. The larger the AUC, the more accurate the prognosis is. H-L test was used to evaluate the calibration degree of models, models with p value>0.1 were considered in good calibration degree. All calculations were based on SPSS version 22.0 software and R.

Results

Clinical information of 1102 stroke patients with cranial MR were collected, including demographic parameters (age, gender, weight, ), vascular risk factors (hypertension, diabetes, coronary artery disease, atrial fibrillation, hyperlipidemia, history of smoking and drinking alcohol), blood biochemistry and vital signs on admission (NIHSS score, MBP, FGB, P2hPG, HbA1c, TC, TG, LDL, HDL, LVEF, WBC, Neutrophil, Lymphocyte, RBC,HB, TP, Albumin, PLT, PT, INR, APTT, FIB,BUN, Cr, HCY, TSH, FT3, FT4), serum biochemical markers of CPM (P, Ca, adjusted Ca, Ca×P, adjusted Ca×P). Patients lack of serum Ca and P data, or with CPM disorders, such as severe chronic kidney disease, pituitary or parathyroid disease, bone metastasis of malignant tumor, were excluded (Fig. 1). In order to check if there was selection bias, we calculated propensity scores of demographic parameters, vascular risk factors, blood biochemistry and vital signs on admission in excluded and included cases respectively. Distributions of propensity score were similar between groups (Fig. 2).

Patients with any one or more image representations of lacunes, WMHs, CMBs were divided into CSVD group, and patients without CSVD signs were divided into no CSVD group. There were 140 cases in no CSVD group and 448 cases in CSVD group. Up to 76.19% of stroke patients, had signs of CSVD on cranial MR images (Fig. 3A). TOAST classification distributions had no statistical difference among no CSVD and CSVD groups (Fig. 4). The general characteristics of two groups were detailed in Table 1. Age, rate of hypertension or diabetes, serum P, Ca, Ca×P, adjusted Ca×P were higher, and WBC, HB were lower in CSVD patients than no CSVD patients (Table 1, Fig. 5).

Univariate and multivariate logistic regression were used to identify factors potentially prognostic factors of CSVD, and further construct models to predict CSVD. Receiver operating characteristic (ROC) curves were used as the discrimination index of models. Hosmer-Lemeshow goodness of fit (H-L) test were used to evaluate the calibration degree of models. All factors with p value <0.2 in table 1 were analyzed by univariate logistic regression (Table 3). Baseline factors with p value <0.1 in table 3, including age, hypertension, diabetes, MBP, WBC, RBC, HB, BUN were analyzed with multivariate logistic regression, and factors with p value <0.05, including age, hypertension, diabetes, P, Ca, adjusted Ca×P were included in model 1 ultimately (Table 2). AUCs is 0.771 and p value of H-L test is 0.873>0.05 (Fig.3).

Based on model 1, we constructed model 2 (including P, Ca or adjusted Ca), model 3 (including Ca×P), model 4 (including adjusted Ca×P) (Table 2). In these three models, only age and hypertension were still remained in the end. P was included in model 2, Ca×P was included in model 3, and adjusted Ca×P was included in model 4 (Table 2). AUCs were similar among three models, 0.855, 0.851, 0.853, respectively, better than model 1 (Fig. 6A). P value of H-L test is 0.689, 0.698, 0.307, respectively (Fig. 6B). Adjusted Ca×P impaired the calibration degree in model 4, and was no better than Ca×P hence, there was no need to use adjusted Ca×P to predict CSVD instead of Ca×P. Model 2 and model 3 had similar discrimination and calibration power, thus, we selected them to establish nomograms to predict the probability of CSVD (Fig. 7). Age, hypertension, P or Ca×P were all positive factors related to CSVD. The OR of P was 3720.401 (646.665:21404.249), and OR of Ca×P was 1.294, 1.222-1.370 (Table 2). P, Ca×P are potential CSVD risk factors.

We explored the incidence of subtypes of CSVD, including lacunes, WMHs, and CMBs, and patients with lacunes were further divided into scattered lacunes and multiple lacunes. Then we validate if model 2 and model 3 were valid in these subtypes. We found that lacunes was the most common types in CSVD, 22.96% of the CSVD patients have two or three types coexisted (Fig. 4A). P and Ca×P were significantly increased in lacunes than no CSVD, and was highest in multiple lacunes. Ca is slightly higher in multiple lacunes than no CSVD. P and Ca×P were significantly increased in WMHs and CMBs, compared with no CSVD. Ca were slightly increased in WMHs than no CSVD (Fig. 4B-2D). P and Ca×P were still positive risk factors in each subgroup of CSVD (Table 4-6). AUCs and H-L test shows good discrimination and calibration power of models in each subgroup (Fig. 3, Table. 7).

Discussion

Using a hospital stroke registry of the First Affiliated Hospital of Wenzhou Medical, we found that 76.19% of relatively normal kidney function stroke patients concomitant with CSVD. Higher serum P and Ca×P levels were positively linked to CSVD, and subtypes of CSVD, including lacunes, WMHs, and CMBs. Most CPM related clinical investigations are conducted in chronic kidney disease population, and takes large blood vessel calcification as research target(Felsenfeld, Levine, & Rodriguez, 2015; Reiss et al., 2018). High P and Ca could promote large vessel calcification, and their influences can mutually reenforcing (Fu, Cui, Ning, Fu, & Liao, 2015; Masumoto et al., 2017; Panizo et al., 2016). CSVD is concomitant with stroke in South China, indicating they may share some common risk factors or pathologies, although they are two different kinds of diseases. Our results indicates that higher serum P and Ca×P may promote CSVD even when they are in normal levels. Studies of CSVD and CPM have some discrepancies in different regions. A study in Taiwan shows that circulating P levels > 3.925 mg/dl were associated with severe WMHs in a community-based longitudinal aging Study(Chung et al., 2019). A study in Turkey follows subjects who underwent brain imaging for any reason, reported that CSVD patients had lower P than normal subjects (Ozelsancak, Micokzakdoglu, Torun, & Tekkarismaz, 2019). In US veterans population, high serum P is associated with CSVD and dementia, and this relationship is more significantly in participants <60 years old (Li, Xie, Bowe, Xian, & Al-Aly, 2017). CPM is significant affected by light and diet in normal population, further large-scale cohort studies on relationship of CPM and CSVD tracking of localized community are need.
The primary underlying initiating cause of CSVD is the derangement of the blood-brain barrier (BBB), this may start some years before the first symptoms, leads to the small vessel structural changes and perivascular changes (Liu, Yang, & Fan, 2020), the cellular structure of the BBB is mainly composed of brain microvascular endothelial cells (MVECs) (Wardlaw, 2010). Junction protein (zona occludens-1, occludin, and claudin-5) expression is downregulated in human brain MVECs after high phosphorus treatment (Chung et al., 2019). Many BBB dysfunction diseases are associated with increasing intracellular Ca levels of MVECs and serum Ca. Many cell membrane ion channels are responsible for regulating intracellular Ca balance. Plasma membrane calcium ATPase could export calcium, and calcium entry from the extracellular space can occur from store-operated cation channels or receptor-operated cation channels (Dalal, Muller, & Sullivan, 2020). In addition, voltage-gated L-type calcium channels and the transient receptor potential superfamily have been identified to comprise calcium influx channels in endothelial cell (Jiang et al., 2018).

High levels of phosphate accelerate the precipitation of calcium and phosphate in the form of hydroxyapatite, accelerating vascular calcification (Reynolds et al., 2004). Calcification is associated independently with high calcium levels, and is synergistic with elevated phosphate levels (Shroff et al., 2010) (Schurgers et al., 2018). High phosphate can directly induces phenotypic transformation of vessel smooth muscular cells (VSMCs) into osteoblasts by stimulating VSMCs to express core binding factor-a1, and repress the production of calcification inhibitors and promote the release of extracellular vesicles (EVs) lacking these inhibitors, but rich in pro-calcic proteins such as tissue-nonspecific alkaline phosphatase (Yang et al., 2019). miR-29b, miR-133b, and miR-211 have direct roles in the vascular smooth muscle calcification induced by high phosphorus (Panizo et al., 2016). In VSMCs, the sodium-phosphorus co-transporter PiT-1 promotes matrix calcification caused by elevated phosphorus, while PiT-2 inhibits its changes (Yamada & Giachelli, 2017). Vitamin K-dependent matrix Gla protein (MGP) is a key inhibitor in the formation of vascular calcification, and elevated phosphate and calcium levels increased MGP levels, as an inherent protective mechanism of vascular calcification (Houben, Neradova, Schurgers, & Vervloet, 2016).

Although this study showed propensity score distribution of the included and excluded data to confirm that the baseline characteristic of the excluded cases was similar to that of the included cases. However, the analyzed data was from a single-center medical institution in Southern China. The population in the hospital had higher health literacy, compared to those who did not seek medical attention; the selected population had geographic and ethnic specificities; awareness rate of vascular risk factors varied. All above factors lead to an inevitable selection bias and recalling bias of the study. Further multi-center, multi-ethnic, community-based research should be conducted.

In conclusion, serum P and Ca×P was higher in CSVD patients, including subtypes of lacunes, especially multiple lacunes, WMHs, and CMBs. Patients with elder age, hypertension, higher P or Ca×P had an increased risk of CSVD in stroke patients who did not have CPM disorder. Higher P or Ca×P are also risk factors of CSVD subtypes, including lacunes, WMHs, and CMBs.

**Abbreviations**

Mean blood pressure (MBP); left ventricular ejection fraction (LVEF); fasting blood glucose (FBG); 2-hour post-meal blood glucose (P2hPG); glycated hemoglobin (HbA1c%); total cholesterol (TC); triglyceride (TG); low density lipoprotein cholesterol (LDL); high density lipoprotein cholesterol (HDL); white blood cell (WBC); red blood cell (RBC); blood platelet (PLT); hemoglobin (HB); total serum protein (TP); total prothrombin time (PT); international normalized ratio (INR); activated partial thromboplastin time (APTT); fibrinogen (FIB); urea nitrogen (BUN); creatinine (Cr); homocysteine (HCY); thyroid stimulating hormone (TSH); free triiodothyronine (FT3); free tetraiodothyronine (FT4); National Institutes of Health Neurological Decit Score (NIHSS); cerebral small vascular disease (CSVD); white matter hyperintensities (WMHs); cerebral microbleeds (CMBs), calcium and phosphorus metabolism (CPM); chronic kidney diseases (CKD); microvascular endothelial cells (MVECs); vessel smooth muscular cells (VSMCs); extracellular vesicles (EVs); matrix Gla protein (MGP)

**Discussion**

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**Ethics approval:** The Ethical Decision Committee of the Research Administration at First Affiliated Hospital of Wenzhou Medical University approved the study (CR2009041).

**Consent to participate:** All patients agreed to participate in the study, and use their clinic data and information for research purpose.

**Consent for publication:** All participants agreed for publications related to this study.

**Availability of data and material:** Data and material can be shared with the consent of corresponding authors.

**Code availability:** SPSS version 22.0 software and R.

**Authors’ contributions:** Wenjing Lv and Zixuan Wang were responsible for data statistics and writing paper. Junqi Jiang collected data. Binbin Deng and Qingxia Lin provided resources and designed the study.

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### Table 1 The general characteristics of no CSVD and CSVD patients.

| Baseline characteristics | No CSVD | CSVD | P-value |
|--------------------------|---------|------|---------|
| N                        | 140     | 445  |         |
| Age, years               | 58.8±2.99 | 70 (62.77) | .000 |
| Femal                    | 45 (32.1%) | 155 (34.59%) | .661 |
| Hypertension             | 100 (71.43%) | 393 (88.51%) | .000 |
| Diabetes                 | 39 (27.86%) | 181 (40.58%) | .007 |
| Coronary artery disease  | 18 (13.95%) | 76 (18%) | .349 |
| Atrial fibrillation      | 19 (13.57%) | 68 (15.21%) | .685 |
| Hyperlipidemia           | 69 (49.64%) | 216 (48.98%) | .923 |
| History of smoking       | 69 (49.29%) | 191 (42.73%) | .174 |
| Drinking alcohol         | 54 (38.57%) | 147 (32.89%) | .222 |
| Weight, kg               | 65 (60.69) | 65 (59.7) | .369 |
| NIHSS score              | 2 (1.4) | 2.5 (1.4) | .864 |
| MBP, mmHg                | 104.04±14.63 | 106.67 (97.12, 67.7) | .000 |
| FBG, mmol/L              | 5.15±(4.5,6.55) | 5.2 (4.5,6.65) | .655 |
| P2hPG, mmol/L            | 8.58±(6.63,11.84) | 8.18 (6.37,11.3) | .281 |
| HaA1C, %                 | 6.1 (5.6,7.18) | 6.1 (5.6,7) | .590 |
| TC, mmol/L               | 4.51±(3.82,5.08) | 4.53 (3.79,5.32) | .612 |
| TG, mmol/L               | 1.6 (1.14,2.23) | 1.5 (1.14,2.06) | .146 |
| LDL, mmol/L              | 2.55 (2.07,3.07) | 2.57 (2.08,3.2) | .521 |
| HDL, mmol/L              | 0.99±(0.84,1.178) | 1.94 (0.8,1.22) | .117 |
| LVEF, %                  | 65 (61.5,68.7) | 65.3 (61.6,68.6) | .984 |
| WBC, ×10⁉/L              | 7.01±(5.66,8.55) | 6.54 (5.43,7.82) | .023 |
| Neutrophil, ×10⁹/L       | 4.34±(3.36,5.64) | 4.02 (3.15,5.33) | .080 |
| Lymphocyte, ×10¹²/L      | 1.7±(1.3,2.18) | 1.69 (1.31,2.05) | .529 |
| RBC, ×10¹²/L             | 4.55±(4.4,4.83) | 4.5 (4.1,4.83) | .080 |
| Hb, g/L                  | 139 (128.25,148.75) | 135 (124,146) | .032 |
| TP, g/L                  | 67.25 (63.25,69.75) | 67.1 (63.7,70.2) | .465 |
| Albumin, g/L             | 37.75 (34.8,39.58) | 38 (35,36,39.9) | .329 |
| PLT, ×10⁹/L              | 218 (172.25) | 224 (184,202) | .135 |
| Pt, s                    | 13.5±(13.1,14.28) | 13.5 (13.14) | .367 |
| INR                      | 1.04±(0.99,1.11) | 1.03 (0.98,1.09) | .483 |
| ApTt, s                  | 36.9±(34.03,39.58) | 37.1 (34.9,39.6) | .358 |
| FIB, g/L                 | 3.48±(2.98,4.24) | 3.63 (2.96,4.41) | .271 |
| BUN, mmol/L              | 4.8±(3.86) | 5 (4.1,6.2) | .083 |
| Cr, μmol/L               | 71 (60.83,75) | 68 (58.8) | .599 |
| Hcy, μg/L                | 10 (8.12) | 10 (8.13) | .055 |
| TSH, mIU/L               | 1.46±(0.99,2.39) | 1.62 (1,05,2.39) | .563 |
| FT3, pmol/L              | 4.4 (4.78) | 4.5 (4.1,4.8) | .235 |
| FT4, pmol/L              | 11.23 (10,12.79) | 11.23 (10,14,12.62) | .588 |
| P, mmol/L                | 0.81±(0.73,0.98) | 1.06 (0.95,1.19) | .000 |
| Ca, mmol/L               | 2.18±0.11 | 2.2 (2.13,2.28) | .022 |
| Adjusted Ca*, mmol/L     | 2.22±0.106 | 2.23 (2.17,2.29) | .135 |
| CaXP, (mg/dl)²           | 21.36 (19.1,26.29) | 28.94 (25.6,32.98) | .000 |
| Adjusted CaXP, (mg/dl)²  | 21.84 (20.09,26.29) | 29.22 (25.98,33.28) | .000 |

Normally distributed data is represented by mean±SD, skewed distribution data is represented by median (IQR), and frequency data is represented by n (%). Adjusted Ca (mmol/L) = Ca (mmol/L)+0.02[40-Albumin (g/L)], serum Ca is adjusted if albumin<35g/L, or >51g/L. Factors with P<0.05 is statistically significant.

### Table 2 Multivariate analysis for models by Binary Logistic Regression.

| P-value | Model 1 OR | 95%CI | Model 2 OR | 95%CI | Model 3 OR | 95%CI | Model 4 OR |
|---------|------------|-------|------------|-------|------------|-------|------------|
| Age     | .000       | 1.080 | 1.099-1.110| .027  | 1.071-1.072| .000  | 1.048      |
| Diabetes| .010       | 1.896 | 1.110-3.384| .002  | 2.745      |       | .001       |
| MBP     | .010       | 1.023 | 1.004-1.043| .017  | 1.037      |       | .001       |
| WBC     | .010       | 0.889 | 0.811-0.974| .000  | 0.811-0.974| .000  | 0.810      |
| P       | .000       | 37.20401 | 64.665-21404.249 | .000  | 1.294      | 1.222-1.370| .000  |

Model 1: adjusted for covariates that p<0.1 in univariate analysis. Model 2= model 1+P, Ca or adjusted Ca. Model3= model 1+ CaXP. Model4= model 1+ adjusted CaXP. A method of Backward LR are used, and factors with p value>0.05 are removed from models. NE: not entry.

### Table 3. Univariate analysis for factors associated with CSVD by Binary Regression.
Baseline characteristics & OR & 95%CI & P-value  
\hline
**N 140/448**  
Age, years & 1.077 & 1.058-1.097 & .000  
Hypertension & 3.090 & 1.934-4.937 & .000  
Diabetes & 1.769 & 1.168-2.679 & .007  
History of smoking & .772 & .527-1.129 & .182  
MBP, mmHg & 1.015 & 1.002-1.029 & .027  
HDL, mmol/L & 1.566 & .796-3.080 & .194  
WBC, x10^9/L & .930 & .857-1.010 & .086  
Neutrophil, x10^9/L & .941 & .862-1.029 & .182  
RBC, ×10^12/L & .714 & .512-995 & .047  
HB, g/L & 1.566 & .796-3.080 & .194  
P, mmol/L & 8473.940 & 1522.947-47150.476 & .000  
Ca, mmol/L & 5.190 & 1.025-26.290 & .047  
Adjusted Ca*, mmol/L & 3.118 & .514-18.928 & .217  
Ca×P, (mg/dL) & 2 & 1.335 & 1.262-1.412 & .000  
Adjusted Ca×P, (mg/dL) & 2 & 1.345 & 1.269-1.425 & .000  

Factors with p value<0.2 are listed.

Table 4. Multivariate analysis for factors associated with lacunes by Multinominal Regression.

| Factors       | Model 2 | Model 3 |
|---------------|---------|---------|
|               | P-value | OR      | 95%CI     | P-value | OR      | 95%CI     |
| Scattered lacunes |        |         |          |        |         |          |
| Age | .000 | 1.051 | 1.027-1.075 | Age | .000 | 1.050 | 1.027-1.074 |
| Hypertension | .024 | 2.281 | 1.116-4.662 | Hypertension | .018 | 2.350 | 1.159-4.766 |
| Diabetes | .128 | 1.497 | .891-2.517 | Diabetes | .121 | 1.501 | .898-2.510 |
| MBP | .161 | 1.014 | .955-1.033 | MBP | .129 | 1.015 | .996-1.034 |
| WBC | .036 | .883 | .786-992 | WBC | .020 | .874 | .780-979 |
| BUN | .100 | 1.118 | .979-1.276 | BUN | .084 | 1.121 | .985-1.275 |
| P | .000 | 161.836 | 22.407-1168.871 | Ca×P | .000 | 1.159 | 1.086-1.237 |
| Ca or adjusted Ca | .592 | .547 | .060-4.982 |    |    |    |
| Multiple lacunes |        |         |          |        |         |          |
| Age | .000 | 1.078 | 1.043-1.113 | Age | .000 | 1.081 | 1.047-1.115 |
| Hypertension | .071 | 2.547 | .922-7.052 | Hypertension | .036 | 2.894 | 1.070-7.824 |
| Diabetes | .685 | 1.153 | .577-2.298 | Diabetes | .588 | 1.204 | .616-2.352 |
| MBP | .017 | 1.031 | 1.005-1.057 | MBP | .013 | 1.031 | .997-1.075 |
| WBC | .602 | .962 | .832-1.112 | WBC | .413 | .945 | .824-1.083 |
| BUN | .019 | 1.210 | 1.031-1.420 | BUN | .016 | 1.213 | 1.036-1.420 |
| P | .000 | 945018125.529 | 36026472.282- | Ca×P | .000 | 1.159 | 1.086-1.237 |
| Ca or adjusted Ca | .768 | 1.593 | .071-34.941 |    |    |    |

Factors with P value<0.05 entered into models.

Table 5. Multivariate analysis for factors associated with WMHs by Binary Regression.

| Factors       | Model 2 | Model 3 |
|---------------|---------|---------|
|               | P-value | OR      | 95%CI     | P-value | OR      | 95%CI     |
| Age | .000 | 1.122 | 1.071-1.176 | Age | .000 | 1.118 | 1.068-1.171 |
| Hypertension | NE | 3.090 | 1.934-4.937 | Hypertension | .039 | 3.767 | 1.068-13.284 |
| Diabetes | NE | 1.769 | 1.168-2.679 | Diabetes | NE | 1.002 | 1.029-1.029 |
| MBP | NE | 3.090 | 1.934-4.937 | MBP | NE | 1.002 | 1.029-1.029 |
| WBC | NE | 1.769 | 1.168-2.679 | WBC | NE | 1.002 | 1.029-1.029 |
| BUN | .036 | 1.256 | 1.014-1.555 | BUN | .023 | 1.298 | 1.036-1.625 |
| P | .000 | 6965.965 | 576.828-84123.332 | Ca×P | .000 | 1.338 | 1.231-1.454 |

NE: not entry. Method: Bachward LR. N=100 in WMHs.
Table 6. Multivariate analysis for factors associated with CMBs by Binary Regression.

|                | Model 2 |          |          | Model 3 |          |          |
|----------------|---------|----------|----------|---------|----------|----------|
|                | P-value | OR       | 95%CI    | P-value | OR       | 95%CI    |
| Age            | .004    | .057     | 1.017-1.097 | .001    | .068     | 1.027-1.121 |
| Hypertension   | NE      |          |          | NE      |          |          |
| Diabetes       | NE      |          |          | NE      |          |          |
| MBP            | NE      |          |          | NE      |          |          |
| WBC            | NE      |          |          | NE      |          |          |
| BUN            | NE      |          |          | NE      |          |          |
| P              | .000    | 558.429  | 49.395-6313.236 | .000    | 1.213    | 1.121-1.312 |
| Ca or adjusted Ca | NE      |          |          | NE      |          |          |

NE: not entry. Method: Bachward LR. N=55 in CMBs.

Table 7. H-L test of model 2 and model 3 in subtypes of CSVD.

|                  | Scattered lacunes | Multiple lacunes | WMHs | CMBs |
|------------------|-------------------|------------------|------|------|
| Model 2          | 0.858             | 0.404            | 0.693| 0.541|
| Model 3          | 0.571             | 0.404            | 0.295| 0.560|

Models with p value>0.1 means good calibration power.

Figures

1102 patients with stroke and cranial MR

Excluded:
- No serum calcium and phosphorus test: 501
- Parathyroid disease: 1
- Pituitary disorders: 1
- Bone metastasis of malignant tumor: 2
- Chronic kidney disease with blood creatinine level>186μmol/L: 9

388 patients included in our study:
- No CSVD: 140; CSVD: 448

Univariate and multivariate analysis for factors associated with CSVD by Regression; establish and verify predictive models for CSVD;

Subgroup CSVD

Lacunes (scattered lacunes: 212; multiple lacunes: 236), WMHs (100), CMBs (55)

Verify models in lacunes, WMHs, CMBs

Figure 1
Flow diagram showing the patient selection and data analysis.

Figure 2
Propensity score of included cases and excluded cases, from demographic and clinical characteristics and routine serum test. Including age, gender, hypertension, diabetes, coronary artery disease, atrial fibrillation, hyperlipidemia, history of smoking and drinking alcohol, weight, NIHSS score at admission and discharge, MBP, LVEF, FBG, P2hPG, HbA1c%, TC, LDL, HDL, cell count of WBC, neutrophil, lymphocyte, RBC, and PLT; albumin, PT, INR, APTT, FIB, BUN, Cr, HCY, TSH, FT3, FT4.

Figure 3
A Percentage of single or combined subtypes in CSVD. B-D Serum P, Ca, Ca×P of subtypes in CSVD, including scattered lacunes, multiple lacunes, WMHs, and CMBs. Brown dotted lines is the upper and lower limits of normal values.
Figure 4

TOAST classification distribution of no CSVD and CSVD groups. TOAST classification: 1. Large artery atherosclerosis. 2. Cardioembolism. 3. Small-artery occlusion. 4. Stroke of other determined cause. 5. Stroke of undermined cause.

Figure 5

Serum P, Ca, Ca×P concentration of no CSVD and CSVD. Blue dotted lines is the upper and lower limits of normal values.
Figure 6

A AUCs of model 1-4. B H-L test of model 1-4.
Figure 7

Nomograms of model 2 and model 3 for predicting the probability of CSVD. A model 2; B model 3

Figure 8

ROCs and AUCs of model2 and model3 in subtypes of CSVD.