Associations of Variants within the PHACTR1, WDR12 and ANRIL Genes with Coronary Artery Disease Severity

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Abstract

Objective: To conduct a stage I study to identify genetic predictors of coronary artery disease severity and develop genomic profiles for accurate classification of high-risk patients in a clinical cohort.

Methods: 719 patients who underwent coronary angiography for myocardial infarction or suspected coronary artery disease were divided into cases with severe disease and controls with non-severe and no disease. Coronary artery disease severity was scored on based on: i) severity of coronary artery luminal stenosis and ii) number of diseased main coronary vessels. A case-control design was employed. We directly assessed for SNP association with coronary artery disease severity as determined by invasive coronary angiography.

Results: A systematic statistical analysis strategy was used to assess the association of all genetic factors with CAD severity. Results of this analysis revealed two SNPs associated with severity of coronary artery stenosis and 3 SNPs associated with number of diseased proximal vessels. Logistic regression models were generated for each SNP factoring in related covariates – age, sex, dyslipidemia and hypertension status.

(rs12526453) in the PHACTR1 gene on chromosome 6, and (rs6725887) in the WDR12 gene on chromosome 2 were predictive of the severity of coronary artery disease luminal stenosis after adjustment for cofactors (OR=1.38 p=0.02 and OR=1.46 p=0.048 respectively). (rs6725887) was again associated with CAD severity as determined by the number of diseased proximal vessels (OR=2.01, p=0.004). Two further SNPs were also associated with CAD severity as defined by the number of diseased proximal vessels: (rs4977574) in proximity to CDK2A & CDK2B on chromosome 9p21.3 (OR=1.64 p=.005); and (rs10953541) in the BCAP29 gene on chromosome 7 (OR=1.814, p=0.004).

Conclusion: We have identified four SNPs to be predictive of CAD severity as assessed by coronary angiography. Two SNPs are predictive of the severity of proximal coronary artery disease based on the extent of luminal stenosis: (rs12526453) in the PHACTR1 gene on chromosome 6 and (rs6725887) in the WDR12 gene on chromosome 2.

(rs6725887) and a further two SNPs are predictive of triple vessel disease: (rs4977574) in the ANRIL gene in proximity to CDK2A & CCK2B on chromosome 9p21.3 and (rs10953541) in the BCAP29 gene on chromosome 7.

This discovery may pave the way for the development of a genetic diagnostic and screening tool for coronary atherosclerosis. It may also help identify targets for future gene therapeutics against atherosclerotic coronary disease.

Introduction

Despite the broad application of proven preventive strategies, coronary artery disease (CAD) remains the leading cause of mortality worldwide. In 2005 it killed more than 7.5 million people [1], and deaths from CAD continue to rise globally [2]. In clinical cardiology, invasive coronary angiography remains the gold standard for establishing the diagnosis and defining the severity of coronary artery disease [3]. Here we report the identification of four SNPs to be associated with the severity of CAD as assessed by coronary angiography in an Australian Caucasian population:
remedy of coronary artery disease (CAD). It is investigating the genetic basis of CAD traits. Several large-scale GWAS of CAD patients [12] with 22 SNPs in 19 database (www.genome.gov/26525384) has catalogued results from chromosome 9p21.3) associated with CAD [11]. Recently, the GWAS identified the first notable genetic variant (located on chromosome 9p21.3 and (rs10953541) in the BCAP29 gene on chromosome 7. Distinct morphological characteristics of CAD show different degrees of heritability. Proximal coronary artery disease has been shown to display higher degree of heritability in comparison to distal disease [14]. Hence our focus was on the degree of stenosis and the plaque distribution in the four major epicardial coronary arteries: left main, left anterior descending, left circumflex and right coronary arteries.

Our research is unique in that it goes beyond the association of SNPs with coronary artery disease traits, directly assessing for SNP association with CAD severity as assessed by invasive coronary angiography. Two SNPs are predictive of the severity of proximal coronary artery disease luminal stenosis: (rs12526453) in the PHACTR1 gene on chromosome 6 and (rs6725887) in the WDR12 gene on chromosome 2. (rs6725887) and a further two SNPs are predictive of triple vessel disease: (rs4977574) in the ARNII gene in proximity to CDK2A & CK2B on chromosome 9p21.3 and (rs10953541) in the BCAP29 gene on chromosome 7.

This discovery furthers the potential of developing a genetic test for screening for the risk of severe and diffuse coronary atherosclerosis permitting early detection and effective prevention by the aggressive control of modifiable risk factors.

Methods

Study population

This study was conducted collaboratively between the cardiology department of the Gold Coast Hospital, Queensland and The Genomics Research Centre (GRC) at Griffith University, Queensland Australia. The study was approved by the institutional ethics committee and all subjects gave informed written consent. 719 Caucasian subjects who underwent coronary angiography between 2006 and 2012 were selected. The study population was homogeneous consisting only white Caucasians.

Coronary angiography

The indications for coronary angiography included acute coronary syndrome (diagnosed by clinical history, electrocardiographic findings and elevated cardiac troponin [15]), pre-surgical cardiac assessment, work-up for valve surgery and stable angina. The full spectrum of clinically manifested coronary artery disease was represented in this population as follows: ST elevation myocardial infarction (STEMI) (5.7%) and Non-ST elevation myocardial infarction (NSTEMI) (26.0%); Unstable Angina (5.0%); Stable Angina (21.1%); and other reasons (42.1%) including positive cardiac stress testing, positive functional imaging and pre-operative assessment. Coronary angiography was carried out using a Siemens Axiom Artis image intensifier (Siemens inc. Sweden) according to the Judkins’ technique using standard 6 French gauge right and left coronary catheters with the images acquired at a frame count of 15 per second and the orthogonal projections were made according to a standard institutional protocol and images of the four main coronary arteries: the left main coronary artery (LMCA), the left anterior descending artery (LAD), the left circumflex artery (LCx) and the right coronary artery (RCA) were acquired. Two independent workers analysed and reported the findings of angiography on two separate occasions for cross-validation. The lesions in the main coronary arteries were recorded as a percentage of the stenosis of the vessel (determined by visual estimation of the diameter of the narrowing compared to the proximal normal vessel). The number of diseased main vessels and the number of lesions in each main vessel were recorded. The reader reporters of the angiograms were blinded to the genotyping results of the patients.
In our study the coronary artery disease severity was measured utilising two steps: In the first step, the vessel score, was defined as the number of diseased main vessels with >50% luminal diameter stenosis. A point was allocated to each diseased vessel: single vessel disease (1 point), two vessel disease (2 points) or triple vessel disease (3 points). Stenosis of the left main coronary artery (LMCA) was considered more severe and was attributed a score of 2 points (LMCA divides into two main branches the left anterior descending and the left circumflex). In the second step the scoring was based on the number of lesions in each vessel. Patient’s were dichotomised into those with no or mild disease (those with less than 30% diameter stenosis) and severe disease (one or more lesion of 50% or more diameter luminal stenosis in a main coronary artery was defined as severe disease). This approach aimed at identifying genetic markers associated with the morphological characteristics of CAD which have clinical significance and which demonstrate the highest inheritability. Therefore by focusing our scoring on proximal disease we intend to enhance our prospect of identifying genetic predictors of inheritability and severity.

Risk factors

All clinical data was obtained during the index admission. Each patient had the standard coronary risk factors recorded. 62.7% of the study population were male, 50.5% were hypertensive, 57.8% had hypercholesterolaemia, 21.3% had diabetes, 5.3% were current smokers and 20.6% of patients had a positive family history (CAD in male first-degree relative, or father less than 55, or female first-degree relative or mother less than 65).

Genotyping

5ml EDTA blood samples were collected from femoral artery or the radial artery at the time of arterial puncture for vascular access. Genomic DNA was extracted and purified from peripheral blood lymphocytes using a standard salting-out procedure [16]. 22 previously identified SNPs [17] were genotyped in the 719 patients.  SNPs were selected from a search of the GWAS database for SNPs associated with CAD in 2011. Genotypes for each SNP was determined by restriction-fragment 140 length polymorphism (RFLP) analysis of restriction enzyme-digested PCR products on agarose gels. A 20-μl PCR reaction mix contained 1× PCR buffer, 1.75 mM MgCl2, 0.2 mM dNTPs, 0.15 “M of each primer, 20–40 ng of genomic DNA and 1.5 U of GoTaq® (Promega). Thermocycler conditions were an initial denaturation at 95°C for 10 min, followed by 35 cycles of 95°C for 45 s, 60°C for 45 s, 72 °C for 45 s and a final extension step of 72 °C for 7 min. Specific primers were used to amplify each region containing the SNPs and restriction enzymes (NEB) chosen that would differentiate the two alleles after separation on 3% agarose gels.

Statistical analysis

For the primary exploratory analysis, a bivariate correlation analysis of all SNPs was performed against severity (as determined by the number of vessels involved and the severity of stenosis) using Spearman’s statistic [18]. 2 SNPs were shown to be associated with severity of stenosis: (rs12526453, OR=1.42 p = 0.01) and (rs6725887, OR=1.51 p = 0.031). Four SNPs were shown to be associated with triple vessel disease in the primary exploratory analysis (rs10953541, OR=1.77 p=0.005), (rs12526453 OR=1.52, p=0.014), (rs4977574, OR=1.66 p=0.003) and (rs6725887 OR=2.05 P=0.003).

For the analysis of the severity of luminal stenosis, the severity was dichotomised into 0 (Normal, which is <30% stenosis) and 1 (significant disease, >50% stenosis in 1 or more proximal vessel). A Logistic regression model was generated for the two SNPs identified in the severity score primary analysis. Age, sex, family history, dylipidaemia, hypertension, smoking and diabetes were factored in as co-variates.

Results

Results of this analysis showed (rs12526453) in the PHACTR1 gene on chromosome 6, and (rs6725887) in the WD repeat domain 12 (WDR12) on chromosome 2 to be predictive for the severity of coronary artery disease (as determined by the severity of luminal stenosis) after adjustment for cofactors (OR=1.38 p=0.02 and OR=1.46 p= 0.048 respectively).

For the analysis of the number of vessels involved, the number of vessels was dichotomised into 0 (no vessels, n=226) and 1 (2 or 3 vessels, n=493). A Logistic regression model was generated for the three SNPs identified in the vessel score primary analysis. Again, age, sex, family history, dylipidaemia, hypertension, smoking and diabetes were factored in as co-variates.

Results of this analysis again showed (rs6725887) in the WDR12 gene on chromosome 2 to be associated with CAD severity as defined by the number of vessels involved (OR=2.01, p=0.004). Two further SNPs were also associated with CAD severity as defined by the vessel score: (rs4977574) in proximity to CDK2A & CDK2B genes on chromosome 9p21.3 (OR=1.64 p=0.005); and (rs10953541) in the B-cell receptor-associated protein 29 (BCAP29) on chromosome 7 (OR=1.814, p=.004).

Discussion

Our study evaluated the association between 22 SNPs and CAD severity as assessed by the gold standard investigation, invasive coronary. Results of our analysis have shown four SNPs in total to be associated with disease severity.

One of the strengths of our study design is that we identify cases based on angiographic evidence of both severity and the most inheritable morphological characteristics of CAD. Other case-control comparison studies often have variability in study enrolment, in particular controls are chosen from healthy individuals in the general population whose coronary anatomy is not known (and therefore may well be harbouring sub-clinical atherosclerosis). Each of our controls has angiographic evidence of the percentage of luminal stenosis in their coronary arteries and our study design excludes...
individuals from the general population who may have CAD but are asymptomatic.

Variants of phosphatase and actin regulator 1 gene (PHACTR1) demonstrated association with early onset MI in a study published by the MI genetics consortium [19]. PHACTR1 has also been associated with severity of coronary stenosis in a Lebanese cohort (OR = 1.37). SNP (rs9349379) in their study was associated with the severity of atherosclerosis in the Lebanese study but not replicated in our study. Although rs12526453 did not show association in the Lebanese exploratory set it did show association in their replication set [20]. Multiple GWASs in the Chinese Han population have not shown rs12526453 and PHACTR1 to be associated with CAD or MI. In the Chinese studies the minor allele frequency (based on HapMap data) of rs12526453 was 0% and none of the 113 SNPs in or near PHACTR1 showed significant association (p <0.01) for coronary artery disease [21,22]. Although their study sizes were small in comparison to the European studies it suggests that PHACTR1 may not play a role in the Chinese population.

The pathophysiology of the PHACTR1 in coronary artery disease remains to be elucidated. PHACTR1 has been proposed to be a key regulator of endothelial cell function [23] and is a regulator protein of protein phosphatase 1 (PP1), an enzyme that regulates endothelial nitric oxide (eNO) [24]. eNO is an important modulator of coronary artery disease and has been shown to influence thrombogenicity and inflammation [25]. Mutations in the PHACTR1 may contribute to atherosclerotic plaque formation by leading to endothelial dysfunction. Our finding that rs12526453 is associated with coronary artery disease severity but not associated with MI is suggestive that the role of PHACTR1 is in atherogenesis (plaque formation) and not plaque rupture or thrombosis.

Rs 4977574 has previously been shown to be associated with coronary artery disease severity in an Italian study [26] and also with MI [19,27]. Rs4977574 is located within the ANRIL gene that encodes a large non-coding RNA, its pathophysiology is unknown but its role in CAD has been implicated in several expression studies. The ANRIL promoter region contains binding sites for zinc-finger proteins that are critical for the transcription of CDKN2A and CDKN2B and in the mouse, deletion of the 9p21 orthologous non-coding region including the 3’ sites for zinc-finger proteins that are critical for the transcription of ANRIL a gene-centric approach to elucidating cardiovascular risk. Circ Cardiovasc Genet. 2009; 2: 3-6.

At least 11 chromosomal regions have been identified as associated with the risk of MI beyond that of traditional risk factors, with each risk allele increasing the risk of MI by a relatively small margin (10-30%) [28]. Myocardial infarction and coronary occlusion most frequently evolve from mild to moderate stenosis whilst severe atherosclerotic coronary artery disease is less likely to be the cause of MI [ENREF_9 8 11]. Severe disease is clinically significant due to the associated symptoms, morbidity, mortality and therapeutic options such as stent insertion or coronary artery bypass surgery.

Conventional risk factors do not explain the full extent of coronary artery disease pathophysiology. Our work shows the importance of using the phenotype information gained from coronary angiography to begin to identify the mechanism of action of the risk genes identified in large scale GWASs. Data from coronary angiography will also improve the development and accuracy of genetic tools for identifying individuals at risk of severe coronary artery disease. This could eventually enable earlier prevention and would be a leap forward in the personalisation of medicine based on one’s genetic risk. The identification of novel genetic links and elucidation of their pathophysiology will be essential for the development for new therapies for the prevention and treatment of coronary artery disease.

Conclusion

In conclusion, we have identified of four SNPs (rs12526453) in the PHACTR1 gene on chromosome 6, (rs6725887) in the WDR12 gene on chromosome 2, (rs4977574) in the ANRIL gene on chromosome 9p21.3 and (rs10953541) in the BCAP29 gene on chromosome 7 to be predictive of CAD severity as assessed by coronary angiography. This discovery may pave the way for the development of a genetic diagnostic and screening tool for coronary atherosclerosis. It may also help identify targets for future gene therapeutics against atherosclerotic coronary disease.

References


