Infra-renal abdominal aortic calcification volume does not predict small abdominal aortic aneurysm growth

Kerolos Hendya, Ronny Gunnarsson, Oliver Cronin, Jonathan Golledge

A R T I C L E   I N F O
Article history:
Received 2 April 2015
Received in revised form 10 July 2015
Accepted 14 July 2015
Available online 16 July 2015

Keywords:
Abdominal aortic aneurysm
Growth
Calcification

A B S T R A C T
Background: Vascular calcification is a common finding in abdominal aortic aneurysms (AAA) however whether it predicts aneurysm expansion is controversial.
Objectives: 1) To establish a reproducible method of assessing AAA calcification using computed tomography (CT); 2) To investigate the association between AAA calcification and growth.
Method: Patients were identified from a prospectively maintained small AAA surveillance database. To be included patients required at least two CT scans a minimum of 6 months apart. All patients had a maximal AAA diameter of ≤55 mm on their initial scan. Infra-renal aortic calcification volume, total infra-renal aortic volume and maximal AAA diameter were measured. Reproducibility was assessed from repeat scans performed on 31 patients. AAA growth, estimated by volume change per year, was compared between patients with baseline infra-renal aortic calcification volumes< and ≥median.
Results: 95% agreement limits (lower, upper) for intra and inter-observer error in measuring infra-renal calcification volume were 0.68, 97 mm³ and −140, 5.8 mm³, respectively. Concordance correlation coefficients for inter and intra-observer variability in measuring infra-renal aortic calcification volume were 0.99 and 0.98, respectively. Patients with infra-renal aortic calcification volume < median (n = 44) and ≥median (n = 44) had an infra-renal aortic volume increase of 6.0 cm³/yr and 7.8 cm³/yr, respectively (p = 0.66). Mean percentage infra-renal aortic volume increase/yr was found to be 4.2 ± 6.4 and 8.9 ± 6.2 for patients with and without diabetes, respectively (p = 0.003).
Conclusion: Infra-renal aortic calcification volume can be assessed reproducibly from CT images. Infra-renal aortic calcification volume did not predict small AAA growth.

Crown Copyright © 2015 Published by Elsevier Ireland Ltd. All rights reserved.

1. Introduction
Abdominal aortic aneurysm (AAA) is an important cause of mortality in older adults. There are currently no established drug therapies to limit AAA growth and surgical intervention does not reduce mortality of patients with small AAAs [1–3]. As a result, most patients with small AAAs undergo imaging surveillance until AAA diameter is ≥50–55 mm (mm). Despite such surveillance, 1–2% of small AAAs rupture per year [4,5]. Additional measures of selecting patients for intervention are needed [6].

AAA rupture represents a mechanical failure of the degenerated aortic wall, thus biomechanical considerations are important to understand this process and to improve our predictions of its occurrence [7]. Patient specific biomechanical profiling has been suggested as a potentially valuable tool in rupture risk assessment [8,9]. Additionally, peak wall stress (PWS) has been reported to be greater in ruptured and symptomatic AAAs compared to asymptomatic AAAs [10], and PWS has been reported to predict location of future rupture [9,11]. Furthermore, finite element analysis (FEA) studies have demonstrated that PWS is significantly greater in AAA regions with calcified plaque compared to regions with no
calciﬁcation [12]. Abdominal aortic calciﬁcation (AAC) score has also been reported to be higher in symptomatic and ruptured AAAs compared to asymptomatic AAAs [13]. A recent study evaluating AAA tissue specimen has implicated AAA calciﬁcation in determining rupture risk [14]. However, a multi-detector computed tomography study suggested that calcified plaques are smaller in AAA compared to normal abdominal aortas [15]. Thus the role of calciﬁcation in AAA pathogenesis has not been clearly elucidated.

A previous study reported that greater AAA calciﬁcation was associated with slower AAA expansion. In that study calciﬁcation was assessed semi-quantitatively by ultrasound (US) and inter-observer reproducibility was not reported. Furthermore analyses did not adjust for baseline AAA size [16]. A further small study which used computed tomography (CT) imaging to assess AAA calciﬁcation reported no association between AAA calciﬁcation volume and growth [17]. The nature of the relationship between infra-renal aortic calciﬁcation volume and small AAA growth is yet to be resolved. Studies investigating AAA calciﬁcation have employed variable methods of quantifying calciﬁcation and frequently not reported the assessment reproducibility [18].

The aims of the current study were:

1. To establish a valid and reproducible method of assessing infra-renal aortic calciﬁcation with CT;
2. To investigate the association between infra-renal aortic calciﬁcation volume and AAA growth rate.

2. Methods

2.1. Patients and clinical deﬁnitions

We performed a retrospective analysis of serial CT images of patients with small AAAs that were under surveillance at The Townsville Hospital (TTH) between June 2003 and November 2013. Such patients had previously consented to their medical information being conﬁdentially stored in a database and used for research purposes. A low, negligible risk ethics application for this study was approved by TTH Human Research and Ethics Committee (HREC) (approval number 13/QTHS/125) and endorsed by James Cook University HREC. To be included patients had to have undergone at least two CT scans a minimum of 6 months apart and images had to be available for retrospective analysis. All patients had a maximal axial AAA diameter of ≤55 mm on their initial scan. Repaired and mycotic AAAs were excluded. Patients with connective tissue diseases, such as Marfan’s syndrome, were excluded. Ischemic heart disease (IHD) was deﬁned by a history of myocardial infarction, angina pectoris or coronary revascularisation. Hypertension and diabetes mellitus (DM) were deﬁned by previous history or treatment for these conditions. Cigarette smoking classiﬁcation was based on smoking history and deﬁned as current smoker (smoked within the last month), ex-smoker (smoked previously but not in the last month) or never smoked.

2.2. CT acquisition

All CTs were performed at TTH using a 64 slice multi scanner (Philips, North Ryde, NSW), under a set acquisition protocol [19]. Abdominal aortic images were obtained in a single breath hold cycle in 3 mm slices at 3 mm intervals. 100 ml Ultravist 300 was delivered intravenously using an automated injection driver system. Image capture commenced when the locater set above the renal arteries detected that Hounsﬁeld units at the centre of the aorta reached 130.

2.3. Assessment of infra-renal aortic calciﬁcation

Original CT images were transferred onto a Philips MxView Visualisation Workstation software for detailed analysis. The region of interest (ROI) was the slice inferior to the origin of the lowest renal artery to the slice superior to the bifurcation of the abdominal aorta. Firstly, axial images throughout the ROI were scouted to carefully demarcate the outer boundary of the aortic wall. The outer aortic wall was traced free hand with the cursor under magniﬁed images to improve accuracy and to exclude adjacent bowel tissue, vertebral body tissue and all non-aortic tissue from the ROI. Secondly, coronal images were also scouted to identify the upper and lower boundaries of the ROI as described above. Calciﬁcation volume was subsequently extracted from the ROI using a workstation tool based on predetermined and validated thresholds of radio density for calciﬁcation [19] as illustrated in Fig. 1. Results were transcribed onto an excel spreadsheet for further analysis.

2.4. Assessment of infra-renal aortic volume and AAA diameters

Infra-renal aortic volume was assessed using methods similar to those used to measure infra-renal aortic calciﬁcation volume. After selecting the ROI and abdominal aortic boundary from axial slices, the total infra-renal aortic volume was estimated based on predefined and validated radio density set up on the Phillips workstation [19]. AAA orthogonal diameter was measured by ﬁrst scouting axial images within the ROI to identify the abdominal aortic centre point in each axial slice. Subsequently, images perpendicular to the AAA centreline were analysed to measure the maximal orthogonal diameter including the outer wall and thrombus. AAA growth was estimated from two CT scans, at least six months apart, for all patients. Growth was calculated by normalising the respective volume and diameter changes relative to the time interval between CTs for each patient and reporting as changes per year.

2.5. Reproducibility

In order to determine the repeatability of calciﬁcation measurements we assessed the agreement of measurements between and within observers on 31 patients. This involved two ﬁnal year medical students, each trained by an experienced doctor, measuring and recording the calciﬁcation volume from CTs whilst being blinded to each other’s results. Measurements were then repeated at least 24 h later by one observer (KH) to assess the intra-observer error.

2.6. Statistical analysis

Statistical analysis for reproducibility was performed according the methods outlined by Bland and colleagues [20]. Furthermore, concordance correlation coefﬁcient were calculated according to Lin [21]. To test our hypothesis that patients with infra-renal aortic calciﬁcation volume below median would experience a greater AAA volume increase compared to those with calciﬁcation volume equal to or above median, patients were divided into two groups. Group 1 has infra-renal aortic calciﬁcation volume less than median and group 2 has an infra-renal aortic calciﬁcation volume greater than or equal to median. The required sample size was calculated based on two assumptions. Firstly, mean AAA volume growth/year in patients with calciﬁcation volume < median was assumed to be 12 cm³/yr, SD = 6.5 cm³/yr based on results from a previous CT study [17]. Secondly we predicted that AAA growth rate would be 42% greater
in patients with calcification volume < median as suggested by results from a study by Lindholt and colleagues [16]. Using the G-power 3.1.9.2 tool, (Two tailed t-test: difference between means \( \alpha = 0.05 \), Power = 0.95), 30 observations in each group were needed.

All results were transcribed onto an excel spreadsheet and later transferred to SPSS version 22 (IBM SPSS) for statistical analysis. Chi squared t-test and Mann–Whitney U test were used to compare nominal and continuous variables, respectively, between groups. Medians and interquartile ranges were reported for variables which were not normally distributed and means and standard deviations were used to describe results for continuous variables following a normal distribution. Correlation analyses were performed between continuous variables where the dependant variable followed a normal distribution and reported as Pearson’s correlation coefficient.

3. Results

Records of 173 patients who underwent serial CT imaging of their AAA between June 2003 and Nov 2013 were screened for inclusion into the study. A total of 88 patients met eligibility criteria and were included. Median time interval between the two CT scans assessed from each patient was 16 months, interquartile range 12–28 months. Thirty percent of the cohort were current smokers at recruitment and 74% were male. The median baseline infra-renal aortic calcification volume for the cohort was 1600 mm\(^3\), interquartile range of 830–2800 mm\(^3\). Median baseline infra-renal aortic volume was 91 cm\(^3\) with an interquartile range of 73–108 cm\(^3\). Mean baseline orthogonal diameter was 43.0 mm, standard deviation was 5.5 mm.

3.1. Reproducibility

31 subjects were involved in the reproducibility study. Inter-observer and intra-observer 95% limits of agreement for infra-renal aortic calcification volume were (lower, upper): \(-140, 5.8 \) mm\(^3\) and \(0.68, 97 \) mm\(^3\), respectively. Ninety-five percent limits of agreement for infra-renal aortic volume were (lower, upper): \(-4.5, 0.027 \) cm\(^3\) and \(-0.007, 3.0 \) cm\(^3\) for inter and intra-observer error respectively. The concordance correlation coefficients (CCC) for inter-observer and intra-observer variability in

**Fig. 1.** Computed tomography images illustrating the method of assessing infra-renal aortic calcification volume. Three images displaying the acquisition of infra-renal aortic calcification volume. The coronal view of the abdominal aorta with the green lines at the upper and lower boundaries set at the slice inferior to the lowest renal artery and the slice above the aortic bifurcation respectively (a). The axial images depict the selection of aortic boundary to include outer wall (b). The extracted calcification volume from the region of interest is calculated based on predefined thresholds for a calcification specific radio-density spectrum. A 3D reconstruction of the calcium volume within the region of interest is depicted in (c).
measuring infra-renal aortic volume were 0.98 and 0.99, respectively. In measuring infra-renal aortic calcification volume the CCCs were 0.99 and 0.99 for inter-observer and intra-observer variability, respectively (Table 1).

### 3.2. Association of aortic calcification with AAA expansion

None of baseline characteristics assessed were significantly different between the groups with <median or ≥median calcification volume (Table 2). Median infra-renal aortic volume increase in the group with <median calcification was 6.0 cm³/yr compared to 7.8 cm³/yr in the group with ≥median (p = 0.66). Median AAA orthogonal diameter increase was 1.6 mm/yr in the group with <median calcification and 1.8 mm/yr for the group with ≥median (p = 0.99). Actual and percentage AAA growth rates were similar in both groups (Table 3). Infra-renal aortic calcification volume had minimal correlation with percentage AAA volume change/yr (Pearson correlation = 0.10, p = 0.36). Mean percentage AAA volume change/yr was 4.2 ± 6.4% and 8.9 ± 6.2% for patients with and without diabetes, respectively (p = 0.003).

### 4. Discussion

The findings from this study suggest that infra-renal aortic calcification volume can be measured from CT with good inter and intra-observer reproducibility compared to other published studies on the subject [17,22]. Results from this cohort suggest that infra-renal aortic calcification volume is not significantly associated with small AAA growth rates. Findings from this cohort of patients are consistent with previous US and CT based studies in reporting that diabetes mellitus is associated with reduced AAA growth rates [1,23,24].

Patients with diabetes are believed to have more marked vascular calcification and thus one possible explanation for reduced AAA expansion in these patients could be through more advanced aortic calcification [25]. Our findings however suggest that aortic calcification does not explain the reduced AAA expansion in patients with diabetes. The changes seen in the walls of aneurysmal aortas include inflammation and the activation of proteolytic pathways associated with loss of elastin and other structural proteins [26]. In contrast, diabetes is associated with increased synthesis and reduced degradation of extracellular matrix. The deposition of advanced glycosylation end products also renders vascular matrix resistant to proteolysis and inflammation in patients with diabetes [25]. It is likely these changes in the extracellular matrix are responsible for the reduced AAA expansion in patients with diabetes rather than any effects of aortic calcification.

Previous studies have alluded to the potential that AAA calcification may be a novel additional tool for AAA rupture risk assessment [13]. This has been supported by FEA findings of increased PWS in AAA regions with calcified plaque [9–11]. However our results did not suggest a significant association between infra-renal aortic calcification volume and AAA growth. This suggests that incorporating AAA calcification in rupture risk assessment is unlikely to improve accuracy of predicting rupture, although a larger long-term prospective study would be needed to confirm this. Thus
we currently have insufficient evidence to necessitate consideration of infra-renal aortic calcification volume when selecting patients for intervention. Additionally, this study contributes to the evidence refuting the theory that suggests heavily calcified AAAs grow slower than less calcified AAAs. This is important in avoiding misconceived alterations to AAA surveillance periods based on the degree of AAA calcification. Our current data suggests that calcified AAAs should have similar imaging surveillance as other AAAs. Moreover, our findings do not support the exploration of pharmacological manipulation of AAA calcification as a means to slow expansion of AAAs.

This study has a number of limitations. Firstly, only 88 patients were included in the current study. We based this on a sample size calculation which assumed a 42% difference in AAA growth rate between the two groups thus we were underpowered to examine smaller differences in AAA growth. We assessed AAA growth from CT scans performed at varying intervals. In order to adjust for these differences in time intervals we had to assume linear AAA growth which is not always the case although we found this to be present in a recent cohort [27]. Furthermore, the possibility of selection bias should be noted. Patients who undergo CT rather than US surveillance are commonly those with one or more of the following attributes: obesity, a particularly tortuous aorta, and those with a large baseline AAA diameter. On the other hand, patients presumably omitted from this cohort include those who may need CT imaging but are at increased risk of contrast induced nephropathy. Finally we only assessed aortic calcification volume. It is conceivable that other aspects of calcification like is distribution or thickness might have separate impact on the growth of small AAAs however we suspect this is unlikely given the lack of association of calcification volume.

In conclusion despite the established association of vascular calcification with increased incidence of cardiovascular events [28], infrarenal aortic calcification volume does not appear to be associated with small AAA growth. The finding of the current study is consistent with findings from a previous study by our group [17].

Conflict of interest

Authors declare no conflict of interest.

Acknowledgements

JG is supported by grants from the National Health and Medical Research Council [project grants 1063476, 1022752, 1021416, 1020955, 1003707; Centre of Research Excellence 1000967; fellowship 1019921], and The Office of Health and Medical Research, Queensland Government. JG holds a Practitioner Fellowships from the National Health and Medical Research Council, Australia (1019921). JG holds a Senior Clinical Research Fellowship from the Office of Health and Medical Research, Queensland. 

References