Gaarder and Seierstad Cardiovascular Ultrasound 2015, 13:5 http://www.cardiovascularultrasound.com/content/13/1/5

RESEARCH



Open Access

Measurements of carotid intima media thickness in non-invasive high-frequency ultrasound images: the effect of dynamic range setting

Mario Gaarder^{1,2*} and Therese Seierstad¹

Abstract

Background: Carotid intima media thickness (CIMT) measured with ultrasound (US) is widely used as biomarker for arteriosclerosis and as surrogate endpoint in interventional studies to assess efficacy of drug therapies. Strict US protocols are necessary to ensure reproducibility. The range of US signal intensities used for image formation, the dynamic range (DR), is rarely reported in studies and little is known about its effect on CIMT measurements in humans. The purpose of this study was to quantify the impact of DR on measurements of CIMT.

Methods: US was used to examine 313 carotid arteries in participants from two different clinical studies. For each artery, images with DR of 40, 55, 70 and 85 dB were captured from the same frozen US frame. Mean CIMT (CIMT_{mean}), maximum CIMT (CIMT_{max}) and standard deviation of CIMT (CIMT_{sd}) were obtained for all images. CIMT for different DRs were compared using student *t*-test.

Results: $CIMT_{mean}$ for 40, 55, 70 and 85 dB were 0.529, 0.564, 0.590 and 0.605 mm respectively. For $CIMT_{max}$ the corresponding values were 0.626, 0.667, 0.698, and 0.716 mm. $CIMT_{mean}$ and $CIMT_{max}$ increased significantly for increasing DR steps (p < 0.01). The relative change in $CIMT_{mean}$ and $CIMT_{max}$ were largest between 40 and 55 dB (6.7% and 7.0%) and smallest between 70 and 85 dB (2.6% and 2.7%) indicating a declining dependency for increasing DR.

Conclusions: DR significantly changes CIMT measurements and the changes are most prominent for lower DRs. The effect of changing DR is larger in human arteries than in phantoms. Reporting the DR will therefore increase the validity of CIMT data.

Keywords: Carotid intima media thickness, Dynamic range, Ultrasound, Carotid arteries

Background

Ultrasound (US) measurements of carotid intima-media thickness (CIMT) are used to assess the extent of arteriosclerosis and are a biomarker for the future risk of cardiovascular disease (CVD) and stroke [1-6]. Rate of CIMT change is also used as a surrogate endpoint for CVD in interventional studies to assess early effects of drug therapies [7,8].

In US images the carotid artery wall is depicted as three separate layers: intima, media and adventitia (Figure 1). The CIMT is defined as the distance from the lumen-

* Correspondence: s.m.gaarder@medisin.uio.no

intima interface to the media-adventitia interface [9]. The intima-media complex can easily be distinguished from the surrounding tissue in US images and the distinct borders allows for manual as well as automatic measurements of the CIMT.

Reproducibility is a prerequisite for a reliable biomarker, and standardization of imaging and measurement of CIMT will minimize variability between measurements and between readers. Development of US protocols for measuring CIMT has been focusing on choice of vessel segment, near or far wall measurements, and frequency setting as well as performance of the technicians and the observers [5]. The updated Mannheim Carotid Intima-Media Thickness and Plaque Consensus [5] suggests that the dynamic range (DR) should be set at approximately 60



© 2015 Gaarder and Seierstad; licensee BioMed Central. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly credited. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

¹Department of Radiology and Nuclear Medicine, Oslo University Hospital, P.O. Box 4950, Nydalen 0424, Oslo, Norway

²Faculty of Clinical Medicine, University of Oslo, P.O. Box 1078, Blindern 0316, Oslo, Norway



decibel (dB). The DR defines the number of US echo intensities included in the image formation process. Increased DR will increase the range of signal intensities resulting in a low-contrast image. Oppositely, decreased DR will decrease the range of signal intensities resulting in a high-contrast image.

It has previously been shown that change in DR has a significant effect on measurements of wall thickness and lumen diameter in phantoms [10], however the documentation on effects in human arteries are lacking.

The purpose of this study was to quantify the effects of changing DR in non-invasive US CIMT measurements in the common carotid arteries (CCA).

Methods

Study population

From August 2012 to January 2014, 165 participants were included in this prospective study. The participants were recruited from the two clinical studies *Arteriosclerosis and childhood diabetes* (the ABD-study, n = 96) and *Impact of insulin sensitivity on cardiovascular risk markers during* 9-19 years of follow-up (the INFO-study, n = 69). The characteristics of the populations are presented in Table 1. The Regional Committee for Medical and Health Research Ethics South East approved both studies (ABD (REC ID 2011/1818), INFO (REC ID 2010/3339)) and written informed consent was obtained from all participants.

US examination

All US examinations were performed using Zonare Z-One ULTRA US equipment (Zonare Medical Systems, Mountain View, CA, USA) equipped with a linear 10–5 transducer and a three lead electrocardiogram (ECG). A frequency of 8 MHz and its harmonic frequencies were used. All examinations were performed using the same protocol

Table 1 Demographic data

	Total	The ABD-study	The INFO-study
Arteries [n]	313	181*	132**
Participants [n]	165	96	69
Males [%]	31	47	100
Age [yrs.]	27.7 (10.9)	19.0 (3.1)	39.9 (3.5)
Diabetics [%]	31	60	1
Systolic BP [mmHg]	123.6 (11.1)	119.6 (9.5)	129.2 (10.7)
Diastolic BP [mmHg]	74.5 (9.5)	69.9 (7.6)	81.0 (8.1)

*missing DR 40 dB data in one participant, **missing DR 40 dB data in two participants, and 55, 70 and 85 dB data in one participant. Only one artery was examined in nine participants in the ABD-study and in five participants in the INFO-study.

(compound imaging, edge enhancement and standard B-mode grey scale images) and by the same operator (MG). The study participants were instructed to abstain from alcohol, caffeine and nicotine 12 hours prior to the examination and to only consume clear liquids on the morning of the examination.

All participants were examined in the supine position with the neck extended and the head tilted slightly towards the opposite of the examined side. Both left and right CCAs were depicted. The three ECG pads were attached to both shoulders and the left hip prior to US examination, and brachial blood pressure was measured after at least ten minutes in the supine position.

The distal two centimetres of the CCAs were depicted in the longitudinal plane, with the diverging of the artery walls towards the carotid bifurcation as the distal limitation. The intima- and media layers were visible in both the near and the far wall. The images were captured in the end diastolic phase (ECG-assisted), when the artery had its smallest diameter. The image plane was most often an oblique sagittal plane, however, depending on the arteries tortuousness and depth, the image plane was shifted towards the coronal plane to better visualize the arteries parallel to the skin surface. The end-diastolic frame of the third cardiac cycle was chosen to explore the effect of DR changes on CIMT measures. For this frame the DR setting was changed in 15 dB steps from 40 dB to 85 dB, and images for the different DR settings were obtained. All other imaging parameters were kept fixed. Thus, four identical images, apart from the DR setting, were acquired from each artery (Figure 2). The examinations were saved in the DICOM format and exported to a remote computer for measurements of the CIMT.

CIMT measurements

The commercially available semi-automatic intima-media thickness measurement software Máth 3.2.0 (Intelligence in Medical Imaging, Paris, France) was used for CIMT measurements. The operator identified the area to measure: a continuous one-centimetre segment of the distal two



and CIMT_{max} were 0.691 and 0.750 millimetre.

centimetres in the CCA far wall. The software automatically detected the intima-lumen and the media-adventitia interfaces and calculated average CIMT (CIMT_{mean}), maximum CIMT (CIMT_{max}) and the local variation of CIMT in the segment (CIMT_{sd}) for DR of 40, 55, 70 and 85 dB, respectively. The coefficient of variance (CV) was calculated for repeated measurements in 10 randomly selected participants, for both CIMTmean and CIMTmax. The CV and the corresponding 95% confidence interval (95% CI) was calculated for 10 repeated measurements for 40, 55, 70 and 85 dB.

Data analyses

The statistical analysis was done in SPSS (IBM SPSS Statistics 21, Armonk, NY, USA). Descriptive statistics was used to describe the study populations. Groups of normally distributed data were compared using the two-way student t-test and the paired student *t*-tests.

Results

A total of 313 CCAs were included in the analysis: 181 CCAs from participants in the ABD-study and 132 CCAs from participants in the INFO-study. Population mean $\text{CIMT}_{\text{mean}}$ was 0.529, 0.564, 0.590 and 0.605 mm with DR of 40, 65, 70 and 85 dB, respectively.

Corresponding results for CIMT_{max} was 0.626, 0.667, 0.698 and 0.716. Differences between the DR steps were all statistically significant (p < 0.01) (Tables 2 and 3). The relative increases in CIMT_{mean} between DR steps were equal to the relative increase in CIMT_{max} for the same steps (p > 0.44). The relative increases for CIMT_{mean} and CIMT_{max} were 6.7% and 7.0% for 40–55 dB, 4.9% and 4.8% for 55–70 dB, and 2.6% and 2.7% for 70–85 dB, respectively.

Average change in $\text{CIMT}_{\text{mean}}$ from 40 to 85 dB was 0.076 mm corresponding to approximately 0.008 mm for each 5 dB increase. For all measurements mean CIMT_{sd} was less than 0.05 mm.

The CV for CIMT_{mean} was 0.65% (95% CI; 0.48-0.83) for 40 dB, 0.42% (95% CI; 0.28-0.56) for 55 dB, 0.86% (95% CI; 0.51-1.21) for 70 dB and 0.59% (95% CI; 0.37-0.82) for 85 dB. For CIMT_{max} the CVs were 0.59% (95% CI; 0.33-0.85), 0.85% (95% CI; 0.16-1.36), 1.36 (95%CI; 0.37-2.37) and 1.04% (95% CI; 0.41-1.67) for 40, 55, 70 and 85 dB, respectively.

Discussion

This study showed that an increase in DR in US image acquisition yields increased CIMT values and that the effect of increasing DR is largest for lower DRs. The DR

Population	CIMT measure	n	DR setting				
			40 dB	55 dB	70 dB	85 dB	
ABD	CIMT _{mean} [mm]	181	0.488 ± 0.066	0.521 ± 0.066	0.547 ± 0.066	0.563 ± 0.066	
	CIMT _{max} [mm]	181	0.578 ± 0.074	0.619 ± 0.072	0.648 ± 0.071	0.669 ± 0.070	
INFO	CIMT _{mean} [mm]	130	0.586 ± 0.091	0.621 ± 0.093	0.648 ± 0.095	0.662 ± 0.095	
	CIMT _{max} [mm]	130	0.691 ± 0.107	0.732 ± 0.107	0.765 ± 0.109	0.780 ± 0.108	
Total	CIMT _{mean} [mm]	311	0.529 ± 0.091	0.564 ± 0.093	0.590 ± 0.094	0.605 ± 0.093	
	CIMT _{max} [mm]	311	0.626 ± 0.105	0.667 ± 0.105	0.698 ± 0.106	0.716±0.104	
INFO Total	CIMT _{max} [mm] CIMT _{mean} [mm] CIMT _{max} [mm] CIMT _{mean} [mm] CIMT _{max} [mm]	181 130 130 311 311	0.578 ± 0.074 0.586 ± 0.091 0.691 ± 0.107 0.529 ± 0.091 0.626 ± 0.105	0.619 ± 0.072 0.621 ± 0.093 0.732 ± 0.107 0.564 ± 0.093 0.667 ± 0.105	0.648 ± 0.071 0.648 ± 0.095 0.765 ± 0.109 0.590 ± 0.094 0.698 ± 0.106	0.669 ± 0 0.662 ± 0 0.780 ± 0 0.605 ± 0 0.716 ± 0	

Table 2 Measurements of CIMT with different DR settings

Data on DR 40 dB was missing in one participant in the ABD group and two in the INFO group.

All measurement values are given as mean and standard deviation.

should therefore be specified in all US protocols where CIMT values are used either directly or as a predictive biomarker, especially in longitudinal follow-up assessments.

As DR increases the range of signal intensities included in the image formation increases. This increase yields reduced image contrast and less defined interfaces between the intima, media and adventitia layers in the artery wall. Consequently, it is anticipated that CIMT would increase with increasing DR. In the interval between 40 and 85 dB the average increase in CIMT_{mean} per 5 dB increase in DR was 0.008 mm. This is approximately 2.5 times higher than the values found by Potter et al. in a phantom study [10]. The phantoms were made of tissue-mimicking agar with wall thicknesses of 0.5, 1.0 and 1.5 mm and acoustic properties similar to tissue. In our study, the changes in CIMT measurements due to DR variation were anticipated to be in the same range as the phantom with 0.5 mm wall thickness. The discrepancy in findings may be related to the composition of the wall (phantom versus human arteries). In contrast to human arteries, the phantom walls were homogenous with defined interfaces and consequently, interface echo signals will be more distinct in phantoms than in human arteries. The relative change in CIMT_{mean} for CCAs was largest between 40 and 55 dB (6.7%) and smallest between 70 and 85 dB (2.6%) indicating a declining dependency for increasing DR. The relative increase found by Potter et al. for DR of 40 to 60 dB was about 3% and as such our findings suggest an underestimation of DR dependency in phantoms, especially for low DRs. The algorithms used for the automatic edge detection as well as larger variation in the tissue composition of human arteries than in the phantoms might have contributed to the differences between the studies. We speculate that the variation in the human tissue causes a broader spectrum of echo signals, and that the effect of widening the DR therefore will be more prominent in this setting. However, despite differences in magnitude both Potter et al. and we found that CIMT increased for increasing DR.

The CIMT_{mean} and CIMT_{max} found for different DRs in our study emphasize the need for reporting DR used in US protocols as CIMT and change in CIMT over time are used as tools for risk-stratification, as surrogate endpoints, and as references for clinical decision-making: European Guidelines on cardiovascular disease prevention in clinical practise 2012 states that CIMT > 0.9 mm is considered abnormal [11]. The extreme DR values (40 and 85 dB) in our study yielded a difference in CIMT_{mean} and CIMT_{max} of 8-10% of the suggested reference value. Knowledge of DR setting is important for patients where CIMT values are close to cut-off values or reference values of clinical importance. Different DR between baseline and follow-up examinations may lead to under- or overestimation of CIMT change. Outcome of multicentre studies

Table 3 Absolute	differences i	in CIMT	between	the	different	DR	steps
------------------	---------------	---------	---------	-----	-----------	----	-------

	DR step	R step ABD		11	NFO	TOTAL		
		Difference	95% CI	Difference	95% CI	Difference	95% CI	
	[dB]	[mm]	[mm]	[mm]	[mm]	[mm]	[mm]	
CIMT _{mean}	40-55	0.032	(0.030, 0.035)	0.037	(0.034, 0.041)	0.034	(0.032, 0.036)	
	55-70	0.026	(0.025, 0.028)	0.027	(0.025, 0.030)	0.027	(0.025, 0.028)	
	70-85	0.016	(0.014, 0.017)	0.014	(0.012, 0,016)	0.015	(0.014,0.016)	
CIMT _{max}	40-55	0.041	(0.037,0.044)	0.043	(0.039, 0.048)	0.042	(0.039, 0.045)	
	55-70	0.029	(0.026,0.032)	0.033	(0.029, 0.038)	0.031	(0.029, 0.033)	
	70-85	0.020	(0.018,0.023)	0.015	(0.012, 0.019)	0.018	(0.016, 0.020)	

For all comparisons p < 0.001. All differences are given as mean values, and in millimetres.

may also be compromised. Thus, by including information of DR in guidelines, reference data and study protocols will increase the validity of the results.

A shortcoming of our study is the large DR steps (15 dB). Smaller DR steps would probably give more precise results of the relation between measured CIMT and DR setting. However, the wide range of DRs included allowed identification of the non-linearity between change in CIMT and change in DR that would probably not have been detected if DR had been restricted to 40 to 60 dB, as in Potter et al. [10]. The large number of participants and the fact that the relative CIMT changes for different DRs were similar for the two study populations differing in age and gender mixture strengthens the external validity of our findings.

Conclusions

In conclusion, DR significantly changes CIMT measurements and the changes are most prominent for lower DRs. The effect of changing DR is larger in human arteries than in phantoms. Reporting the DR will therefore increase the validity of CIMT data, especially in prospective clinical trials.

Abbreviations

CCA: Common carotid artery; CIMT: Carotid intima media thickness; DR: Dynamic range; US: Ultrasound.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

MG designed the study, collected and substantially contributed to the data analyses and drafted the manuscript. TS contributed to data analyses and critically reviewed the manuscript. Both authors have read and approved the final manuscript.

Acknowledgements

We gratefully acknowledge the principal investigators of the two clinical trials, Sigrid Skårn and Martin Heier, for including the participants in this study and the Norwegian Society for Radiographers for financial support.

Received: 1 November 2014 Accepted: 22 January 2015 Published: 27 January 2015

References

- O'Leary DH, Polak JF, Kronmal RA, Manoloio TA, Burke GL, Wolfson SK, et al. Carotid-artery intima and media thickness as a risk factor for myocardial infarction and stroke in older adults. N Engl J Med. 1999;340(1):14–22.
- Lorenz MW, Polak JF, Kavousi M, Mathiesen EB, Volzke H, Tuomainen TP, et al. Carotid intima-media thickness progression to predict cardiovascular events in the general population (the PROG-IMT collaborative project): a meta-analysis of individual participant data. Lancet. 2012;379(9831):2053–62.
- 3. Cobble M, Bale B. Carotid intima-media thickness: knowledge and application to everyday practice. Postgrad Med. 2010;122(1):10–8.
- Coll B, Feinstein SB. Carotid intima-media thickness measurements: techniques and clinical relevance. Curr Atheroscler Rep. 2008;10(5):444–50.
- Touboul PJ, Hennerici MG, Meairs S, Adams H, Amarenco P, Bornstein N, et al. Mannheim carotid intima-media thickness and plaque consensus (2004-2006-2011). An update on behalf of the advisory board of the 3rd, 4th and 5th watching the risk symposia, at the 13th, 15th and 20th European Stroke Conferences, Mannheim, Germany, 2004, Brussels, Belgium, 2006, and Hamburg, Germany, 2011. Cerebrovasc Dis. 2012;34(4):290–6.

- Doneen AL, Bale BF. Carotid intima-media thickness testing as an asymptomatic cardiovascular disease identifier and method for making therapeutic decisions. Postgrad Med. 2013;125(2):108–23.
- Huang Y, Li W, Dong L, Li R, Wu Y. Effect of statin therapy on the progression of common carotid artery intima-media thickness: an updated systematic review and meta-analysis of randomized controlled trials. J Atheroscler Thromb. 2013;20(1):108–21.
- Davidson MH, Rosenson RS, Maki KC, Nicholls SJ, Ballantyne CM, Mazzone T, et al. Effects of Fenofibric Acid on Carotid Intima-Media Thickness in Patients With Mixed Dyslipidemia on Atorvastatin Therapy: Randomized, Placebo-Controlled Study (FIRST). Arterioscler Thromb Vasc Biol. 2014;34(6):1298–306.
- Pignoli P, Tremoli E, Poli A, Oreste P, Paoletti R. Intimal plus medial thickness of the arterial wall: a direct measurement with ultrasound imaging. Circulation. 1986:74(6):1399–406.
- Potter K, Reed CJ, Green DJ, Hankey GJ, Arnolda LF. Ultrasound settings significantly alter arterial lumen and wall thickness measurements. Cardiovasc Ultrasound. 2008;6:6.
- Perk J, De Backer G, Gohlke H, Graham I, Reiner Z, Verschuren WMM, et al. European guidelines on cardiovascular disease prevention in clinical practice (version 2012). Eur J Prevent Cardiol. 2012;19(4):585–667.

doi:10.1186/1476-7120-13-5

Cite this article as: Gaarder and Seierstad: **Measurements of carotid** intima media thickness in non-invasive high-frequency ultrasound images: the effect of dynamic range setting. *Cardiovascular Ultrasound* 2015 13:5.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

) BioMed Central

Submit your manuscript at www.biomedcentral.com/submit