Effect of AK098656 on association of hypertension with coronary stenosis severity in coronary heart disease patients

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Abstract

**Background:** AK098656 may be an adverse factor for coronary heart disease (CHD), especially in patients with hypertension. This study aimed to analyze the effect of AK098656 on CHD and CHD with various complications.

**Methods:** A total of 117 CHD patients and 27 healthy control subjects were enrolled in the study. Plasma AK098656 expression was determined using the quantitative real-time polymerase chain reaction. Student’s *t*-test was used to compare AK098656 expression levels in different groups. Receiver operating characteristic (ROC) curve and area under the curve (AUC) were used to quantify the discrimination ability between CHD patients and health controls and between CHD and CHD+complications patients. The relationship between AK098656 and coronary stenosis was analyzed using Spearman's correlation.

**Results:** AK098656 expression was remarkably higher in CHD patients than in healthy controls (*P*=0.03). The ROC curve revealed an effective predictive AK098656 expression value for CHD risk, with an AUC of 0.656 (95% CI: 0.501–0.809). Moreover, AK098656 expression was increased in CHD+complications patients compared to CHD patients alone (*P*=0.005), especially in patients with hypertension (CHD+HHTN, *P*=0.030). The ROC curve revealed a predictive AK098656 prognostic value for discriminating between CHD and CHD+HHTN patients, with an AUC of 0.666 (95% CI: 0.528–0.805). There was no significant difference in AK098656 expression in CHD patients with diabetes mellitus compared to CHD patients alone. In addition, AK098656 expression in CHD patients was positively correlated with stenosis severity (*R*=0.261, *P*=0.006).

**Conclusion:** AK098656 expression was significantly increased in patients with CHD, especially those with hypertension, and its expression level was positively correlated with the degree of coronary stenosis. This implied that AK098656 may be a risk factor for CHD and can potentially be applied in clinical diagnosis or provide a novel target for treatment.

Introduction

Cardiovascular disease is the leading cause of death worldwide. Coronary atherosclerotic heart disease (CHD) is the most harmful disease and leading cause of hospitalization and death among adult heart disease patients (1–3). Multiple studies have shown that controlling the risk factors can significantly reduce CHD morbidity and mortality (4–8). Many traditional risk factors, such as family history of CHD, hypertension, diabetes, obesity, and smoking, already have been targeted for early interventions in clinical work (9,10). However, current research has shown that some nontraditional factors, such as autoimmune diseases, high sensitivity C-reactive protein, and homocysteine, also play important roles in the occurrence and development of CHD and need to be given sufficient attention.

Long non-coding (Inc) RNA has a length of more than 200 nucleotides and limited protein-coding ability and can be considered as “transcriptional junk” (11). However, growing evidence has suggested that IncRNA has important regulatory functions and is associated with the development of complex diseases,
such as cardiovascular diseases and cancer (12–15). Accumulating studies have revealed that multiple lncRNAs are implicated in CHD. An earlier study has shown that lncRNA-p21 enhances p53 activity via the p53 signaling pathway, thereby inhibiting the formation of atherosclerosis (16). CoroMarker is very stable in the plasma of CHD patients and may become a new plasma biomarker for CHD diagnosis (17). H19 is re-expressed in atherosclerotic patients, and its polymorphism is related to the risk of CHD (18).

A new human vascular smooth muscle cell (VSMC)-dominant lncRNA AK098656 can accelerate the degradation of contractile proteins, increase the synthesis phenotype in VSMCs, and eventually lead to narrowing of resistant arteries and hypertension (19). VSMS phenotype conversion has crucial pathophysiological significance in different stages of atherosclerosis, while lncRNAs are important factors that regulate VSMC function and phenotype, as well as occurrence and development of related diseases. However, because AK098656 is a newly discovered lncRNA specifically expressed in VSMCs, its relationship with CHD has not been reported.

The present study found that AK098656 expression in CHD patients was significantly higher than that in healthy controls. To further analyze the effect of AK098656 on CHD complications, AK098656 expression levels in groups with different CHD complications, including CHD with diabetes mellitus (CHD + HDM), CHD with hypertension (CHD + HHTN), and CHD + HDM + HHTN, were analyzed. The results showed that AK098656 expression in CHD + HHTN and CHD + HDM + HHTN patients was significantly increased compared to CHD alone patients. AK098656 expression was also positively correlated with the degree of coronary artery stenosis. Thus, this study revealed a novel molecule that regulates CHD development.

Materials And Methods

Patients and samples

Due to the limited data on AK098656 expression in CHD patients, plasma and clinical data for healthy controls (n = 27) and CHD patients (CHD: n = 25, CHD + HDM: n = 24, CHD + HHTN: n = 41, and CHD + HDM + HHTN: n = 27) were collected in Beijing Luhe Hospital, Capital Medical University (Beijing, China) between September 2019 and January 2020. All participants were Han individuals from the same region. Patient inclusion criteria were as follows: 1) patients diagnosed with CHD. CHD was defined by coronary artery stenosis of ≥ 50% as determined by coronary angiography. Hypertension was defined by systolic blood pressure of at least 140 mm Hg or diastolic blood pressure of at least 90 mm Hg. Diabetes was defined by fasting blood glucose level of at least 7.0 mmol/L or blood glucose level of 11.0 mmol/L or more 2 h after an oral glucose load of 75 g; and 2) age of 35–80 years. Exclusion criteria included other clinically detectable cardiovascular diseases. The study complied with the Helsinki Declaration for investigation of human subjects and obtained ethical approval from the competent Institutional Review Boards of Beijing Luhe Hospital, Capital Medical University. All participants provided written informed consent.

RNA isolation and qPCR
Total RNA from human plasma was isolated using the TRIzol LS (Invitrogen, USA) and converted to cDNA using a cDNA synthesis kit (BioRad, USA) according to manufacturer instructions. The expression levels of lncRNA-AK098656 were determined using SYBR (BioRad, USA) and normalized using 18 s cDNA. The primer sequences were as follows: AK098656, forward: 5'-CCTCATTTGCTGGCACCTG-3', reverse: 5'-GGAGGCAAGGTAAGGGGA-3'; 18S, forward, 5'-GTAACCCGTTGAACCCATT-3', and reverse: 5'-CCATCCAATCGGTAGTGCG-3'.

**Biochemical Measurements**

Blood samples were collected for triglyceride, cholesterol, and high- and low-density lipoprotein cholesterol analysis using a Hitachi 7600-020 clinical analyzer (Hitachi, Tokyo, Japan).

**Coronary Angiography**

Coronary angiography methods have been previously described (20). Patients with coronary artery stenosis of ≥ 50% were diagnosed with CHD. CHD severity was further divided into three groups: mild, moderate, and severe. Mild refers to the degree of stenosis < 70% for all coronary arteries, moderate refers to the degree of stenosis ≥ 70% for one coronary artery, and severe refers to the degree of stenosis ≥ 70% for two or more coronary arteries.

**Statistical analysis**

To explore the relationship between AK098656 and CHD, Student’s t-test between healthy control and CHD groups was used. Furthermore, receiver operating characteristic (ROC) curve was utilized to estimate the discrimination ability of AK098656 in CHD and to quantify the discrimination ability using an area under the curve (AUC), which is widely used in pattern recognition. The distribution of AUC values was between 0 and 1. AUC > 0.5 meant that the indicator had the ability to distinguish between conditions in question. The larger the AUC value, the stronger the ability to distinguish.

Student’s t-test was then used between the CHD and CHD + complications groups (CHD + HDM, CHD + HHTN, and CHD + HDM + HHTN, n = 92) to explore the relationship between AK098656 and complications with CHD. The ROC curve and AUC were used to quantify the discrimination ability between CHD and CHD + complications. In addition, Student’s t-test was utilized between the CHD and CHD + HDM groups and with the CHD + HHTN group separately to explore which complication would result in a greater effect on AK098656. The ROC curve and AUC were also used to quantify the differences.

Hypertension is one of the most potent risk factors for CHD development, which increases the risk of stenosis in CHD patients. A recent study has demonstrated that AK098656 is a novel risk factor for hypertension (19). Therefore, correlation between AK098656 and CHD stenosis was evaluated using Spearman’s correlation analysis.

**Results**
General characterization of study subjects

Table 1 provides the general characteristics for participants with or without CHD and CHD accompanied by complications. The study included 27 non-CHD participants with an average age of 49 years (48.1% men). For CHD patients (65% men), the average age was 64 years. Compared to healthy controls, CHD and CHD + complications patients had a higher heart rate and increased systolic blood pressure. CHD patients diagnosed with diabetes had a higher fasting glucose level.

![Image](image.png)

| Characters | Healthy controls (n=27) | CHD (n=25) | CHD+HDM (n=24) | CHD+HHTN (n=41) | CHD+HDM+HHTN (n=27) |
|-----------|-------------------------|------------|----------------|----------------|---------------------|
| Age, years | 48.9±14.9               | 62.2±13.8  | 68.0±10.4      | 62.4±11.4      | 68.3±11.3           |
| Gender, M/F | 13/14                  | 13/8       | 13/11          | 29/12          | 15/12               |
| Heart rate, beats/min | 60.7±41.9              | 73.8±15.6  | 81.9±17.2      | 73.8±17.4      | 81.9±21.0           |
| Hypertension history | no                     | no         | no             | yes            | yes                 |
| Drug use | Aspirin, n(%) | 22(88%)    | 24(100%)       | 3(80.5%)       | 21(77.8%)           |
| Clotidogrel, (%) | 0                      | 9(36%)     | 15(62.5%)      | 18(43.9%)      | 15(55.6%)           |
| Statin, n(%) | 0                      | 17(68%)    | 23(95.8%)       | 35(85.4%)     | 20(74.1%)           |
| Calcium antagonist, (%) | 4(14.8%) | 2(8%)       | 2(8.3%)         | 17(41.5%)     | 1(48.1%)           |
| β-blocker, n(%) | 3(11.1%) | 13(52%)    | 16(66.7%)       | 24(58.5%)     | 12(44.4%)           |
| Total cholesterol, mmol/L (SCHO) | 4.48±1.88 | 4.78±1.21  | 4.64±1.09      | 4.21±1.12      | 3.93±0.89           |
| Total trigly (STG) | 1.19±0.71 | 1.62±0.87  | 1.94±1.28       | 1.50±0.80      | 1.65±0.97           |
| HDL cholesterol, mmol/L (SHDL) | 1.37±0.39 | 1.14±0.25  | 1.06±0.24       | 1.09±0.23      | 1.12±0.24           |
| LDL cholesterol, mmol/L (SLDL) | 2.67±0.67 | 3.09±0.84  | 3.15±0.83       | 2.73±0.80      | 4.24±0.74           |
| Fasting glucose, mmol/L (SFGB) | 5.49±1.20 | 5.55±0.99  | 8.54±3.37      | 5.77±1.37      | 8.98±2.99           |
| Coronary stenosis | Mild - | 4 | 2 | 4 | 3 |
| | Moderate - | 7 | 4 | 14 | 5 |
| | Severe - | 8 | 8 | 12 | 8 |
| | Not available - | 6 | 10 | 11 | 11 |

Demographics and clinical characteristics. Data was presented as mean value ± standard deviation or count (%). CHD, coronary heart disease; HDM, diabetes mellitus; HHTN, hypertension.

AK098656 expression correlation for CHD risk

AK098656 expression level was significantly higher in CHD patients compared to healthy controls (P = 0.038; Fig. 1A). In order to quantify the discrimination ability between healthy controls and CHD patients, ROC curve analysis revealed that AK098656 was a valuable predictor for CHD risk (AUC = 0.656, 95% CI: 0.501–0.809; Fig. 1B).

Independent predictive value for AK098656 expression for CHD complication risk

Because CHD is often accompanied by other chronic complications, such as hypertension and diabetes, AK098656 expression level in CHD and CHD + complications patients was further analyzed. The AK098656 expression level in CHD + complications patients was significantly higher than that in CHD alone patients (P = 0.005; Fig. 2A). The ROC curve analysis revealed that AK098656 had the ability to distinguish between the CHD and CHD + complications groups (AUC = 0.673, 95% CI: 0.552–0.793; Fig. 2B). Furthermore, the AK098656 expression level in different CHD complications (HDM, HHTN, and
HDM + HHTN) was analyzed. Expression in CHD + HHTN patients was significantly higher than that in CHD alone patients (P = 0.030). In CHD patients with diabetes mellitus, AK098656 expression was slightly higher than that in CHD patients, although no statistical difference was present (P = 0.365). In addition, there was no significant difference in AK098656 expression level between CHD + HDM and CHD + HHTN patients (P = 0.135; Fig. 2C). The ROC analysis implied that AK098656 had the potential to distinguish between the CHD and CHD + HHTN groups (AUC = 0.666, 95% CI: 0.528–0.805; Fig. 2D).

**AK098656 increases coronary stenosis risk in CHD patients.**

CHD severity was divided into three degrees: mild (stenosis degree for all coronary arteries was < 70%), moderate (stenosis degree of one coronary artery was ≥ 70%), and severe (the stenosis degree of two or more coronary arteries was ≥ 70%; Fig. 3A–C). It is worth noting that AK098656 expression was gradually upregulated with the increase in stenosis degree (Fig. 3D) and was positively associated with stenosis severity (R = 0.261, P = 0.006). It is possible that a higher AK098656 expression may indicate a danger of coronary stenosis.

**Discussion**

In this study, human-specific lncRNA AK098656 was significantly increased in CHD patients, especially in those with hypertension. In addition, AK098656 was positively correlated with coronary stenosis severity in CHD patients. This study not only suggested that AK098656 could predict CHD severity, but also proposed that elevated AK098656 in hypertensive individuals may enhance the risk of CHD. The results also confirmed a previous report that AK098656 is upregulated in hypertensive patients compared to normotensive controls (19). A great deal of research shows that diabetes has a significant effect on coronary arteries and cardiac microcirculation. Both insulin resistance and hyperglycemia inflict significant damage to the cardiovascular system (21, 22). Thus, AK098656 expression was further investigated in CHD + HDM patients. However, AK098656 expression in the CHD + HDM group was not greater than that in the CHD group. An explanation for this observation cannot be suggested based on the current experiment data. However, the insignificant difference between the CHD + HDM and CHD groups may be due to the study's small sample size.

Differentiation and phenotypic transformation of VSMCs are key initial steps in cardiovascular diseases, such as hypertension, atherosclerosis, and restenosis after angioplasty (23). Many studies have confirmed that under the influence of different external factors, VSMCs show different phenotypic characteristics. Thus, transformation between different VSMC phenotypes plays an important role in the occurrence and development of a variety of vascular diseases (24). Therefore, it is essential to investigate the molecular mechanisms of VSMC differentiation and phenotypic transformation for prevention and treatment of these diseases. Differentiated and mature VSMCs mainly maintain their vascular morphology, while dedifferentiated “secretory” VSMCs secrete a large amount of extracellular matrix (ECM) through migration and proliferation. They also play an important role in the occurrence and development of a variety of vascular diseases, including CHD [24–25]. Angiotensin II and epidermal growth factor are widely recognized to induce proliferation and migration of mature VSMCs [26, 27].
Insulin-like growth factor-1 (IGF-1) is one of the most intensively studied factors that can inhibit phenotypic VSMC transformation. IGF-1 activates the downstream PI3K-AKT signaling pathway by binding to the IGF receptor, thereby increasing the expression of VSMC systolic protein (28). In addition, tumor growth factor, rapamycin, and microRNA (miR-221, -222, -21, -146A, and −133) are all involved in the regulation of cell proliferation, apoptosis and migration (29, 30). According to Jin et al. study, transgenic AK098656 rats spontaneously developed hypertensive phenotypes accompanied by reduced lumen diameter and enhanced media/lumen ratio (). Then, AK098656 directly bound to MYH11, which is a smooth muscle cell-specific contractile protein. Fibronectin is another AK098656 binding protein that is an essential ECM component (19). Both MYH11 and fibronectin promote VSMC plasticity and synthetic phenotype in hypertension pathogenesis (31, 32). To some extent, the above evidence may explain the mechanism of increased AK098656 expression aggravating coronary artery stenosis, which eventually affects the stenosis score in CHD patients.

Increasing evidence has demonstrated that IncRNAs are not “transcriptional junk”. In fact, IncRNAs play various regulatory roles in each step of gene expression and translation via epigenetic regulation, alternative splicing regulation, or acting as a molecular sponge (33). A large number of IncRNAs may be part of a complex regulatory network in physiopathology in the cardiovascular system. Consequently, IncRNAs may open up new avenues for diagnosis or serve as therapeutic targets for CHD treatment.

There were some limitations in this study. (1) A total of 144 subjects were analyzed in this study, resulting in a relatively small sample size, which may have resulted in insufficient statistical power. (2) Because the clinical data and samples were all collected from a general population at a single center, selection bias may be present due to regional restrictions. Therefore, the results should be further confirmed via large, multi-center, prospective clinical trials. (3) Detailed IncRNA-AK098656 mechanisms in CHD or CHD + complications patients were not investigated.

In conclusion, AK098656 may be a risk factor for CHD and can potentially be applied in clinical diagnosis and provide a novel target for treatment.

Declarations

Acknowledgments

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Availability of data and material

All data generated or analyzed during this study are included in this published article.

Author contributions

XW, YLW, and DZ designed the experiments. YLW, YYZ, BYZ, and YM were responsible for sample collection and database management. XW performed all RNA isolations and qPCR testing, as well as
data analysis. XW and YLW wrote the manuscript. DZ, LYY, and JK supervised the experiments and revised the manuscript.

**Conflict of interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**Figures**
Figure 1

Diagnostic value of AK098656 for CHD. (a) AK098656 expression level in healthy control and CHD groups. (b) Discrimination ROC for healthy control and CHD groups.
Figure 2
Predictive AK098656 value for CHD patients with complications. (a) AK098656 expression level boxplot for the CHD and CHD+complications groups. (b) Discrimination ROC for the CHD and CHD+complications groups. (c) An AK098656 expression boxplot for the CHD, CHD+HDM, and CHD+HHTN groups. (d) Discrimination ROC for the CHD and CHD+HHTN groups.
Figure 3

Correlation between AK098656 expression and coronary stenosis. Coronary angiography for (a) normal, (b) mild coronary stenosis, (c) and severe coronary stenosis. (d) The AK098656 expression level in different stenosis degrees (normal, n=27; mild+moderate, n=43; and severe, n=36).