Integrity Check for the Epidemiology of Diabetes Interventions and Complications Study (EDIC) One Year Intima-media Wall Thickness (IMT) Datasets

As a partial check of the integrity of the EDIC One Year IMT datasets archived in the NIDDK data repository, a set of tabulations was performed to verify that published results can be reproduced using the archived datasets. Analyses were performed to duplicate published results for the data reported by the EDIC Research Group [1] in *Diabetes* in February 1999. The results of this integrity check are described below. The full text of the *Diabetes* article can be found in Attachment 1, and the SAS code for our tabulations is included in Attachment 2.

The intent of this DSIC is to provide confidence that the data distributed by the NIDDK repository is a true copy of the study data. Our intent is *not* to assess the integrity of the statistical analyses reported by study investigators. As with all statistical analyses of complex datasets, complete replication of a set of statistical results should not be expected on a first (or second) exercise in secondary analysis. This occurs for a number of reasons including differences in the handling of missing data, restrictions on cases included in samples for a particular analysis, software coding used to define complex variables, etc. Experience suggests that most discrepancies can ordinarily be resolved by consultation with the study data coordinating center (DCC), however this process is labor-intensive for both DCC and Repository staff. It is thus not our policy to resolve every discrepancy that is observed in an integrity check. Specifically, we do not attempt to resolve minor or inconsequential discrepancies with published results or discrepancies that involve complex analyses, *unless staff of the NIDDK Repository suspect that the observed discrepancy suggests that the dataset may have been corrupted in storage, transmission, or processing by repository staff.* We do, however, document in footnotes to the integrity check those instances in which our secondary analyses produced results that were not fully consistent with those reported in the target publication.

Background. The Diabetes Control and Complications Trial (DCCT), 1983-1993, was a multi-center trial designed to determine whether intensive therapy to maintain blood glucose and glycosolated hemoglobin concentrations as close to the normal range as possible would prevent or delay long-term complications in patients with type 1 diabetes. This trial showed a markedly reduced risk of microvascular complications as compared with conventional therapy. Most participants were then enrolled in the EDIC study, a long-term observational study. One of the objectives of EDIC was to study the development and progression of atherosclerotic cardiovascular disease in type 1 diabetes. The published data describe the role of cardiovascular risk factors and antecedent therapy in the DCCT on carotid IMT in type 1 diabetes [1].

In summary, approximately 18 months after the end of the DCCT, high-resolution B-mode ultrasonography was used to assess the carotid arteries of 1,325 patients. An age- and sex-matched nondiabetic population (n=153) was studied with the same protocol [1].

Baseline Data. Table 1 of the 1999 *Diabetes* article reports on baseline characteristics. Variables summarized in this baseline table (Table 1. Baseline clinical characteristics of EDIC participants by sex and original treatment group assignment during the DCCT) are taken from the EDIC One Year IMT analysis dataset, EDICIMT1, created for this study. Original treatment group assignment was verified against the DCCT baseline dataset, BASELINE. Table A lists the variables used for replication of the Table 1 variables.

Table 1 Variable	Variables Used in Replication		
Sample size	group, sex		
Attained age (years)	att_age		
Attained duration of type 1 diabetes (months)	att_durn		
Height (cm)	height		
BMI (kg/m ²)	bmi		
BMI > 27 (%)	bmi_27		
Natural WHR	nwst_hip		
sBP (mmHg)	sbp		
dBP (mmHg)	dbp		
Hypertensive (%)	ht		
Total cholesterol (mg/dl)	tchol		
HDL cholesterol (mg/dl)	hdl		
LDL cholesterol (mg/dl)	ldl		
Triglycerides (mg/dl)	trig		
AER (mg/24 h)	aer		
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	aer_40		
GFR	gfr		
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	std_ins		
Cigarette smoking (current) (%)	smoking		
HbA _{1c}	hbalc		
Mean HbA _{1c} during DCCT	avg_alc		
Framingham score	fscore		
Note: The EDIC baseline (1994-1995) is after an average of 6.5 years of intensive treatment in DCCT. Hypertensive is defined as sitting $sBP \ge 140$ mmHg and/or $dBP \ge 90$ mmHg or the use of antihypertensive medication. Framingham score is defined as described by Anderson et al. (An Updated Coronary Risk Profile. <i>Circulation</i> 83:355-365, 1991).			

Table A: Variables Used to Replicate Table 1

In Table B, we compare the results for characteristics calculated from the archived dataset to the results published in the 1999 *Diabetes* article. As Table B shows, the results obtained from the archived data are the same as those in the published tabulations, with a few minor exceptions.

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Articl	е
Values	

Table 1 Variable	Women, Treatment: Intensive		
	Diabetes (1999)	Integrity Check	Difference
Sample size	321	321	0
Attained age (years)	35 ± 7	35 ± 7	0
Attained duration of type 1 diabetes (months)	168 ± 58	168 ± 58	0
Height (cm)	165 ± 6	165 ± 6	0
BMI (kg/m^2)	26.5 ± 4.6	26.5 ± 4.6	0
BMI > 27 (%)	39.7	39.7	0
Natural WHR	0.76 ± 0.07	0.76 ± 0.07	0
sBP (mmHg)	114 ± 12	114 ± 12	0
dBP (mmHg)	74 ± 9	74 ± 9	0
Hypertensive (%)	9.7	9.7	0
Total cholesterol (mg/dl)	188 ± 36	188 ± 36	0
HDL cholesterol (mg/dl)	59 ± 14	59 ± 14	0
LDL cholesterol (mg/dl)	113 ± 29	113 ± 29	0
Triglycerides (mg/dl)	84 ± 74	84 ± 74	0
AER (mg/24 h)	21.1 ± 63.2	21.1 ± 63.2	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	6.9	6.9	0
GFR	117.5 ± 23.5	117.5 ± 23.5	0
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	0.63 ± 0.21	0.63 ± 0.21	0
Cigarette smoking (current) (%)	19.7	19.8	0.1
HbA _{1c}	7.9 ± 1.4	7.9 ± 1.4	0
Mean HbA _{1c} during DCCT	7.3 ± 0.9	7.3 ± 0.9	0
Framingham score	0.018 ± 0.025	0.018 ± 0.025	0
Note: Data are means \pm SD, n, or %			

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Article
Values (cont.)

Table 1 Variable	Women, Treatment: Conventional		
	Diabetes (1999)	Integrity Check	Difference
Sample size	313	313	0
Attained age (years)	34 ± 7	34 ± 7	0
Attained duration of type 1 diabetes (months)	170 ± 61	170 ± 61	0
Height (cm)	165 ± 6	165 ± 6	0
BMI (kg/m ²)	25.1 ± 3.6	25.1 ± 3.6	0
BMI > 27 (%)	27.5	27.5	0
Natural WHR	0.76 ± 0.07	0.76 ± 0.07	0
sBP (mmHg)	114 ± 13	114 ± 13	0
dBP (mmHg)	72 ± 10	72 ± 10	0
Hypertensive (%)	12.9	12.9	0
Total cholesterol (mg/dl)	188 ± 39	188 ± 39	0
HDL cholesterol (mg/dl)	59 ± 14	59 ± 14	0
LDL cholesterol (mg/dl)	113 ± 32	113 ± 32	0
Triglycerides (mg/dl)	82 ± 73	82 ± 73	0
AER (mg/24 h)	63.5 ± 313.9	63.5 ± 313.9	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	16.0	16.0	0
GFR	119.3 ± 26.1	119.3 ± 26.1	0
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	0.64 ± 0.19	0.64 ± 0.19	0
Cigarette smoking (current) (%)	18.1	18.1	0
HbA _{1c}	8.2 ± 1.5	8.2 ± 1.5	0
Mean HbA _{1c} during DCCT	9.0 ± 1.4	9.0 ± 1.4	0
Framingham score	0.017 ± 0.028	0.017 ± 0.028	0
Note: Data are means \pm SD, n, or %			

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Article
Values (cont.)

Table 1 Variable	Women		
	Diabetes (1999)	Integrity Check	Difference
Attained age (years)	0.042	0.042	0
Attained duration of type 1 diabetes (months)	0.722	0.722	0
Height (cm)	0.682	0.680	0.002
BMI (kg/m ²)	< 0.001	< 0.001	0
BMI > 27 (%)	0.001	0.001	0
Natural WHR	0.731	0.731	0
sBP (mmHg)	0.739	0.739	0
dBP (mmHg)	0.101	0.101	0
Hypertensive (%)	0.210	0.210	0
Total cholesterol (mg/dl)	0.495	0.495	0
HDL cholesterol (mg/dl)	0.737	0.737	0
LDL cholesterol (mg/dl)	0.542	0.542	0
Triglycerides (mg/dl)	0.427	0.427	0
AER (mg/24 h)	0.003	0.003	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	< 0.001	< 0.001	0
GFR	0.069	0.069	0
Insulin dose (U • $kg^{-1} • day^{-1}$)	0.801	0.800	0.001
Cigarette smoking (current) (%)	0.605	0.605	0
HbA _{1c}	0.009	0.009	0
Mean HbA _{1c} during DCCT	< 0.001	< 0.001	0
Framingham score	0.037	0.037	0
Note: Data are p-values	•	•	

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Article	е
Values (cont.)	

Table 1 Variable	Men, Treatment: Intensive		
	Diabetes (1999)	Integrity Check	Difference
Sample size	340	340	0
Attained age (years)	36 ± 7	36 ± 7	0
Attained duration of type 1 diabetes (months)	167 ± 59	167 ± 59	0
Height (cm)	178 ± 7	178 ± 7	0
BMI (kg/m^2)	26.8 ± 4.2	26.8 ± 4.2	0
BMI > 27 (%)	41.2	41.2	0
Natural WHR	0.88 ± 0.08	0.88 ± 0.08	0
sBP (mmHg)	119 ± 11	119 ± 11	0
dBP (mmHg)	77 ± 9	77 ± 9	0
Hypertensive (%)	21.2	21.2	0
Total cholesterol (mg/dl)	187 ± 35	187 ± 35	0
HDL cholesterol (mg/dl)	49 ± 13	49 ± 13	0
LDL cholesterol (mg/dl)	119 ± 30	119 ± 30	0
Triglycerides (mg/dl)	98 ± 77	98 ± 77	0
AER (mg/24 h)	28.6 ± 112.9	28.6 ± 112.9	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	7.4	7.4	0
GFR	115.3 ± 19.7	115.3 ± 19.7	0
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	0.67 ± 0.24	0.67 ± 0.24	0
Cigarette smoking (current) (%)	19.3	19.4	0.1
HbA _{1c}	7.9 ± 1.2	7.9 ± 1.2	0
Mean HbA _{1c} during DCCT	7.2 ± 0.9	7.2 ± 0.9	0
Framingham score	0.039 ± 0.038	0.039 ± 0.038	0
Note: Data are means \pm SD, n, or %			

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Article	е
Values (cont.)	

Table 1 Variable	Men, Treatment: Conventional		
	Diabetes (1999)	Integrity Check	Difference
Sample size	351	351	0
Attained age (years)	36 ± 7	36 ± 7	0
Attained duration of type 1 diabetes (months)	159 ± 55	159 ± 55	0
Height (cm)	179 ± 7	179 ± 7	0
BMI (kg/m ²)	25.9 ± 3.2	25.9 ± 3.2	0
BMI > 27 (%)	34.5	34.5	0
Natural WHR	0.87 ± 0.09	0.87 ± 0.09	0
sBP (mmHg)	120 ± 12	120 ± 12	0
dBP (mmHg)	77 ± 8	77 ± 8	0
Hypertensive (%)	17.6	17.6	0
Total cholesterol (mg/dl)	184 ± 36	184 ± 36	0
HDL cholesterol (mg/dl)	50 ± 12	50 ± 12	0
LDL cholesterol (mg/dl)	115 ± 31	115 ± 31	0
Triglycerides (mg/dl)	97 ± 78	97 ± 78	0
AER (mg/24 h)	46.1 ± 123.2	46.1 ± 123.2	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	17.9	18.0	0.1
GFR	116.1 ± 24.8	116.1 ± 24.8	0
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	0.65 ± 0.20	0.65 ± 0.20	0
Cigarette smoking (current) (%)	18.4	18.4	0
HbA _{1c}	8.4 ± 1.3	8.4 ± 1.3	0
Mean HbA _{1c} during DCCT	9.0 ± 1.1	9.0 ± 1.1	0
Framingham score	0.037 ± 0.035	0.037 ± 0.035	0
Note: Data are means \pm SD, n, or %			

Table B: Comparison of Baseline Table Values Computed in Integrity Check to Reference Article Values (cont.)

Table 1 Variable	Men		
	Diabetes (1999)	Integrity Check	Difference
Attained age (years)	0.912	0.912	0
Attained duration of type 1 diabetes (months)	0.096	0.096	0
Height (cm)	0.076	0.076	0
BMI (kg/m^2)	0.016	0.016	0
BMI > 27 (%)	0.069	0.069	0
Natural WHR	0.101	0.101	0
sBP (mmHg)	0.186	0.186	0
dBP (mmHg)	0.446	0.446	0
Hypertensive (%)	0.240	0.240	0
Total cholesterol (mg/dl)	0.166	0.166	0
HDL cholesterol (mg/dl)	0.742	0.742	0
LDL cholesterol (mg/dl)	0.145	0.145	0
Triglycerides (mg/dl)	0.756	0.755	0.001
AER (mg/24 h)	0.038	0.038	0
$AER \ge 40 \text{ mg}/24 \text{ h} (\%)$	< 0.001	< 0.001	0
GFR	0.335	0.335	0
Insulin dose $(U \cdot kg^{-1} \cdot day^{-1})$	0.269	0.269	0
Cigarette smoking (current) (%)	0.750	0.750	0
HbA _{1c}	< 0.001	< 0.001	0
Mean HbA _{1c} during DCCT	< 0.001	< 0.001	0
Framingham score	0.704	0.704	0
Note: Data are p-values			

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Nondiabetic Control Subject Data. Table 2 of the 1999 *Diabetes* article presents the IMT data including the average maximum of the common and internal carotid arteries stratified by sex and age decades for nondiabetic and type 1 diabetic subjects. Variables summarized in this follow-up table (Table 2. Comparison of IMT in nondiabetic subjects with EDIC type 1 diabetes cohort) are taken from both analysis datasets created for this study, EDICIMT1X (for the nondiabetic cohort) and EDICIMT1 (for the diabetic cohort). Table C lists the variables used for replication of the Table 2 variables.

Table 2 Variable	Variables Used in Replication		
	EDICIMT1X dataset: include all subjects		
Sampla siza	EDICIMT1 dataset : include all subjects where		
Sample size	'decade' variable has a value of '20 - 29', '30 - 39'		
	or '40 - 49'		
Sov	EDICIMT1X dataset: gender		
Sex	EDICIMT1 dataset: sex		
	EDICIMT1X dataset: common		
Common carotic artery	EDICIMT1 dataset: common		
Internal constid artemy	EDICIMT1X dataset: internal		
	EDICIMT1 dataset: internal		
A go (by doordo)	EDICIMT1X dataset: decade		
Age (by decade)	EDICIMT1 dataset: decade		
Note: Nondiabetic data were obtained from six no	rmal subjects without diabetes at each of the 29		
clinics using the EDIC ultrasound scanning protocol and read at the Central Ultrasound Reading Unit.			
Data exclude six EDIC patients who were 19 years old when scanned and five who were aged 50 or			
51 years old			

In Table D, we compare the follow-up results calculated from the archived dataset to the results published in the 1999 *Diabetes* article. As Table D shows, the results obtained from the archived data are the same as those in the published tabulations, with a few minor exceptions.

Table 2 Variable	Women, Cor	Women, Common Carotid Artery IMT (mm)			
		Nondiabetic			
	Diabetes (1999)	Integrity Check	Difference		
20-29	$25, 0.603 \pm 0.062$	$25, 0.603 \pm 0.062$	0		
30-39	$27, 0.663 \pm 0.074$	$27, 0.663 \pm 0.074$	0		
40-49	$25, 0.704 \pm 0.078$	$25, 0.704 \pm 0.078$	0		
		Type 1 diabetes			
	Diabetes (1999)	Integrity Check	Difference		
20-29	$172, 0.616 \pm 0.073$	$172, 0.616 \pm 0.073$	0		
30-39	$278, 0.657 \pm 0.081$	$278, 0.657 \pm 0.081$	0		
40-49	$178, 0.696 \pm 0.079$	$178, 0.696 \pm 0.079$	0		
		p-value			
	Diabetes (1999)	Integrity Check	Difference		
20-29	0.471	0.473	0.002		
30-39	0.453	0.448	0.005		
40-49	0.622	0.622	0		
Note: Data are sample size, mea rank-sum test after linear adjusti	$n \pm standard$ deviation and p nent for covariance with ag	p-values. P-values are f e.	rom Wilcoxon		

Table D: Comparison of Nondiabetic Control Subject Table Values Computed in Integrity Check to Reference Article Values

Table 2 Variable	Women Internal Carotid Artery IMT (mm)			
	Nondiabetic			
	Diabetes (1999)	Integrity Check	Difference	
20-29	$25, 0.547 \pm 0.053$	$25, 0.547 \pm 0.053$	0	
30-39	$27, 0.632 \pm 0.153$	$27, 0.632 \pm 0.153$	0	
40-49	$25, 0.655 \pm 0.074$	$25, 0.655 \pm 0.074$	0	
		Type 1 diabetes	·	
	Diabetes (1999)	Integrity Check	Difference	
20-29	$172, 0.583 \pm 0.092$	$172, 0.583 \pm 0.092$	0	
30-39	$278, 0.632 \pm 0.147$	$278, 0.632 \pm 0.147$	0	
40-49	$178, 0.719 \pm 0.226$	$178, 0.719 \pm 0.226$	0	
		p-value		
	Diabetes (1999)	Integrity Check	Difference	
20-29	0.065	0.064	0.001	
30-39	0.859	0.856	0.003	
40-49	0.704	0.697	0.007	
Note: Data are sample size, mean \pm s rank-sum test after linear adjustment	standard deviation and p- for covariance with age	values. P-values are fi	rom Wilcoxon	

Table 2 Variable	Men, Com	Men, Common Carotid Artery IMT (mm)			
		Nondiabetic			
	Diabetes (1999)	Integrity Check	Difference		
20-29	$21, 0.648 \pm 0.098$	$21,0.648\pm 0.098$	0		
30-39	$25, 0.657 \pm 0.076$	$25, 0.657 \pm 0.076$	0		
40-49	$30, 0.741 \pm 0.094$	30, 0.741 ± 0.094	0		
		Type 1 diabetes			
	Diabetes (1999)	Integrity Check	Difference		
20-29	$125, 0.636 \pm 0.059$	$125, 0.636 \pm 0.059$	0		
30-39	$350, 0.684 \pm 0.083$	$350, 0.684 \pm 0.083$	0		
40-49	$211, 0.745 \pm 0.104$	$211, 0.745 \pm 0.104$	0		
		p-value			
	Diabetes (1999)	Integrity Check	Difference		
20-29	0.971	0.973	0.002		
30-39	0.103	0.104	0.001		
40-49	0.925	0.924	0.001		
Note: Data are sample size, mea rank-sum test after linear adjusti	$n \pm standard$ deviation and p nent for covariance with ag	o-values. P-values are f e.	rom Wilcoxon		

Table D: Comparison of Nondiabetic Control Subject Table Values Computed in Integrity Check to Reference Article Values (cont.)

Table 2 Variable	Men, Internal Carotid Artery IMT (mm)			
	Nondiabetic			
	Diabetes (1999)	Integrity Check	Difference	
20-29	$21, 0.588 \pm 0.085$	$21,0.588\pm 0.085$	0	
30-39	$25, 0.645 \pm 0.104$	$25, 0.645 \pm 0.104$	0	
40-49	$30, 0.741 \pm 0.147$	$30, 0.741 \pm 0.147$	0	
		Type 1 diabetes		
	Diabetes (1999)	Integrity Check	Difference	
20-29	$125, 0.629 \pm 0.083$	$125, 0.629 \pm 0.083$	0	
30-39	$350, 0.684 \pm 0.114$	$350, 0.684 \pm 0.114$	0	
40-49	$211, 0.806 \pm 0.261$	$211, 0.806 \pm 0.261$	0	
	p-value			
	Diabetes (1999)	Integrity Check	Difference	
20-29	0.032	0.032	0	
30-39	0.071	0.071	0	
40-49	0.437	0.437	0	
Note: Data are sample size, mean \pm standard deviation and p-values. P-values are from Wilcoxon rank-sum test after linear adjustment for covariance with age.				

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Univariate Correlation Data. Table 3 of the 1999 *Diabetes* article presents univariate Spearman correlations of various clinical characteristics with average maximum wall thickness for the common and internal carotid arteries. Variables summarized in this follow-up table (Table 3. Spearman correlation coefficients of clinical characteristics with average maximum IMT) are taken from the EDICIMT1 analysis dataset created for this study. Table E lists the variables used for replication of the Table 3 variables.

Table 3 Variable	Variables Used in Replication
Sex	sex
Attained age (years)	att_age
Attained duration of type 1 diabetes (months)	att_durn
Height (cm)	height
BMI (kg/m ²)	bmi
WHR	nwst_hip
sBP (mmHg)	sbp
dBP (mmHg)	dbp
Total cholesterol (mg/dl)	tchol
HDL cholesterol (mg/dl)	hdl
LDL cholesterol (mg/dl)	ldl
Triglycerides (mg/dl)	trig
Smoking (total pack-years)	cum_smok
Insulin dose (U • kg ⁻¹ • day ⁻¹)	std_ins
HbA _{1c}	hba1c
Mean HbA _{1c} during the DCCT	avg_a1c
Framingham risk score	fscore
AER (mg/24 h)	aer
GFR	gfr

Table E: Variables Used to Replicate Table 3

In Table F, we compare the follow-up results calculated from the archived dataset to the results published in the 1999 *Diabetes* article. As Table F shows, the results obtained from the archived data are the same as those in the published tabulations, with a few minor exceptions.

Table 3 Variable	Common Carotid Artery				
	Diabetes (1999)	Difference			
	M: 0.46, <0.001	M: 0.46, <0.001	M: 0		
Attained age (years)	F: 0.42, <0.001	F: 0.42, <0.001	F: 0		
Attained duration of type 1 diabetes	M: 0.13, <0.001	M: 0.13, <0.001	M: 0		
(months)	F: 0.08, 0.04	F: 0.08, 0.04	F: 0		
	M: 0.17, <0.001	M: 0.17, <0.001	M: 0		
Height (cm)	F: 0.12, 0.003	F: 0.12, 0.002	F: 0, 0.001		
$\mathbf{D}\mathbf{M}(\mathbf{l},\mathbf{z}/m^2)$	M: 0.15, <0.001	M: 0.15, <0.001	M: 0		
BMI (kg/m)	F: 0.06, 0.12	F: 0.06, 0.11	F: 0, 0.01		
МЛІР	M: 0.22, <0.001	M: 0.22, <0.001	M: 0		
WHK	F: 0.01, 0.89	F: 0.01, 0.90	F: 0, 0.01		
aDD (mmHa)	M: 0.21, <0.001	M: 0.21, <0.001	M: 0		
sBP (mmHg)	F: 0.14, <0.001	F: 0.14, <0.001	F: 0		
	M: 0.11, 0.004	M: 0.11, 0.003	M: 0, 0.001		
dBP (mmHg)	F: 0.07, 0.08	F: 0.07, 0.08	F: 0		
$T_{1} = \{1, 1, 2, 3, 4, 3, 5, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 2, 3, 4, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,$	M: 0.21, <0.001	M: 0.21, <0.001	M: 0		
Total cholesterol (mg/dl)	F: 0.09, 0.04	F: 0.09, 0.03	F: 0, 0.01		
UDL shalastanal (ma/dl)	M: -0.03, 0.37	M: -0.03, 0.36	M: 0, 0.01		
HDL cholesterol (mg/dl)	F: 0.00, 0.96	F: 0.00, 0.96	F: 0		
LDL shalastaral (mg/dl)	M: 0.24, <0.001	M: 0.24, <0.001	M: 0		
LDL cholesterol (mg/dl)	F: 0.08, 0.06	F: 0.08, 0.06	F: 0		
Triglyaaridas (mg/dl)	M: 0.07, 0.06	M: 0.07, 0.05	M: 0, 0.01		
The second secon	F: 0.06, 0.14	F: 0.06, 0.14	F: 0		
Smalting (total neals years)	M: 0.21, <0.001	M: 0.21, <0.001	M: 0		
Shloking (total pack-years)	F: 0.15, <0.001	F: 0.15, <0.001	F: 0		
Insulin doss $(U \bullet ka^{-1} \bullet day^{-1})$	M: -0.06, 0.10	M: -0.06, 0.09	M: 0, 0.01		
Insum dose (0 • kg • day)	F: -0.08, 0.06	F: -0.08, 0.05	F: 0, 0.01		
	M: 0.07, 0.07	M: 0.07, 0.06	M: 0, 0.01		
HUA _{1c}	F: -0.02, 0.62	F: -0.02, 0.62	F: 0		
Moon Uh A during the DCCT	M: 0.06, 0.12	M: 0.06, 0.11	M: 0, 0.01		
Mean HDA _{1c} during the DCC I	F: -0.11, 0.008	F: -0.11, 0.007	F: 0, 0.001		
Framingham right goorg	M: 0.47, <0.001	M: 0.47, <0.001	M: 0		
Framingham fisk score	F: 0.42, <0.001	F: 0.42, <0.001	F: 0		
A E B (ma/24 h)	M: 0.10, 0.01	M: 0.10, 0.01	M: 0		
AER (Ing/24 II)	F: -0.01, 0.76	F: -0.01, 0.75	F: 0, 0.01		
CEP	M: -0.01, 0.76	M: -0.01, 0.77	M: 0, 0.01		
	F: -0.05, 0.25	F: -0.05, 0.24	F: 0, 0.01		
Note: Data are Spearman correlations and p-values.					

Table F: Comparison of Univariate Correlation Table Values Computed in Integrity Check to Reference Article Values

Table 3 Variable	Internal Carotid Artery				
	Diabetes (1999)	Difference			
Attained are (warma)	M: 0.38, <0.001	M: 0.38, <0.001	M: 0		
Attained age (years)	F: 0.35, <0.001	F: 0.35, <0.001	F: 0		
Attained duration of type 1 diabetes	M: 0.12, 0.002	M: 0.12, 0.002	M: 0		
(months)	F: 0.12, 0.004	F: 0.12, 0.004	F: 0		
Height (am)	M: 0.05, 0.19	M: 0.05, 0.19	M: 0		
Height (cm)	F: -0.04, 0.36	F: -0.04, 0.36	F: 0		
$\mathbf{DML}(\mathbf{l}_{ra}/\mathbf{m}^2)$	M: 0.09, 0.02	M: 0.09, 0.02	M: 0		
Divit (kg/iii)	F: 0.02, 0.56	F: 0.02, 0.56	F: 0		
W/HD	M: 0.14, <0.001	M: 0.14, <0.001	M: 0		
WIIK	F: 0.01, 0.85	F: 0.01, 0.85	F: 0		
aDD (mmHa)	M: 0.14, <0.001	M: 0.14, <0.001	M: 0		
SBP (mmrg)	F: 0.11, 0.007	F: 0.11, 0.007	F: 0		
dDB (mmHz)	M: 0.13, <0.008	M: 0.13, 0.008	M: 0		
dbp (mmrg)	F: 0.07, 0.09	F: 0.07, 0.08	F: 0, 0.01		
Total shalastaral (mg/dl)	M: 0.23, <0.001	M: 0.23, <0.001	M: 0		
Total cholesterol (ling/dl)	F: 0.13, 0.002	F: 0.13, 0.001	F: 0, 0.001		
UDL shalastaral (ma/dl)	M: 0.00, 0.92	M: 0.00, 0.92	M: 0		
HDL cholesterol (hig/di)	F: 0.03, 0.41	F: 0.03, 0.41	F: 0		
I DL abalastaral (mg/dl)	M: 0.26, <0.001	M: 0.26, <0.001	M: 0		
LDL cholesteror (hig/di)	F: 0.13, 0.001	F: 0.13, 0.001	F: 0		
Triglycorides (mg/dl)	M: 0.05, 0.19	M: 0.05, 0.18	M: 0, 0.01		
(ling/di)	F: 0.00, 0.94	F: 0.00, 0.94	F: 0		
Smaking (total nack years)	M: 0.09, 0.02	M: 0.09, 0.02	M: 0		
Shloking (total pack-years)	F: 0.16, <0.001	F: 0.16, <0.001	F: 0		
Insulin dose $(U \bullet ka^{-1} \bullet day^{-1})$	M: -0.10, 0.02	M: -0.10, 0.01	M: 0, 0.01		
Insum dose (0 ° kg ° day)	F: -0.05, 0.20	F: -0.05, 0.20	F: 0		
	M: 0.02, 0.56	M: 0.02, 0.56	M: 0		
HUA _{1c}	F: 0.02, 0.68	F: 0.02, 0.68	F: 0		
Mean HbA during the DCCT	M: -0.02, 0.57	M: -0.02, 0.56	M: 0, 0.01		
Mean HDA _{1c} during the DCC I	F: -0.07, 0.07	F: -0.07, 0.06	F: 0, 0.01		
Framingham risk soore	M: 0.38, <0.001	M: 0.38, <0.001	M: 0		
	F: 0.35, <0.001	F: 0.35, <0.001	F: 0		
AEP (mg/24 h)	M: 0.07, 0.09	M: 0.07, 0.09	M: 0		
AER (IIIg/24 II)	F: 0.01, 0.85	F: 0.01, 0.84	F: 0, 0.01		
GER	M: 0.02, 0.54	M: 0.02, 0.54	M: 0		
	F: -0.02, 0.65	F: -0.02, 0.65	F: 0		
Note: Data are Spearman correlations and p-values.					

Table F: Comparison of Univariate Correlation Table Values Computed in Integrity Check to Reference Article Values (cont.)

Norma Pugh February 12, 2008

Multivariate Analyses Data. Table 4 of the 1999 *Diabetes* article presents multiple linear regression analyses. Variables summarized in this follow-up table (Table 4. Summary of multiple regression models for common carotid IMT and internal carotid IMT) are taken from the EDICIMT1 analysis dataset created for this study. Selected results were spot-checked and found to be consistent with the published results.

Treatment Group Difference Data. Table 5 of the 1999 *Diabetes* article presents IMT by sex, age, and treatment group. Variables summarized in this follow-up table (Table 5. Average maximum IMTs by age and randomized treatment assignment) are taken from the EDICIMT1 analysis dataset created for this study. Selected results were spot-checked and found to be consistent with the published results. There were, in fact, no differences in any of the means or standard deviations.

The repository has high confidence in the integrity of the EDIC One Year IMT analysis datasets.

Notes

1. Both analysis datasets related to the EDIC One Year Intima-media Wall Thickness (IMT) follow-up were examined in this replication analysis: EDICIMT1X for the nondiabetic cohort and EDICIMT1 for the diabetic cohort.

References

 The EDIC Research Group, Effect of Intensive Diabetes Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes Interventions and Complications, Diabetes, February 1999; Volume 48: 383-390.

ATTACHMENT 1

Full Text of Article

The EDIC Research Group, Effect of Intensive Diabetes Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes Interventions and Complications, Diabetes, February 1999; Volume 48: 383-390.

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Effect of Intensive Diabetes Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes Interventions and Complications

Epidemiology of Diabetes Interventions and Complications (EDIC) Research Group

The Epidemiology of Diabetes Interventions and Complications (EDIC) is a multicenter longitudinal observational study of the Diabetes Control and Complications Trial (DCCT) cohort. One of the major objectives of EDIC is to study the development and progression of atherosclerotic cardiovascular disease in type 1 diabetes. In this study, we evaluated the role of cardiovascular risk factors and antecedent therapy in the DCCT on carotid intima-media wall thickness (IMT) in type 1 diabetes. At ~18 months after the end of the DCCT, high-resolution B-mode ultrasonography was used to assess the carotid arteries of 1,325 patients with type 1 diabetes, 19-51 years of age, with duration of diabetes ranging from 6.3 to 26.1 years. An age- and sex-matched nondiabetic population (n = 153) was studied with the same protocol. The ultrasound protocol was carried out in 28 EDIC clinics by centrally trained and certified sonographers using one of three scanning systems. Determination of IMT from videotaped images was performed by a single reader at the Central Ultrasound Reading Unit. Univariate associations with greater IMT were strongest for older age and longer diabetes duration, greater waist-to-hip ratio (men only), higher blood pressure, higher LDL cholesterol, and smoking. The DCCT therapy group (intensive versus conventional) and HbA_{1c}, measured at the time of the ultrasound or the mean HbA_{1c} during the DCCT, were not significantly related to IMT. Multivariate analyses suggest that age, height, smoking, and BMI were the major predictors of common carotid IMT, whereas age, smoking, and LDL cholesterol predicted internal carotid IMT. There were significant differences between the IMT values of the internal carotid artery in the EDIC male cohort and similarly aged male nondiabetic control subjects. There were no significant differences between the IMT values in the EDIC female cohort and similarly aged female nondiabetic control

subjects. At this point in the planned 10-year follow-up of the DCCT cohort, neither intensive therapy nor HbA_{1c} level appears to influence the early signs of atherosclerosis. Traditional risk factors, including age, smoking, and LDL cholesterol, were related to IMT. As the cohort is only now entering the age interval during which rapid progression and clinical expression of atherosclerosis are expected, further follow-up will help to determine the role of hyperglycemia, and its interaction with other risk factors, on the development of atherosclerosis. *Diabetes* 48:383–390, 1999

ype 1 diabetes is well recognized as a risk factor for early cardiovascular disease (CVD), leading to a more than 10-fold increase in risk in young adults (1–3) and greatly reducing the sex differential in CVD seen in the general population. The mechanism that underlies this effect of diabetes is unclear, and whether type 1 diabetes initiates the atherosclerotic process early or merely hastens the process once started is controversial (2,4). Furthermore, few studies have delineated the risk factors for CVD in type 1 diabetes, although data confirm a strong relationship with renal disease (5), particularly in men (6). Other risk factors of interest include the lipoprotein profile, blood pressure (BP), fibrinogen level, waist-to-hip ratio (WHR), and depressive symptomatology, particularly in women (6).

Carotid ultrasonography, measuring both the presence of stenosis and intima-media wall thickness (IMT), has provided a powerful noninvasive technique to determine atherosclerosis (7-13). IMT has been extensively used as an outcome measure in clinical trials (14–20). Strong correlations between IMT and cardiovascular risk factors and coronary artery disease (CAD) have been demonstrated in the general population (21,22). However, few studies have been performed in diabetes, particularly in type 1 diabetic populations. The Atherosclerosis Risk in Communities Study demonstrated greater IMT in both diabetic subjects (mainly those with type 2 diabetes) and nondiabetic participants with moderate hyperglycemia compared with subjects with normal glucose levels (23). Similarly, IMT was demonstrated to be increased in type 1 and type 2 diabetic Japanese subjects compared with those without diabetes (24). Two studies from Italy (25,26) have also reported increased IMT and a higher presence of carotid plaques and stenoses in type 2 diabetes (25). More relevant to the current study, young type 1 diabetic subjects (10-25 years of age) in Japan have increased IMT compared with control subjects (27).

The EDIC Research Group (see APPENDIX) is sponsored by the Division of Diabetes, Endocrinology, and Metabolic Diseases of the National Institute of Diabetes and Digestive and Kidney Diseases through research contracts and the General Research Center Program, National Center for Research Resources, the National Institutes of Health.

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AER, albumin excretion rate; BP, blood pressure; CAD, coronary artery disease; CVD, cardiovascular disease; dBP, diastolic blood pressure; DCCT, Diabetes Control and Complications Trial; ECG, electrocardiogram; EDIC, Epidemiology of Diabetes Interventions and Complications; GFR, glomerular filtration rate; IMT, intima-media wall thickness; sBP, systolic blood pressure; WHR, waist-to-hip ratio.

Data concerning determinants of IMT in type 1 diabetes are also limited. In a Japanese population (27), lipoproteins, BP, and HbA_{1c} were not related to IMT in 105 type 1 diabetic subjects. In contrast, in an even smaller study of type 1 diabetes (n = 31), age and HbA_{1c} predicted IMT (28). The role of hyperglycemia (or glycemic control) as a risk factor for CVD or atherosclerosis in type 1 diabetes is poorly understood. Recent data in type 1 diabetes suggest little, if any, effect of HbA_{1c} on CVD (6,8,29). In long-term follow-up of the original 102 subjects in the Stockholm Diabetes Intervention Study, the patients originally assigned to intensive therapy had decreased IMT of the left but not the right common carotid artery, compared with the patients originally assigned to conventional treatment (30).

To address the limited studies and contradictory results with regard to the course of CVD in type 1 diabetes and the role of risk factors for atherosclerosis in type 1 diabetes, carotid ultrasonography was performed on prior participants of the Diabetes Control and Complications Trial (DCCT) (31) who are taking part in a longitudinal follow-up study called the Epidemiology of Diabetes Interventions and Complications (EDIC) (31a). The DCCT cohort (31), which excluded patients with hypertension, hyperlipidemia, and known CAD at baseline, provides an excellent opportunity to examine a large population of patients with type 1 diabetes without obvious CVD risk at baseline who have had careful prospective measurements of many of the putative risk factors for CVD previously established in type 2 diabetes and in nondiabetic populations. The DCCT cohort also provides the opportunity to examine the effects of diabetes treatment on CVD. The randomized interventions during the DCCT might influence the development of CVD, either directly by altering glucose levels or indirectly by altering lipid levels, the development of nephropathy or other risk factors. In addition, other side effects of intensive therapy documented in the DCCT, such as weight gain and increased rates of hypoglycemia, might also affect CVD risk.

Here we present the results of carotid arterial sonographic measurements of 1,325 patients during the baseline data collection of the EDIC study. The primary aims of this baseline analysis are to 1) determine the feasibility of implementing the protocol in 28 sites; 2) compare the EDIC cohort results with normative data from individuals matched for age and sex; 3) examine the intercorrelations of IMT with demographic, clinical, and biochemical covariates that have been reported previously by other investigators; and 4) determine whether the antecedent DCCT therapy (intensive versus conventional) and the different levels of glycemia achieved resulted in differences in IMT.

RESEARCH DESIGN AND METHODS

Subject population. The original cohort of the DCCT consisted of 1,441 men and women who were 13–40 years of age and had type 1 diabetes for 1–15 years at entry (31). They entered the DCCT between 1983 and 1989 and were studied for an average of 6.5 years. A total of 730 patients were randomly assigned to conventional diabetes treatment and 711 to intensive diabetes treatment. In 1993, the DCCT was stopped because of evidence of a powerful salutary effect of intensive therapy on retinal, renal, and neurological long-term complications (31). At study close-out, DCCT subjects were informed of and invited to join EDIC, a multicenter longitudinal observational study. Of the 1,425 living members of the original cohort, 1,388 (96%) elected to participate in some or all aspects of EDIC. The carotid ultrasound protocol was carried out in 1,325 patients (92% of the original DCCT cohort) as part of the EDIC baseline examination. Table 1 presents the clinical characteristics of these 1,325 patients at EDIC baseline, stratified by original treatment group and sex.

TABLE 1

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	Women			Men		
	Intensive	Conventional	P value	Intensive	Conventional	<i>P</i> value
п	321	313	_	340	351	_
Attained age (years)	35 ± 7	34 ± 7	0.042	36 ± 7	36 ± 7	0.912
Attained duration of type 1 diabetes (months)	168 ± 58	170 ± 61	0.722	167 ± 59	159 ± 55	0.096
Height (cm)	165 ± 6	165 ± 6	0.682	178 ± 7	179 ± 7	0.076
BMI (kg/m ²)	26.5 ± 4.6	25.1 ± 3.6	<.001	26.8 ± 4.2	25.9 ± 3.2	0.016
BMI >27 (%)	39.7	27.5	0.001	41.2	34.5	0.069
Natural WHR	0.76 ± 0.07	0.76 ± 0.07	0.731	0.88 ± 0.08	0.87 ± 0.09	0.101
sBP (mmHg)	114 ± 12	114 ± 13	0.739	119 ± 11	120 ± 12	0.186
dBP (mmHg)	74 ± 9	72 ± 10	0.101	77 ± 9	77 ± 8	0.446
Hypertensive (%)	9.7	12.9	0.210	21.2	17.6	0.240
Total cholesterol (mg/dl)	188 ± 36	188 ± 39	0.495	187 ± 35	184 ± 36	0.166
HDL cholesterol (mg/dl)	59 ± 14	59 ± 14	0.737	49 ± 13	50 ± 12	0.742
LDL cholesterol (mg/dl)	113 ± 29	113 ± 32	0.542	119 ± 30	115 ± 31	0.145
Triglycerides (mg/dĺ)	84 ± 74	82 ± 73	0.427	98 ± 77	97 ± 78	0.756
AER (mg/24 h)	21.1 ± 63.2	63.5 ± 313.9	0.003	28.6 ± 112.9	46.1 ± 123.2	0.038
AER 40 mg/24 h (%)	6.9	16.0	<.001	7.4	17.9	<0.001
GFR	117.5 ± 23.5	119.3 ± 26.1	0.069	115.3 ± 19.7	116.1 ± 24.8	0.335
Insulin dose (U · kg ⁻¹ · day ⁻¹)	0.63 ± 0.21	0.64 ± 0.19	0.801	0.67 ± 0.24	0.65 ± 0.20	0.269
Cigarette smoking (current) (%)	19.7	18.1	0.605	19.3	18.4	0.750
HbA _{1c}	7.9 ± 1.4	8.2 ± 1.5	0.009	7.9 ± 1.2	8.4 ± 1.3	<0.001
Mean HbA _{1c} during DCCT	7.3 ± 0.9	9.0 ± 1.4	<0.001	7.2 ± 0.9	9.0 ± 1.1	<0.001
Framingham score	0.018 ± 0.025	0.017 ± 0.028	0.037	0.039 ± 0.038	0.037 ± 0.035	0.704

Data are means ± SD, *n*, or %. Baseline is that at EDIC baseline (1994–1995) after an average of 6.5 years of intensive treatment in DCCT. Hypertensive is defined as sitting sBP 140 mmHg and/or dBP 90 mmHg or the use of antihypertensive medication. Framingham score is defined as described by Anderson et al. (An Updated Coronary Risk Profile. *Circulation* 83:355–365, 1991).

Ultrasonography and image analysis. Carotid ultrasonography was performed between June 1994 and April 1995 (1–2 years after the close of the DCCT) at 28 clinical centers using one of three machines (Toshiba, American Medical Systems, Tustin, CA; ATL Ultra Mark 9; Advanced Technology Laboratory, Bothell, WA; and Acuson XT 128; Mt. View, CA). The criteria used to select these machines included the following:

- · Accurate delineation of near and far wall boundaries,
- Accurate plaque detection and sizing,
- · Simultaneous Doppler, preferably with color,
- · Detection and quantification of early subintimal change,
- Accurately reproduced images on videotape using S-VHS recorders.

The EDIC Ultrasound Scanning Protocol was adapted from procedures used in the Cardiovascular Health Study (7), the Insulin Resistance Atherosclerosis Study (32), and the Mexican-American Heart Study (33).

The imaging protocol involved obtaining a single longitudinal lateral view of the distal 10 mm of the right and left common carotid arteries and three longitudinal views in different imaging planes of each internal carotid artery. The internal carotid artery was defined as including both the carotid bulb, identified by the loss of the parallel wall present in the common carotid artery, and the 10mm segment of the internal carotid artery distal to the tip of the flow divider that separates the external and internal carotid arteries (Fig. 1).

Centrally trained and certified sonographers conducted the studies. Studies were recorded on S-VHS tapes and sent weekly to the Central Ultrasound Reading Unit. The maximum IMT of the common carotid artery was defined as the mean of the maximum IMT for near and far walls in both right and left sides. The maximum IMT of the internal carotid artery was defined in the same way, and the results from the three scans were averaged (anterior, lateral, and posterior views on both sides).

Normative data. During the first year of the EDIC, each of the 28 clinics performed six carotid ultrasounds on nondiabetic subjects who were between the ages of 20 and 50 years with no history of CVD, hypertension, or stroke (n = 153). The mean age (35.3 ± 8.5) and percentage of women (49.7%) were not significantly different than those of the EDIC cohort.

Other procedures. On the anniversary of enrolling in the DCCT, each EDIC subject has a standardized annual history and physical examination, including a detailed evaluation of overall health, diabetes management, occurrence of diabetic complications, development of new disease, and medications used. Annual evaluations also include resting electrocardiograms (ECGs), Doppler ultrasound measurements of ankle/arm BP ratios, and arm BPs. Serum creatinine and HbA_{1c} are determined as they were in the DCCT (34). Lipid profiles and 4-h urine collections for measurement of albumin excretion rate and creatinine clearance are obtained in alternate years using the same methods as in the DCCT (35).

Statistical analysis. Quality scores for carotid ultrasound scans were based on the number of lines visualized from the eight views. The proportion of lines with quality scores of good or excellent were compared across the 28 clinics.

To evaluate the possible association with other covariates, multiple linear regression models were fit to the average maximum IMTs of the common and internal carotid arteries. Both models were fit by ordinary least squares, but a reciprocal transformation had to be applied to internal IMT to obtain a homoscedastic and approximately normal residual distribution. The semipar-

tial R^2 for each variable measures the increase in R^2 obtained by introducing that variable into a model already including all the others.

Wilcoxon rank-sum tests were used to make separate comparisons of the distributions of common IMT and internal IMT between EDIC patients and nondiabetic control subjects within strata defined by age and sex. Separate linear models for each segment were used to adjust for covariance with age before testing; EDIC patients and control subjects were adjusted jointly but stratified by sex. Comparison of IMTs between the DCCT randomized treatment groups was made using the Wei-Lachin test of stochastic ordering (36), which has greatest power against alternatives in which both segments tend to be thicker in the same treatment group. Tests were stratified by sex and decade of age, and an overall test adjusted for between-strata differences (36) was also performed. Based on the results of the multiple linear regression models, the EDIC patients' IMT values were first adjusted for covariance with age and cumulative pack-years of cigarette smoking. All *P* values are reported at their nominal levels, without adjustment for multiple comparisons.

RESULTS

Data quality. Baseline reproducibility analyses of 140 replicate measures resulted in absolute mean differences of 0.04 mm for both the common carotid and the intimal carotid. Intraclass correlation between original and re-readings of maximum wall thickness were 0.71 and 0.89 for the common and for internal carotid arteries, respectively.

Carotid artery data by clinic were generally of uniform high quality. The percentage of scans of the common carotid artery in which all six lines were legible and graded as good or excellent ranged from 90 to 100% across clinics. Visualization of the internal carotid artery was less uniform, with the percentage of scan with all six lines legible ranging from 15 to 94% of scans. Scans that did not meet the criteria (mainly segments of the internal carotid) had only four or five lines legible. A process for continuous quality control was instituted so that information was fed back to each center in an effort to improve scan quality.

Comparison of nondiabetic control subjects with EDIC subjects. Table 2 presents the IMT data including the average maximum of the common and internal carotid arteries stratified by sex and age decades for nondiabetic and type 1 diabetic subjects. A Wilcoxon test of the difference between sexes for both diabetic and nondiabetic groups indicates highly significant (P 0.0001) differences, with men having greater IMT mean thickness. Adjusting for covariance with height reduces but does not eliminate the significance (P =



FIG. 1. Schematic drawing of the carotid artery: the bifurcation of the common into the internal and external carotid arteries, the location of the segments scanned in the EDIC, and the six lines to be measured in the sonographic images.

TABLE 2	
$Comparison \ of \ IMT \ in \ nondiabetic \ subjects \ with \ EDIC$	type 1 diabetes cohort

		Women					Men				
	Ν	Nondiabetic		Type 1 diabetes		N	Nondiabetic		Type 1 diabetes		
Age (years)	n	IMT (mm)	n	IMT (mm)	Р	n	IMT (mm)	n	IMT (mm)	Р	
Common carotid artery											
20-29	25	0.603 ± 0.062	172	0.616 ± 0.073	0.471	21	0.648 ± 0.098	125	0.636 ± 0.059	0.971	
30-39	27	0.663 ± 0.074	278	0.657 ± 0.081	0.453	25	0.657 ± 0.076	350	0.684 ± 0.083	0.103	
40-49	25	0.704 ± 0.078	178	0.696 ± 0.079	0.622	30	0.741 ± 0.094	211	0.745 ± 0.104	0.925	
Internal carotid artery											
20–29	25	0.547 ± 0.053	172	0.583 ± 0.092	0.065	21	0.588 ± 0.085	125	0.629 ± 0.083	0.032	
30–39	27	0.632 ± 0.153	278	0.632 ± 0.147	0.859	25	0.645 ± 0.104	350	0.684 ± 0.114	0.071	
40–49	25	0.655 ± 0.074	178	0.719 ± 0.226	0.704	30	0.741 ± 0.147	211	0.806 ± 0.261	0.437	

Data are means \pm SD or *n*. Nondiabetic data were obtained from six normal subjects without diabetes at each of the 29 clinics using the EDIC ultrasound scanning protocol and read at the Central Ultrasound Reading Unit. Data exclude six EDIC patients who were 19 years old when scanned and five who were aged 50 or 51 years old. *P* values are from Wilcoxon rank-sum test after linear adjustment for covariance with age. *P* < 0.0001 for both common and internal IMT, type 1 men versus type 1 women; *P* = 0.59 for common IMT, *P* = 0.004 for internal IMT, type 1 men versus nondiabetic men; *P* = 0.89 for common IMT, *P* = 0.11 for internal IMT, type 1 women versus nondiabetic women.

0.0007). A test of trend of the increasing maximum wall thickness over the age decades is significant in all strata (e.g., nondiabetic men, type 1 diabetic men, etc.).

Comparing the nondiabetic population with the sexmatched control subjects revealed that overall the type 1 diabetic men had greater IMT thickness for the internal carotid (0.713 ± 0.184 vs. 0.664 ± 0.130 mm, P = 0.004) but not for the common carotid (0.694 ± 0.10 vs. 0.686 ± 0.10 mm, P = 0.59). Within the different age-groups, only the youngest group of men had significantly different internal carotid IMT than their age-matched control subjects (0.629 vs. 0.588, P =0.03). The type 1 diabetic women's mean IMT thickness was similar to the nondiabetic women's for both the common and internal carotid.

Univariate correlation. Table 3 presents univariate Spearman correlations of various clinical characteristics with average maximum wall thickness for the common and internal carotid arteries. Attained age and Framingham risk scores were the variables most strongly associated with IMT in both the common and internal carotid arteries. Attained duration of type 1 diabetes had a weaker, but statistically significant, association with wall thickness that, in men, persisted after adjusting for age. Current HbA_{1c} was not correlated with wall thickness; however, HbA_{1c} at the time of carotid studies did not reflect glycemic exposure during the DCCT, since all subjects were encouraged to adopt intensive therapy at the end of the DCCT. Therefore, correlations between mean HbA_{1c} during the DCCT and IMTs were also computed. Mean HbA_{1c} during the DCCT was not significantly correlated with either common or internal carotid IMT among men or with internal IMT among women. The correlation of mean HbA_{1c} with common IMT in women, while nominally significant, was weak (r = -0.11, P = 0.008) and indicated an inverse relationship. Total and LDL cholesterol levels were correlated with wall thickness, but HDL cholesterol and triglyceride levels were not. Systolic blood pressure (sBP) was associated with IMT in both men and women, and diastolic blood pressure (dBP) was associated with IMT in men only. WHR correlated with IMT in men, but not women, while total packyears of cigarette smoking was correlated with both common and internal carotid IMT in both sexes. Correlations were similar in the two treatment groups.

Multivariate analyses. Table 4 shows the multiple linear regression analysis. Apart from age, the major predictors of common carotid IMT in women were height, glomerular filtration rate (GFR), BMI, and smoking, while in men, height, smoking, BMI, sBP, duration of type 1 diabetes, and average LDL cholesterol during the DCCT were independent predictors. Overall, 29 and 20% of the variance was explained in men and women, respectively. For internal carotid IMT, age, sBP, and LDL were the major predictors in both sexes, and smoking had a significant but reduced effect compared with the model for the common carotid, especially in men. GFR was also a predictor in men. Again, more of the variance was explained in men (24%) than in women (17%).

Tests for treatment group difference. Table 5 presents IMT by sex, age, and treatment group. There are no strata in which intensive treatment group patients had significantly different age- and smoking-adjusted wall thicknesses than conventional treatment group patients. The difference between intensive and conventional treatment from the N-weighted test of stochastic ordering for the combined strata was not statistically significant (P = 0.39).

DISCUSSION

The current report confirms the feasibility of using carotid ultrasonography in a large multicenter study with excellent reproducibility for common carotid and acceptable reproducibility for internal carotid IMT measurements (37,38). The quality of scans was good or excellent for the vast majority of centers, especially for the common carotid artery.

There was no significant difference in mean IMT between EDIC subjects and age- and sex-matched control subjects, except for internal carotid IMT among men. Because two major determinants of IMT are age and BP, the similarity in IMT may reflect the young age of the EDIC subjects and the

TABLE 3

Spearman correlation coefficients of clinical characteristics with average maximum IMT

	Sex	Common caro	tid artery	Internal carotid artery	
		Correlation	Pvalue	Correlation	<i>P</i> value
Attained age (years)	М	0.46	<0.001	0.38	<0.001
	F	0.42	<0.001	0.35	<0.001
Attained duration of type 1 diabetes (months)	М	0.13	<0.001	0.12	0.002
31	F	0.08	0.04	0.12	0.004
Height (cm)	Μ	0.17	<0.001	0.05	0.19
• • •	F	0.12	0.003	-0.04	0.36
BMI (kg/m ²)	Μ	0.15	<0.001	0.09	0.02
-	F	0.06	0.12	0.02	0.56
WHR	Μ	0.22	< 0.001	0.14	<0.001
	F	0.01	0.89	0.01	0.85
sBP (mmHg)	Μ	0.21	<0.001	0.14	<0.001
	F	0.14	<0.001	0.11	0.007
dBP (mmHg)	Μ	0.11	0.004	0.13	<0.008
	F	0.07	0.08	0.07	0.09
Total cholesterol (mg/dl)	Μ	0.21	<0.001	0.23	<0.001
	F	0.09	0.04	0.13	0.002
HDL cholesterol (mg/dl)	М	-0.03	0.37	0.00	0.92
	F	0.00	0.96	0.03	0.41
LDL cholesterol (mg/dl)	Μ	0.24	<0.001	0.26	<0.001
	F	0.08	0.06	0.13	0.001
Triglycerides (mg/dl)	Μ	0.07	0.06	0.05	0.19
	F	0.06	0.14	0.00	0.94
Smoking (total pack-years)	Μ	0.21	< 0.001	0.09	0.02
	F	0.15	< 0.001	0.16	<0.001
Insulin dose (U \cdot kg ⁻¹ \cdot day ⁻¹)	Μ	-0.06	0.10	-0.10	0.02
	F	-0.08	0.06	-0.05	0.20
HbA _{1c}	Μ	0.07	0.07	0.02	0.56
	F	-0.02	0.62	0.02	0.68
Mean HbA _{1c} during the DCCT	Μ	0.06	0.12	-0.02	0.57
	F	-0.11	0.008	-0.07	0.07
Framingham risk score	Μ	0.47	<0.001	0.38	<0.001
C C	F	0.42	< 0.001	0.35	<0.001
AER (mg/24 h)	Μ	0.10	0.01	0.07	0.09
	F	-0.01	0.76	0.01	0.85
GFR	Μ	-0.01	0.76	0.02	0.54
	F	-0.05	0.25	-0.02	0.65

initial exclusion of hypertensive subjects in the DCCT. These data are in contrast to those of Yamasaki et al. (27), who reported significantly greater IMT for 10- to 25-year-old type 1 diabetic subjects compared with nonmatched control subjects, despite the smaller number of type 1 diabetic patients and control subjects (<10% and 30%, respectively) in the Japanese study. The discrepancy between the Japanese and EDIC studies may be secondary to the unmatched control group in the Japanese study, racial differences in the pathogenesis of atherosclerosis, or other factors. The finding of increased IMT in type 2 diabetic subjects (23,26) may be secondary to older age or the high prevalence of other CVD risk factors found in type 2 diabetes that were not present in the EDIC population.

Another possible explanation of the lack of a marked difference in IMT in our type 1 diabetic subjects is the possibility that IMT does not reflect the aspects, or stages, of atherosclerosis that are enhanced or exacerbated by type 1 diabetes. Although IMT is thought to be an intermediate biological marker of atherosclerosis and correlates with the presence of plaque or clinical events (39), it does not provide does it directly measure the hemostatic (and possibly inflammatory) disturbances that may play a vital role in atherosclerotic disease and that are disturbed in diabetes, such as abnormal fibrinogen levels (40) and platelet function (41). The true predictive power of IMT for future events is uncertain, and the data supporting a correlation with existent disease, while encouraging, are still limited (21,22,39,42,43). These potential limitations, however, do not negate the value of carotid ultrasonography in diabetes, which may throw more light on the pathogenesis of artherosclerotic disease generally, as well as on the specific enhanced risk in diabetes. Only careful follow-up with repeated sonography and assessment of clinical outcomes will allow definitive determination of the effect of diabetes on the overall natural history of atherosclerotic CVD.

a direct measure of occlusive disease or plaque stability. Nor

The current study has limitations, the major one being that the DCCT cohort was a selected trial population and not necessarily representative of type 1 subjects in the general population. However, the DCCT conventional treatment group was found to be generally comparable to a subset of a

TABLE 4

Summary of multiple linear regression models for common carotid IMT and 1/internal carotid IMT

	Common 1/Internal			nal		
	Slope	Р	Semipartial R ² (%)	Slope	Р	Semipartial R ² (%)
Women						
Age	0.00435	<0.001	11.95	-0.01176	<0.001	7.02
Type 1 diabetes duration (months)	0.00006	0.267	0.21	-0.00034	0.078	0.54
Height	0.00131	0.011	1.11	0.00188	0.315	0.18
Smoking (pack-years)	0.00076	0.062	0.60	-0.00367	0.013	1.07
BMI	0.00155	0.050	0.67	0.00122	0.670	0.03
sBP	0.00028	0.307	0.18	-0.00217	0.027	0.85
LDL cholesterol (DCCT average)	0.00009	0.507	0.08	-0.00172	<0.001	2.00
GFR	0.00025	0.053	0.65	-0.00056	0.230	0.25
HbA _{1c} (DCCT average)	-0.00310	0.274	0.21	0.01192	0.247	0.23
Intensive therapy	-0.00522	0.515	0.07	0.02407	0.407	0.12
Men						
Age	0.00451	<0.001	9.35	-0.01167	<0.001	7.19
Type 1 diabetes duration (months)	0.00018	0.001	1.53	-0.00046	0.006	1.13
Height	0.00161	<0.001	2.15	-0.00051	0.684	0.03
Smoking (pack-years)	0.00149	<0.001	2.52	-0.00263	0.015	0.89
BMI	0.00322	<0.001	2.06	-0.00131	0.612	0.04
sBP	0.00085	0.002	1.38	-0.00201	0.016	0.88
LDL cholesterol (DCCT average)	0.00033	0.009	1.03	-0.00220	<0.001	5.07
GFR	0.00025	0.072	0.49	-0.00105	0.012	0.94
HbA _{1c} (DCCT average)	0.00226	0.481	0.08	0.00894	0.350	0.13
Intensive therapy	-0.00985	0.241	0.21	0.01190	0.635	0.03

Total women in the common group is 20.5%; total women in the 1/internal group is 17.4%; total men in the common group is 29.4%; total men in the 1/internal group is 24.3%.

population-based type 1 diabetes cohort (44). A further limitation to the study was the exclusion at baseline of patients with hypertension and hyperlipidemia. As noted, the relatively young age of the EDIC cohort and the exclusions noted above may have reduced the occurrence of CVD and the ability to detect risk factor associations and differences in IMT at the current time. It should also be noted that the current analysis is essentially cross-sectional (albeit with a well-documented and randomized historical prospective measure of glycemic exposure) and is subject to the limitations of these studies, in particular, the absence of baseline (carotid) studies. Cardiovascular risk profile, however, did not differ by DCCT treatment group at baseline (31).

An increased number of CVD events is very likely to occur in the DCCT cohort during the 10 years of the EDIC. By the end of the 10-year follow-up, the mean age of the study population, the major predictor of CAD, will approach 43 years, and type 1 diabetes duration will average 22 years. Based on

TABLE 5 Average maximum IMTs by age and randomized treatment assignment

	Age (years)		Common c	arotid ar	Internal carotid artery			
Sex		n	Intensive	п	Conventional	Intensive	Conventional	Р
Women								
	20-29	78	0.626 ± 0.078	94	0.608 ± 0.068	0.576 ± 0.071	0.589 ± 0.107	0.809
	30-39	137	0.651 ± 0.086	141	0.663 ± 0.076	0.628 ± 0.129	0.636 ± 0.164	0.437
	40-49	102	0.698 ± 0.079	76	0.693 ± 0.080	0.704 ± 0.207	0.738 ± 0.250	0.954
Men								
	20–29	65	0.637 ± 0.062	60	0.635 ± 0.056	0.636 ± 0.091	0.622 ± 0.072	0.743
	30-39	167	0.680 ± 0.079	183	0.687 ± 0.086	0.684 ± 0.107	0.685 ± 0.120	0.807
	40-49	104	0.728 ± 0.084	107	0.761 ± 0.118	0.791 ± 0.211	0.820 ± 0.302	0.149
Combined								
(stratified-	adjusted)							0.388

Data are means \pm SD or *n*. *P* values are from the Wei-Lachin test of stochastic ordering after adjustment of IMT values for covariance with age and pack-years of cigarette smoking. The "combined" test aggregates the within-stratum results while accounting for the differences between age-groups and sex. Means and SDs were calculated from the unadjusted IMTs. The table excludes 11 patients who were <20 or >49 years old when tested.

estimates derived from previous studies (1–6), the prevalence of CAD, as manifested clinically and/or as detected by ECG or exercise tolerance testing, is likely to be 15–40%. It seems likely that we shall have sufficient power to determine the relationship of IMT to CVD events occurring over the subsequent 10 years.

Documented risk factors for increased IMT include age, hypertension, lipid profile, and smoking. Our data suggest that these factors relate to IMT in type 1 diabetes. Among established lipid risk factors, only LDL level correlated with IMT. Some risk factors appeared to affect IMT in the common carotid, while others affected the internal carotid wall. For example, in multivariate analyses, smoking was more strongly related to common carotid IMT than to internal carotid, while LDL cholesterol was stronger for the internal carotid.

The large sample size in the current study allowed the multivariate analyses to establish a variety of factors including smoking, sBP, GFR, BMI, and LDL cholesterol as predictors of IMT. Some studies in type 2 diabetes have confirmed some of these associations, e.g., BP (24–26) and hyperlipidemia (24), while other studies in type 2 diabetes have failed to find such associations (27,28), possibly owing to their small sample size. The relationship between GFR and IMT may reflect the recognized association between renal disease and CVD in type 1 diabetes (5,6).

The role of glycemic control (or glycemic level) is of particular interest. The few data available relating glycemia to CVD in type 1 diabetes are controversial. The DCCT results suggested a borderline salutary effect of intensive treatment on combined macrovascular events (45). A follow-up report by the Stockholm Diabetes Intervention Study suggested that intensive treatment leads to reduced IMT thickness and less stiffening of the carotid arteries (30), although the results were not uniformly supportive. The current study does not support an association between HbA_{1c} at the time of carotid ultrasonography or the mean HbA_{1c} during the 6.5 years of the DCCT and IMT. In addition, treatment assignment during the DCCT did not have an apparent effect on IMT. It may be argued that the absence of a major difference in HbA_{1c} at the time of ultrasonography between the two groups might explain the absence of an effect, and that 1.5 years of similar control after the DCCT may obliterate any benefit that 6.5 years of better control may have had. However, as much of the adverse effect of hyperglycemia in atherosclerosis is thought to be a chronic effect, e.g., cumulative advanced glycosylation end product formation (46), some differential effect of DCCT exposure might reasonably be expected. However, the current findings do not support an obvious association between glycemia and CVD in type 1 diabetes, consistent with several previous studies (3,6,28). Further follow-up will permit more definitive assessments as age, IMT, atherosclerosis, and clinical events all increase. Future studies will also help determine whether recent American Diabetes Association guidelines on the management of diabetic dyslipidemia in type 2 diabetes (47) should apply to type 1 diabetic subjects as well. The predictive power of LDL cholesterol for IMT in the current study provides some further support for aggressive cholesterol lowering (47). Intensive glycemic control in type 1 diabetes, nonetheless, remains the central component of diabetes management in view of its undoubted benefit in delaying or preventing microvascular complications and improving the CVD risk profile.

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APPENDIX: THE EDIC RESEARCH GROUP

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REFERENCES

- Dorman JS, LaPorte RE, Kuller LH, Cruickshanks KJ, Orchard TJ, Wagener DK, Becker DJ, Cavender DE, Drash AL: The Pittsburgh insulin-dependent diabetes mellitus (type 1 diabetes) morbidity and mortality study: mortality results. *Dia betes* 33:271–277, 1984
- Krolewski AS, Kosinski EJ, Warram JH, Leland OS, Busick EJ, Asmal AC, Rand LI, Christlieb AR, Bradley RF, Kahn CR: Magnitude and determinants of coronary artery disease in juvenile-onset insulin-dependent diabetes mellitus. *Am* J Cardiol 59:750–755, 1987
- Maser RE, Wolfson SK, Stein EA, Drash AL, Becker DJ, Dorman JS, Ellis D, Orchard TJ: Cardiovascular disease and arterial calcification in insulin-dependent diabetes mellitus: interrelationships and risk factor profiles: Pittsburgh Epidemiology of Diabetes Complications Study-VI. Arterioscler Thromb 11:958–965, 1991
- 4. Orchard TJ: Summary and comment: magnitude and determinants of coronary

artery disease in juvenile-onset insulin-dependent diabetes mellitus. *Diabetes Spectrum* 4:344–345, 1991

- Borch-Johnsen K, Andersen PK, Deckert T: The effect of proteinuria on relative mortality in type 1 (insulin-dependent) diabetes mellitus. *Diabetologia* 28:590–596, 1985
- Lloyd CE, Kuller LH, Becker DJ, Ellis D, Wing RR, Orchard TJ: Coronary artery disease in type 1 diabetes: gender differences in risk factors, but not risk. Arterioscler Thromb Vasc Biol 16:720–726, 1996
- 7. Fried LP, Borhani NO, Enright P, Furberg CD, Gardin JM, Kronmal RA, Kuller LH, Manolio TA, Mittelmark MB, Newman A, O'Leary DH, Psaty B, Rautaharju P, Tracy RP, Weiler PG: The Cardiovascular Health Study: design and rational. *Ann Epidemiol* 1:263–276, 1991
- The ARIC Investigators: The Atherosclerosis Risk in Communities (ARIC) Study: design and objectives. Am J Epidemiol 129:687–702, 1989
- Salonen R, Salonen JT: Progression of carotid atherosclerosis and its determinants: a population-based ultrasonography study. *Atherosclerosis* 81:33–40, 1990
- Burke GL, Evans GW, Riley WA, Sharrett AR, Howard G, Barnes RW, Rosamond W, Crow RS, Rautaharju PM, Geiss G, for the ARIC Study Group: Arterial wall thickness is associated with prevalent cardiovascular disease in middle-aged adults: the Artherosclerosis Risk in Communities (ARIC) Study. *Stroke* 26:386–391, 1995
- Allan PL, Mowbray PI, Lee AJ, Fowkes GR: Relationship between carotid intima-media thickness and symptomatic and asymptomatic peripheral arterial disease the Edinburgh Study. *Stroke* 28:348–353, 1997
- Bots ML, Hoes AW, Koudstaal PJ, Hofman A, Grobbee DE: Common carotid artery intima-media thickness and risk of stroke and myocardial infarction: the Rotterdam Study. *Circulation* 96:1432–1437, 1997
- Chambless LE, Heiss G, Folsom AR, Rosamond W, Szklo M, Sharrett AR, Clegg X: Association of coronary heart disease incidence with carotid arterial wall thickness and major risk factors: the Artherosclerosis Risk in Communities (ARIC) Study, 1987–1993. Am J Epidemiol 146:483–494, 1997
- 14. ACAPS Group: Rationale and design for the Asymptomatic Carotid Artery Plaque Study (ACAPS). *Control Clin Trials* 13:293–314, 1992
- Crouse JR, Byington RP, Bond MG, Espeland MA, Sprinkle JW, McGovern M, Furberg CD: Pravastatin, lipids, and atherosclerosis in the carotid arteries: design features of a clinical trial with carotid atherosclerosis outcome. *Con trol Clin Trials* 13:495–506, 1992
- Hennerici M, Klephas W, Gries FA: Regression of carotid plaques during lowdensity lipoprotein cholesterol elimination. *Stroke* 22:989–992, 1991
- Furberg CD, Borhani NO, Byington RP, Gibbons ME, Sowers JR: Calcium antagonists and atherosclerosis: the multicenter Isradipine/Diuretic Atherosclerosis Study. Am J Hypertens 6:24S–29S, 1993
- Blankenhorn DH, Selzer RH, Crawford DW, Barth JD, Liu C-R, Liu C-H, Mack WJ, Alaupovic P: Beneficial effects of colestipol-niacin therapy on the common carotid artery: two- and four-year reduction of intima-media thickness measured by ultrasound. *Circulation* 88:20–28, 1993
- Hodis HN, Mack WJ, LaBree L, Selzer RH, Liu C, Liu C, Alaupovic P, Kwong-Fu H, Azen SP: Reduction in carotid arterial-wall thickness using lovastatin and dietary therapy: a randomized controlled clinical trial. *Ann Intern Med* 124:548–556, 1996
- 20. Furberg CD, Adams HP, Applegate WB, Byington RP, Espeland MA, Hartwell T, Hunninghake DB, Lefkowitz DS, Probstfield J, Riley WA, Young B for the Asymptomatic Carotid Artery Progression Study (ACAPS) Research Group: Effect of Iovastatin on early carotid atherosclerosis and cardiovascular events. *Circulation* 90:1679–1687, 1994
- Heiss G, Sharrett AR, Barnes R, Chambless LE, Szklo M, Alzola C, ARIC Investigators: Carotid atherosclerosis measured by B-mode ultrasound in populations: associations with cardiovascular risk factors in the ARIC Study. *Am J Epidemiol* 134:250–256, 1991
- 22. O'Leary DH, Polak JF, Kronmal RA, Kittner SJ, Bond G, Wolfson SK Jr, Bommer W, Price TR, Gardin JM, Savage PJ: Distribution and correlates of sonographically detected carotid artery disease in the Cardiovascular Health Study. *Stroke* 23:1752–1760, 1992
- Folsom AR, Eckfeldt JH, Weitzman S, Ma J, Chambless LE, Barnes RW, Cram KB, Hutchinson RG: Relation of carotid artery wall thickness to diabetes mellitus, fasting glucose and insulin, body size, and physical activity. *Stroke* 25:66–73, 1994
- Kawamori R, Yamasaki Y, Matsushima H, Nishizawa H, Nao K, Hougaku H, Maeda H, Handa N, Matsumoto M, Kamada T: Prevalence of carotid atherosclerosis in diabetic patients. *Diabetes Care* 15:1290–1294, 1992
- 25. Pujia A, Gnasso A, Irace C, Colonna A, Mattioli PL: Common carotid arterial

wall thickness in type 1 diabetes subjects. *Diabetes Care* 17:1330–1336, 1994 26. Bonora E, Tessari R, Micciolo R, Zenere M, Targher G, Padovani R, Falezza

- G, Muggeo M: Intimal-medial thickness of the carotid artery in nondiabetic and type 1 diabetes patients. *Diabetes Care* 20:627–631, 1997
- Yamasaki Y, Kawamori R, Matsushima H, Nishizawa H, Kodama M, Kajimoto Y, Morishima T, Kamada T: Atherosclerosis in carotid artery of young type 1 diabetes patients monitored by ultrasound high-resolution B-mode imaging. *Diabetes* 43:634–639, 1994
- Kanters SDJM, Algra A, Banga J: Carotid intima-media thickness in hyperlipidemic type 1 and type 2 diabetic patients. *Diabetes Care* 20:276–280, 1997
- Koivisto VA, Stevens LK, Mattock MB, Ebeling P, Stephenson J, Idzior-Walus B, EURODIAB Type 1 Diabetes Complications Study Group: Cardiovascular disease and its risk factors in type 1 diabetes in Europe. *Diabetes Care* 19:689–697, 1996
- Jensen-Urstad KJ, Reichard PG, Rosfors JJS, Lindblad LEL, Jensen-Urstad MT: Early atherosclerosis is retarded by improved long-term blood glucose control in patients with type 1 diabetes. *Diabetes* 45:1253–1258, 1996
- 31. The Diabetes Control and Complications Trial Research Group: The effect of intensive treatment of diabetes on the development and progression of longterm complications in insulin-dependent diabetes mellitus. N Engl J Med 329:977–986, 1993
- 31a.EDIC Research Group: Epidemiology of diabetes interventions and complications (EDIC): design, implementation, and preliminary results of a long-term follow-up of the Diabetes Control and Complications Trial cohort. *Diabetes Care* 22:99–111, 1999
- D'Agostino RB, Burke G, O'Leary D, Rewers M, Selby J, Savage PJ, Saad MF, Bergman RN, Howard G, Wagenknecht L, Haffner SM: Ethnic differences in carotid wall thickness: the Insulin Resistance Atherosclerosis Study. *Stroke* 27:1744–1749, 1996
- 33. Wei M, Gonzalez C, Haffner SM, O'Leary DH, Stern MP: Ultrasonographyassessed maximum carotid wall thickness in Mexico City residents and Mexican-Americans in San Antonio, TX: association with diabetes and cardiovascular risk factors. *Atheroscler Thromb Vasc Biol* 16:1184–1188, 1996
- 34. DCCT Research Group: The Diabetes Control and Complications Trial (DCCT): design and methodologic considerations for the feasibility phase. *Dia betes* 35:530–545, 1986
- DCCT Research Group: DCCT Manual of Operations. Springfield, VA, National Technical Information Service 93-183382, 1993
- Lachin JM: Some large-sample distribution-free estimation tests for multivariate partially incomplete data from the populations. *Stat Med* 11:1151–1170, 1992
- 37. Crouse JR: B-mode ultrasound in clinical trials. Circulation 88:319-320, 1993
- Espeland MA, Craven TE, Riley WA, Corson J, Romont A, Furberg CD: Reliability of longitudinal ultrasonographic measurements of carotid intimalmedial thickness. *Stroke* 27:480–485, 1996
- 39. Burke GL, Evans GW, Riley WA, Sharrett AR, Howard G, Barnes RW, Rosamond W, Crow RS, Rautaharju PM, Heiss G: Arterial wall thickness is associated with prevalent cardiovascular disease in middle-aged adults: the Atherosclerosis Risk in Communities (ARIC) Study. *Stroke* 26:386–391, 1995
- 40. Ganda OP, Arkin CF: Hyperfibrinogenemia: an important risk factor for vascular complications in diabetes. *Diabetes Care* 15:1245–1250, 1992
- Mustard JF, Packham MA: Platelets and diabetes mellitus. N Engl J Med 297:1345–1347, 1977
- 42. Chambless LE, Heiss G, Folsom AR, Rosamond W, Szklo M, Sharrett AR, Clegg LX: Association of coronary heart disease incidence with carotid arterial wall thickness and major risk factors: the Atherosclerosis Risk in Communities (ARIC) Study, 1987–1993. Am J Epidemiol 146:483–494, 1997
- Hodis HN, Mack WJ, LaBree L, Selzer RH, Liu C, Liu C, Azen SP: The role of carotid arterial intima-media thickness in predicting clinical coronary events. *Ann Intern Med* 128:262–269, 1998
- 44. Klein R, Santiago J, Cleary P, Moss S, Nathan D: A comparison of the study populations in the Diabetes Control and Complications Trial (DCCT) and the Wisconsin Epidemiologic Study of Diabetic Retinopathy (WESDR). Arch Intern Med 155:745–754, 1995
- 45. Diabetes Control and Complications Trial Research Group: Effect of intensive diabetes management on macrovascular events and risk factors in the Diabetes Control and Complications Trial. Am J Cardiol 75:894–903, 1995
- Lyons TJ, Lopes-Virella MF: *Glycosylation-Related Mechanisms in Diabetes* and Atherosclerosis: Molecular Basis and Clinical Aspects. Drazmiz B, Eckel RH, Eds. New York, Elsevier, 1993, p. 191–212
- American Diabetes Association: Management of dyslipidemia in adults with diabetes (Position Statement). *Diabetes Care* 21 (Suppl. 1):S36–S44, 1998

Author Queries (please see Q in margin and underlined text)

Q1: Is "w" the correct symbol for equations such as "w = 153"? Does it need explanation at first mention? Q2: The last line of the first footnote has been moved to Acknowledgments. Q2a: Please supply a running title of 40 characters. Q3: In Table 1, please supply title of article by Anderson et al. Q4: Please list the name of the machine from Acuson. Q5: In Table 2, last line of footnote, P = 11 correct? Q6: Should CHD be changed to CAD or CVD? Q7: Stockholm Diabetes Intervention "Society" in discussion vs. "Study" in introduction. OK as is or change needed? Q8: Appendix—The term "(past)" appears after "A. Kowarski"; does this mean "past member"? Q9: For ref. 33, please list volume, page range, and year of publication. Q10: For ref. 35, please list location of publisher.

Table 1: First column, row six: should the number after \pm begin with a decimal point (0.07)?

ATTACHMENT 2

SAS Code for Baseline, Nondiabetic Control Subject, Univariate Correlation, Multivariate Analyses, and Treatment Group Difference Tabulations from EDIC One Year IMT Datasets in the NIDDK Repository

/* /* Program: R:\05_Users\Norma\EDIC\Carotid Artery\getdata.sas /* Author: Norma Pugh /* Date: 01 February 2008 /* Purpose: Create SAS datasets from the SAS transport files provided to the repository. /* NEW LOCATION FOR EDIC SAS FILES */ libname newlib 'R:\05_Users\Norma\EDIC\Carotid Artery'; /* ORIGINAL LOCATION OF EDIC TRANSPORT FILES */ filename file1 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMTdata\edcimt1x.xpt'; filename file2 'R:\03_Data_And_Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMTdata\edicimt1.xpt'; /* CREATE THE DATASETS */ proc cimport library=newlib infile=file1; run;

proc cimport library=newlib infile=file2; run;

```
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\verify_trt.sas
/* Author: Norma Pugh
       01 February 2008
/* Date:
/* Purpose: Verify that treatment assignment matches DCCT Baseline dataset.
/************************
/* Libnames and formats */
/****************************/
libname trt 'R:\03 Data And Tools\Studies\DCCT-EDIC\DCCT-
EDIC FINAL\DCCT\Analyses\Baseline\SAS DATA';
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
/* Get treatment assignment from DCCT baseline */
data trt;
set trt.baseline;
keep mask_pat group;
rename group=baseline_group;
run;
proc sort data=trt; by mask_pat; run;
/* Get 1-year-IMT-data dataset treatment assignments */
data temp1;
set data.edicimt1;
keep mask pat group;
rename group=temp1_group;
run:
proc sort data=temp1; by mask pat; run;
proc freq data=temp1; tables temp1_group / missing; title'edicimt1'; run;
/* Merge project data with treatment assignment */
data table1; merge temp1(in=x1) trt(in=x2); by mask pat; if x1 &
baseline group^=temp1 group; run;
```

```
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\table1.sas
/* Author: Norma Pugh
/* Date: 01 February 2008
/* Purpose: Replicate Table 1 results from Diabetes article: Effect of Intensive Diabetes
       Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes
/*
/*
         Interventions and Complications (EDIC).
/*
          (1999)
/****************************
/* Libnames and formats */
/*************************/
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
/********************************/
/* Get Table 1 variables */
/****************************/
data table1; set data.edicimt1; run;
/* Table 1 Replication Analysis */
title'Table 1: Baseline clinical characteristics of EDIC participants';
title2'by sex and original treatment group assignment during the DCCT';
title3'Categorical Counts/Percentages & p-values'; run;
proc freq data=table1;
tables sex*group;
run;
proc freq data=table1;
tables sex*group*bmi 27 / chisq;
tables sex*group*ht / chisq;
tables sex*group*aer_40 / chisq;
tables sex*group*smoking / chisq;
run;
title2'Quantitative Means & Standard Deviations'; run;
proc sort data=table1; by sex group; run;
proc means data=table1 n mean std;
by sex group;
var att age att durn height bmi nwst hip sbp dbp tchol hdl ldl trig aer gfr std ins
hba1c avg a1c fscore;
run;
proc sort data=table1; by group; run;
proc npar1way data=table1(where=(sex='F')) wilcoxon;
class group;
var att_age att_durn height bmi nwst_hip sbp dbp tchol hdl ldl trig aer gfr std_ins
hba1c avg_a1c fscore;
```

```
title3'Women';
run;
proc npar1way data=table1(where=(sex='M')) wilcoxon;
class group;
var att_age att_durn height bmi nwst_hip sbp dbp tchol hdl ldl trig aer gfr std_ins
hba1c avg_a1c fscore;
title3'Men';
run;
```

```
options mprint;
             /********
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\table2.sas
/* Author: Norma Pugh
/* Date: 01 February 2008
/* Purpose: Replicate Table 2 results from Diabetes article: Effect of Intensive Diabetes
/*
        Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes
/*
        Interventions and Complications (EDIC).
         (1999)
/*
/*****************************
/* Libnames and formats */
/************************/
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
/* Get Table 2 population and variables */
/* Nondiabetics */
data nondiab;
set data.edcimt1x;
/* Flag dataset as nondiabetic (diab=0) */
diab=0;
```

```
label diab = 'Diabetic? (0=n,1=y)';
```

```
/* Recode 'gender' to match 'sex' variable from diabetic dataset */
if gender=1 then sex='M'; else if gender=2 then sex='F';
```

```
/* Type 1 diabetics */
data diab;
set data.edicimt1(where=(decade in('20 - 29','30 - 39','40 - 49')));
```

/* Flag dataset as diabetic (diab=1) */
diab=1;
label diab = 'Diabetic? (0=n,1=y)';

```
keep mask_pat diab sex decade common internal;
run;
```

/* Merge data */
data table2; set nondiab diab;; run;

```
/*********************************/
/* Table 2 Replication Analysis */
/******************************/
title'Table 2: Comparison of IMT in nondiabetic subjects with EDIC type 1 diabetes
cohort';
```

```
title2'Quantitative Means & Standard Deviations'; run;
proc sort data=table2; by sex diab decade; run;
proc means data=table2 n mean std;
by sex diab decade;
var common;
run;
proc means data=table2 n mean std;
by sex diab decade;
var internal;
run;
%macro pval(sex,age,var,title);
proc sort data=table2; by diab; run;
proc npar1way data=table2(where=(sex="&sex" & decade="&age")) wilcoxon;
 class diab;
 var &var;
 title3"&title";
run;
%mend pval;
%pval(F,20 - 29,common,%str(Women, 20-29));
%pval(F,30 - 39,common,%str(Women, 30-39));
%pval(F,40 - 49,common,%str(Women, 40-49));
%pval(F,20 - 29,internal,%str(Women, 20-29));
%pval(F,30 - 39,internal,%str(Women, 30-39));
%pval(F,40 - 49,internal,%str(Women, 40-49));
%pval(M,20 - 29,common,%str(Men, 20-29));
%pval(M,30 - 39,common,%str(Men, 30-39));
%pval(M,40 - 49,common,%str(Men, 40-49));
%pval(M,20 - 29,internal,%str(Men, 20-29));
%pval(M,30 - 39,internal,%str(Men, 30-39));
%pval(M,40 - 49,internal,%str(Men, 40-49));
```

```
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\table3.sas
/* Author: Norma Pugh
/* Date: 01 February 2008
/* Purpose: Replicate Table 3 results from Diabetes article: Effect of Intensive Diabetes
      Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes
/*
/*
        Interventions and Complications (EDIC).
/*
        (1999)
/****************************
/* Libnames and formats */
/**************************/
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
```

/*************************/
/* Get Table 3 variables */
/***********************/
data table3; set data.edicimt1; run;

/**********************/
/* Table 3 Replication Analysis */
/**********************/
title'Table 3: Spearman correlation coefficients of clinical';
title2'characteristics with average maximum IMT';
title3'Common & Internal Carotid Artery Results by Gender'; run;
proc sort data=table3; by sex; run;
proc corr data=table3 spearman;
by sex;
var att_age att_durn height bmi nwst_hip sbp dbp tchol hdl ldl trig cum_smok std_ins
hba1c avg_a1c fscore aer gfr;
with common internal;
run;

```
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\table4.sas
/* Author: Norma Pugh
/* Date: 01 February 2008
/* Purpose: Replicate Table 4 results from Diabetes article: Effect of Intensive Diabetes
      Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes
/*
/*
        Interventions and Complications (EDIC).
/*
        (1999)
/****************************
/* Libnames and formats */
/*************************/
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
/* Get Table 4 population and variables */
data table4; set data.edicimt1;
if group='EXPERIMENTAL' then intense=1; else intense=0;
run;
/* Table 4 Replication Analysis */
title'Table 4: Summary of multiple linear regression models for common carotid IMT and
internal carotid IMT';
proc reg data=table4(where=(sex='F'));
title'Women';
model common = att age att durn height cum smok bmi sbp ldl gfr avg a1c intense;
run;
proc reg data=table4(where=(sex='M'));
title'Men';
model common = att_age att_durn height cum_smok bmi sbp ldl gfr avg_a1c intense;
```

run;

```
/*
/* Program: R:\05_Users\Norma\EDIC\Carotid Artery\table5.sas
/* Author: Norma Pugh
/* Date: 01 February 2008
/* Purpose: Replicate Table 5 results from Diabetes article: Effect of Intensive Diabetes
      Treatment on Carotid Artery Wall Thickness in the Epidemiology of Diabetes
/*
/*
        Interventions and Complications (EDIC).
/*
        (1999)
/****************************
/* Libnames and formats */
/**************************/
libname data 'R:\05 Users\Norma\EDIC\Carotid Artery';
%include 'R:\03 Data And Tools\Studies\DCCT-EDIC\NEW-Studies\1-year-IMT-data\fmt.sas';
/* Get Table 5 population and variables */
data table5;
set data.edicimt1(where=(decade in('20 - 29','30 - 39','40 - 49')));
run;
/* Table 5 Replication Analysis */
title'Table 5: Average maximum IMTs by age and randomized treatment assignment';
proc sort data=table5; by sex decade group; run;
proc means data=table5 n mean std;
by sex decade group;
var common;
run;
proc means data=table5 n mean std;
by sex decade group;
var internal;
run;
```