Cardiovascular Diabetology



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Is type 2 diabetes mellitus a vascular disease (atheroscleropathy) with hyperglycemia a late manifestation? The role of NOS, NO, and redox stress.

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Abstract

Background: Cardiovascular disease accounts for at least 85 percent of deaths for those patients with type 2 diabetes mellitus (T2DM). Additionally, 75 percent of these deaths are due to ischemic heart disease.

Hypothesis: Is type 2 diabetes mellitus a vascular disease (atheroscleropathy) with hyperglycemia a late manifestation? The role of NOS, NO, and redox stress.

Testing of the hypothesis: The vulnerable three arms of the eNOS reaction responsible for the generation of eNO is discussed in relation to the hypothesis: (1). The L-arginine substrate. (2). The eNOS enzyme. (3). The BH4 cofactor.

Implications of the hypothesis: If we view T2DM as a vascular disease initially with a later manifestation of hyperglycemia, we may be able to better understand and modify the multiple toxicities associated with insulin resistance, metabolic syndrome, prediabetes, overt T2DM, and accelerated atherosclerosis (atheroscleropathy). The importance of endothelial nitric oxide synthase, endothelial nitric oxide, tetrahydrobiopterin (BH4), L-arginine, and redox stress are discussed in relation to endothelial cell dysfunction and the development and progression of atheroscleropathy and T2DM. In addition to the standard therapies to restore endothelial cell dysfunction and stabilization of vulnerable atherosclerotic plaques, this article will discuss the importance of folic acid (5MTHF) supplementation in this complex devastating disease process.

Atheroscleropathy and hyperglycemia could be early and late manifestations, respectively, in the natural progressive history of T2DM.

Background

The historical discovery of the enzyme family nitric oxide synthases (NOS) in 1989 is relatively recent and we now know there are three distinct isoforms of NOS: neural NOS-1 (nNOS); inducible NOS-2 (iNOS); and endothelial NOS-3 (eNOS). These three isoforms were discovered in the above order from 1991-1994 [1]. eNOS, the rate limiting enzyme responsible for the production of endothelial derived nitric oxide (eNO) was sequenced and cloned in 1992 [2,3]. Each NOS enzyme has its own

Table I: Comparisons of the human NOS enzymes

nNOS NOS-I	Chromosome 12	Amino acids 1434 aa 161 kDa	high Ca++ requirement
iNOS NOS-2	Chromosome 17	Amino acids 1153 aa 131 kDa	low Ca++ requirement
eNOS NOS-3	Chromosome 7	Amino acids 1203 aa 133 kDa	high Ca++ requirement

Table 2: The positve protective effects of eNOS

- I. Promotes vasodilatation of vascular smooth muscle.
- 2. Counteracts smooth muscle cell proliferation.
- 3. Decreases platelet adhesiveness.
- 4. Decreases adhesiveness of the endothelial layer to WBCs "Teflon effect".
- 5. Anti-inflammatory.
- 6. Anti-oxidant. It scavenges reactive oxygen species, locally. Acts as a chain breaking antioxidant to scavenge ROS.
- Anti-fibrotic. When NO is normal or elevated MMPs are low and conversely if NO is low MMPs are elevated and active.
- NO has diverse anti-atherosclerotic actions on the arterial vessel wall: including antioxidant effects by direct scavenging of ROS

 RNS acting as chain breaking antioxidants.

Table 3: Janus - faced properties of NO

GOOD	BAD
eNO, nNO	iNO *
Chromosomes 7,12	Chromosome 17
Duration: Seconds to minutes	Duration: Hours to days
Signaling molecule.	Killer molecule.
Regulated: small bursts	Unregulated: larger bursts and longer duration
Cytoprotective	Cytotoxic

^{*}In host defense mechanisms iNO does just what we want it to, i.e. it kills invading organisms. Inducible NO can also be thought of as inflammatory NO. Paradoxically, in both acute and chronic inflammatory states iNO also kills the native hosts cells in the immediate surrounding area.

unique amino acid structure sharing some 50 percent homology indicating a common ancestral background. Additionally, each NOS isoform is transcribed from a separate gene (table 1).

Nitric oxide, voted the molecule of the year by Science Magazine in 1992 [4] has been shown to be an extremely important signaling molecule in the cardiovascular system and in 1998 Furchgott RF, Ignarro LJ, and Murad F. were awarded the Nobel Prize in Medicine and Physiology "for their discoveries concerning nitric oxide as a signaling molecule in the cardiovascular system" (tables 2,3).

Table 4: Lab - garbage acronym

Factors related to eNOS dysfunction with decreased NO. factors known to uncouple the eNOS enzyme.

L – Lipids LDL – Cholesterol elevated both native and oxidized. Elevated Triglycerides of metabolic syndrome. Decreases or impairs dimethylarginine dimethylaminohydrolase (DDAH). Decline in DDAH activity. Increase in ADMA activity.

A – Arginine (L-arginine) decreased or impaired: By competitive interaction with ADMA (asymmetric dimethylarginine) an endogenous nitric oxide synthase inhibitor. Elevated in proatherogenic conditions. ACE, Ang II, endothelin, hypertension, insulin resistance, prediabetes, T2DM, and hyperhomocysteinemia.

B-BH4 Decreased or impaired Folic acid regeneration of the cofactor BH4 i.e. BH2 \rightarrow BH4 with 5 methyl tetrahydrofolate (5-MTHF) active form of folic acid.

G-Glucose elevation. T2DM. DDAH effect increasing ADMA similar to lipids and elevated homocysteine.

A – Arginine (L-arginine) decreased or impaired: By nitrosylation (nitroarginine). By ONOO' peroxynitrite similar to nitrosylation of tyrosine (nitrotyrosine).

R-ROS reactive oxygen species. Specifically O2' and ONOO' Decline in DDAH activity results in increased ADMA.

B-BH4 decreased or impaired. BH4 oxidized to BH2. Loss of a naturally occurring antioxidant with diminished ROS scavenging capability. A - ADMA Homocysteine impairs DDAH enzyme and increases ADMA.

 $G-Glucose\ elevated-glycated\ eNOS.$

 $E-Endothelial\ NOS\ enzyme\ decreased\ and\ or\ dysfunctional.\ Gene\ polymorphism\ Glu298 \to Asp,\ additional\ unidentified\ gene\ polymorphisms,\ CRP\ causing\ marked\ down\ regulation\ of\ eNOS\ mRNA\ and\ protein\ expression,\ eNOS\ enzyme\ glycation.\ Last\ but\ most\ importantly$

eNOS enzyme UNCOUPLING

This relatively brief and recent history of NOS and NO is in contrast to the ancient history of atherosclerosis which dates to the time of the Egyptians. The theories of atherosclerosis are legion and the atherosclerologists of today have a proud history upon which to build for the future [5].

The endothelial cell, such a central player, responsible for remodeling of the arterial vessel wall in atherosclerosis (intimopathy) and responsible for the synthesis of eNOS and subsequent eNO production will be discussed and related to the accelerated atherosclerosis (atheroscleropathy) associated with type 2 diabetes mellitus (T2DM) [5,8–11].

This review will discuss dysfunction of the eNOS enzyme and its potential to uncouple, resulting in the endothelial cell becoming a net producer of damaging superoxide (O2') instead of the protective eNO (tables 4,5) [6]. The lack of eNO from the endothelial cell is related to the development of atheroscleropathy and is related to the underproduction and / or the excessive consumption of eNO

Table 5: eNOS reaction the three critical arms

IF THE eNOS REACTION IS UNCOUPLED: THEN THE REACTION IS:

$$NAD(P)H \xrightarrow{NAD(P)H \text{ Oxidase}} NAD(P) + + O2'$$

(L-arginine is not converted to NO and L-citrulline).

TREATMENT PARADIGM:

- (I) Prevent competitive inhibition by ADMA (Reduce substrates: LDL-C, Triglycerides, Homocysteine, and Glucotoxicity formation. Prevent L-arginine from being converted to nitroarginine by nitrosylation. Reducing Redox Stress A-FLIGHT toxicities reduction of ROS. ARGININE SUPPLEMENTATION.
- (2) Prevent hs-CRP from decreasing eNOS by lowering hs-CRP and preventing glycation of Enos enzyme. Aggressive treatment of elevated substrates.
- (3) Add FOLIC ACID not only to lower Hcy and ADMA levels but also to restore BH2, BH3 to the active catalytic cofactor BH4. Use the RAAS acronym and prevent the A-FLIGHT Toxicities (ROS) from developing i.e. decrease the manifold toxicities of MS, IR, PD and overt T2DM.

BE AGGRESSIVE AND TREAT TO KNOWN GOALS:

by redox stress. The uncoupling of this complex enzyme will aid in the understanding of the development of atheroscleropathy and the role of oxidative and reductive stress (redox stress) in patients with T2DM [7,8].

Definitions

It is important to set forth the authors' working definitions before proceeding with this review.

ATHEROSCLEROSIS is a systemic dysfunctional endothelial, focal occurring, chronic inflammatory, fibroprolifertive, angiogenic, prothrombotic, multifactorial disease of the arterial intima caused by the retention of modified low density lipoproteins, hemodynamic stress, and accelerated by redox stress [5,8–11].

ATHEROSCLEROPATHY: The term used to describe the unique accelerated atherosclerosis observed in and associated with metabolic syndrome, insulin resistance, prediabetes, and overt T2DM which is driven by reactive oxygen species associated with multiple toxicities (table 6) [8].

T2DM is a heterogeneous, multifactorial, polygenic disease characterized by a defect in insulin's secretion (the beta cell secretory defect) and action (insulin resistance) which results in elevated glucose [8,12].

Table 6: The manifold toxicities of insulin resistance, metabolic syndrome and T2DM.

	A-FLIGHT toxicities	
Α	Amylin (hyperamylinemia)/amyloid toxicity	ROS
	Ang II (also induces PKC)	ROS
	AGEs/AFEs (advanced Glycosylation / fructosylation products)	ROS
	Antioxidant reserve compromised	ROS
	Absence of antioxidant network	ROS
	Ageing	ROS
	Angiogenesis (induced redox stress) Arteriogenesis (impaired PAI-I)	ROS
	Atherosclerosis – Atheroscleropathy. [ROS beget ROS]	ROS
F	Free fatty acid toxicity	ROS
L	Lipotoxicity	ROS
I	Insulin toxicity (hyperinsulinemia-hyperproinsulinemia) (endogenous)	ROS
	Inflammation toxicity	ROS
G	Glucotoxicity (compounds peripheral insulin resist-	ROS
	ance) reductive stress	plus
	Sorbitol / 'olyol pathway	PKC
	Pseudohypoxia (NADH/NAD increased)	
Н	Hypertension toxicity	ROS
	t homocysteine toxicity	ROS
Т	Triglyceride toxicity	ROS

See reference [8,9]

T2DM may be viewed as a dynamic continuum of five stages (table 9) [12].

eNOS and eNO

Chromosome number 7 contains the vascular eNOS gene that encodes the transcription and synthesis of the 1203 amino acid protein enzyme eNOS. eNOS is located uniquely within caveolae of the plasma membrane which allows the monolayer of endothelial cells to compensate if its fellow cellmates become dysfunctional in order to maintain eNO production.

Once conceptualized, that a specific and unique gene (eNOS, iNOS, nNOS) encodes the transcription of NO in a specific organ system, the opposing faces of this highly specialized, small, bioactive, and readily diffusible gas NO can be better understood (table 2, 3) [13].

eNO is a natural occurring, endogenous, chain breaking antioxidant synthesized by the endothelial cell. eNO and superoxide are related to a healthy and dysfunctional endothelium, respectively [8].

THE HEALTHY ENDOTHELIUM IS A NET PRODUCER OF ENDOTHELIAL NITRIC OXIDE [eNO].

THE ACTIVATED, DYSFUNCTIONAL ENDOTHELIUM IS A NET PRODUCER OF SUPEROXIDE [O2'] ASSOCIATED WITH T2DM AND ATHEROSCLEROPATHY

There are at least three arms in the eNOS reaction capable of producing endothelial cell dysfunction with resultant redox stress to the intima and islet resulting in atheroscleropathy and a progressive decline in beta cell function within the vulnerable islet.

Presentation of the hypothesis

Is type 2 diabetes mellitus a vascular disease (atheroscleropathy) with hyperglycemia a late manifestation? The role of NOS, NO, and redox stress.

Testing of the hypothesis

The following eNOS reaction demonstrates the location of the vulnerable three arms of the eNOS reaction responsible for the generation of eNO.

IF THE eNOS REACTION IS UNCOUPLED: THEN THE REACTION IS:

$$NAD(P)H \xrightarrow{NAD(P)H \text{ Oxidase}} NAD(P) + + \underline{O2'}$$

L-arginine is uncoupled and not converted to NO and L-citrulline (figure 1) (tables 3,4).

THE L-ARGININE ARM

The substrate L-arginine is an essential amino acid and is usually not thought to be rate limiting (with the exception of high demands in a dysfunctional endothelial state such as atherosclerosis or the atheroscleropathy associated with T2DM).

Arginine deficiency is a rare occurrence, however, there can be competitive inhibition by the endogenously produced asymmetrical dimethylarginine (ADMA) and nitroarginine. ADMA is emerging as an important cause of endothelial cell dysfunction.

It is an endogenously produced competive inhibitor of Larginine which results in a relative deficiency of the natural substrate for the eNOS enzyme to produce eNO. ADMA accumulates by either a process of increased synthesis by the protein arginine N-methyltransferases

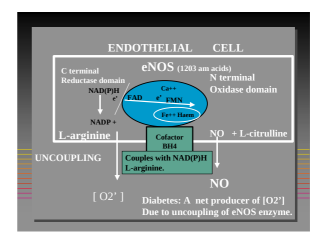


Figure I

Oxygen reacts with the eNOS enzyme in which the BH4 cofactor has coupled NAD(P)H with L-arginine to be converted to NO and L-citrulline. When uncoupling occurs the NAD(P)H reacts with O2 and the endothelial cell becomes a net producer of superoxide (O2').

(PRMTs) or decreased elimination by the enzyme dimethylarginine dimethylaminohydrolose (DDAH) [14–17].

DDAH selectively hydrolyzes ADMA to L-citrulline and dimethylamine, so that the higher the DDAH activity, the lower the ADMA levels. Importantly, inhibition of DDAH activity results in increased ADMA levels. Both native and oxidized LDL-cholesterol have been shown to up-regulate PRMT while causing a decrease in the activity of DDAH [18]. This results in an elevation of ADMA levels contributing to endothelial cell dysfunction and increased oxidative stress to the intima with subsequent accelerated atherosclerosis.

There are multiple causes for the elevation in ADMA levels (figure 2). Each of these toxicities relate to the A-FLIGHT acronym responsible for the production of ROS (table 6). Recently, Faldetta CM, Laurenti O, and Desideri G [19] were able to demonstrate in both normal human subjects and T2DM human patients that acute L-arginine infusion decreased plasma total homocysteine concentrations, counteract oxidative stress, and increased the availability of nitric oxide.

In summary, the relative deficiency of L-arginine due to elevation of ADMA levels contribute to oxidative stress and result in the atheroscleropathy associated with insulin resistance, metabolic syndrome, prediabetes, and overt T2DM.

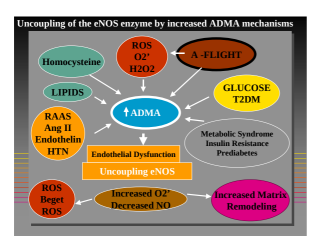


Figure 2

The multiple stressors causing uncoupling of the eNOS enzyme by decreased DDAH and the increased endogenous inhibitor of eNOS: ADMA.

Table 7: eNOS AGONISTS

- I. Bradykinin (BK).
- 2. Phosphorylation by PK-B also called Akt.
- 3. Mechanical forces Sheer stress.
- 4. Acetylcholine (muscarinic receptors).
- 5. ADP ATP (purinergic receptors).
- 6. Vascular Endothelial Cell Growth Factor (VEGF).
- 7. Histamine.
- 8. HSP 90.
- 9. Acetylation by myristate (N-terminal) by palimitate (cys I $5-\mbox{cys}26)$ Stimulated by BK.

THE eNOS ARM

Important agonists for the activation of the eNOS enzyme (table 7) play an important role in the production of eNO.

If there is a complete loss of eNOS enzyme as in the eNOS -/- knockout mouse model of Duplain and Sherrer [20] insulin resistance, hypertension, and hyperlipidemia develop. Their findings indicate that eNOS plays a major role in the regulation of insulin sensitivity. To have an effect on atheroscleropathy and T2DM it would not require a complete knockout of the gene but just one or two changes in the 1203 amino acid protein structure in the eNOS enzyme.

Such a gene polymorphism does exist and may become a major player in the polygenic causes of T2DM and atheroscleropathy:

 $Glu298 \rightarrow Asp Gene Polymorphism (Glu298 \rightarrow Asp GP)$

eNOS may be impaired by abnormal structure and or function that occurs in gene polymorphisms.

One of the first significant eNOS gene polymorphisms described to date is Glu 298 \rightarrow Asp. This amino acid substitution is responsible for a statistically significant increased frequency of hypertension and spastic angina [21]. Also, described in a different set of patients is the occurrence of increased abnormal coronary angiographic findings of atherosclerosis and an association of increased events [22–24].

The Glu298 \rightarrow Asp GP is responsible for a decrease in basal NO production in healthy subjects according to the publication by Veldman BA et al [25].

This gene polymorphism then could interact with other gene polymorphisms and especially environmental conditions such as smoking, obesity, and the A-FLIGHT toxicities of IR, MS, PD, and overt T2DM. Or as in the hypothesis, could be responsible for the development of IR and MS which then would progress to PD or overt T2DM. We now have a Japanese population, a UK population, and an Italian population that correlates Glu298 → Asp GP to atherosclerosis and CAD events [23–25].

These findings indicate that the eNOS gene is important to the development of accelerated atherosclerosis → atheroscleropathy. Furthermore, these three distinct patient populations indicate that this gene polymorphism is widely distributed throughout diverse patient populations and according to Veldman [25] is present in approximately 10 % of patient populations.

Recently, Noiri E and colleagues have shown that there is a significant association of the Glu298 \rightarrow Asp GP and end stage renal disease in T2DM which once again points to the importance of the endothelial cell and the eNOS reaction [26].

Since eNOS contains 1203 amino acids there is the possibility of other gene polymorphisms within the eNOS enzyme that have not been identified at this time. Additionally, this gene polymorphism could interact with other gene polymorphisms of other endogenous antioxidant enzymes in the T2DM patient as we know their antioxidant reserve is compromised (tables 6,8) [7,8,12].

Table 8: Antioxidants: catalytic/enzymatic inactivation of free radicals.

Enzymatic antioxidants

SUPER OXIDE DISMUTASE (SOD) - Location: mitochondrion

 $[O_2^- + SOD \rightarrow H_2O_2 + O_2]$

ecSOD (extracellular)

MnSOD (mitochondrial)

CuZnSOD (intracellular)

CATALASE – Location: peroxisome $[2H_2O_2 + catalase \rightarrow 2 H_2O + O_2]$

GLUTATHIONE PEROXIDASE – Location: mitochondrion/

($\mbox{\sc Glutamyl-cysteinyl-glycine}$ tripeptide) glutathione reduced -SH to the oxidized disulfide GSSG.

(Glutathione peroxidase) [GSH + $2H_2O_2 \rightarrow GSSG + H_2O + O_2$] (Glutathione reductase) [GSSG \rightarrow GSH] at the expense of [NADH \rightarrow NAD⁺] and/or [NAD(P)H \rightarrow NAD(P)⁺]

*NOS (nitric oxide synthase). – Location: membrane Isoforms:

(e) NOS (endothelial): good (importance of eNOS uncoupling) LDL native and oxidized.

(n) NOS (neuronal): good

(i) NOS (inducible-inflammatory): **GOOD** in host defense. **BAD** in chronic inflammation, ischemia – ischemia reperfusion injury, acute and chronic as in autoimmunity – TIDM.

 $O_2^{\text{-}}$ and nitric oxide (NO) are consumed in this process with the creation of reactive nitrogen species (RNS).

 O_2^- + NO \to ONOO- (peroxynitrite) + tyrosine \to nitrotyrosine. (also causes eNOS uncoupling)

Nitrotyrosine reflects redox stress and leaves a measurable footprint. NO: the good; O_2 : the bad; ONOO: the ugly [122]

eNO: A chain breaking antioxidant see (table 2) item 6. eNOS uncoupling causes the generation of O2' instead of NO induced by LDL-C, Glucose, O2', and ONOO'.

Nonenzymatic antioxidants

URIC ACID

VITAMIN A, VITAMIN C, VITAMIN E

THIOLS

<u>APOPROTEINS:</u> Ceruloplasmin and transferrin. Bind copper and iron in forms which cannot participate in the Fenton reaction. [7,8,12]

As the human genome unfolds there may be other related important gene polymorphisms identified.

The BH4 ARM

Tetrahydrobiopterin (BH4) is the necessary cofactor for coupling L-arginine to the NAD(P)H oxidase enzyme in order for L-arginine to be oxidized to NO and L-citrulline. The importance of this cofactor and its mechanism is just starting to emerge.

If this cofactor is not functioning properly the entire eNOS reaction uncouples and the endothelium becomes a net producer of superoxide [O2'] through the uncoupled NAD(P)H oxidase reaction. In addition to BH4 being the necessary cofactor it also serves as a naturally occurring

Table 9: The five stages of T2DM: the natural progressive history of T2DM.

I. LATENT STAGE:[EARLY]

Insulin Resistance:

*Genetic Component.

*Environmental Component. [Modifiable] Obesity / Sedentary life style. [Nonmodifiable] Ageing.

Beta Cell Defect: (Dysfunction)

*Genetic...... Abnormal processing, storage, or secretion.

*Intracellular extracellular amylin fibril toxicity. Abnormal processing, storage, or secretion.

Intra-Islet Endothelial Absorptive Defect:

*Heparan sulfate proteoglycan (HSPG) PERLECAN of the capillary endothelial cells avidly attracts amylin (IAPP) and islet amyloid forms an envelope around the capillary. This is in addition to the increase in basement membrane associated with the pseudohypoxia (associated with glucotoxicity) and the redox stress within the capillary.

II. TRANSITION STAGE: [MIDDLE]

PERSISTENT HYPERINSULINEMIA

PERSISTENT **HYPERAMYLINEMIA** – Continued remodeling of endocrine pancreas. (**AMYLOID**) Beta cell displacement, dysfunction, mass reduction, and diffusion barrier.

III. <u>IGT STAGE</u>: *"Pre-diabetes" HHS ADA term (Impaired Glucose Tolerance)

[LATE] *[Start treatment at this time] [Diagnose earlier: Rejuvenation of the 2 hour glucose tolerance blood sugar 140–199 mg.]

*Increased insulin resistance [Feeds forward] > Glucotoxicity [Feeds forward] > Insulin resistance [Feeds forward] > Glucotoxicity: creating a vicious cycle.

*Islet amyloid. Increasing beta cell defect. Loss of beta cell mass with displacement. [Remodeling of islet architecture including ECM] Beta cell loss centrally.

IV. IFG STAGE: *"Pre-diabetes" HHS ADA term (Impaired Fasting Glucose)

[LATER] Blood sugar > 110 mg./dl. < 126 mg./dl.

[Impaired Hepatic Glucose Production (HGP)]. Increasing global insulin resistance (**HEPATIC**) with subsequent gluconeogenesis. Feeding forward in the vicious cycle to accelerate insulin resistance globally

V. <u>OVERT STAGE</u>: [TOO LATE] 50% loss of Beta Cell Function.

<u>Va. Vb. Vc. Phases I, II, III</u> Mild / Moderate / Severe / Complete. Use medications that do not increase insulin or AMYLIN. Use combination therapy. **Start treatment at stage III-IV (IGT)** (**IFG**)

antioxidant capable of scavenging ROS such as superoxide O2' and perioxynitrite [ONOO']. If BH4 is deficient in supply or function the eNOS reaction not only is capable of uncoupling with subsequent excess [O2'] and [ONOO'] but also is at a disadvantage in that there would be less BH4 to scavenge these local occurring reactive oxygen species being produced in excess amounts [27].

It is important to note that a direct effect of folic acid is to keep this important cofactor BH4 intact so that it can maintain coupling of the eNOS reaction.

^{*}Paradigm Shift. Start treatment at the earlier stage of IGT

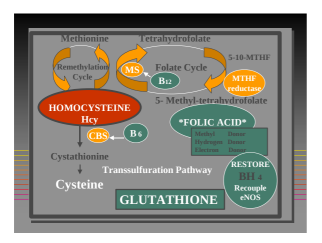


Figure 3

Hcy (demethylated Methionine) can enter the remethylation cycle with the aid of Folic Acid (5-Methyl-tetrahydrofolate) and the Folate cycle, the enzyme Methionine Synthase, and the necessary Cofactor B12. Folic Acid in addition to being a methyl-donor can serve as an electron or hydrogen donor to restore the required BH4 Cofactor to recouple the eNOS enzyme necessary for the production of eNO from Larginine. Hcy may also enter the Transsulfuration Pathway with the assistance of Cystathionine Beta Synthase (CBS) enzyme and the necessary Cofactor B6 to form Cystathionine and Cysteine which can then be excreted in the urine or converted to the very important antioxidant Glutathione. The substrate Folic Acid and the Cofactors B6 and B12 are very important not only for improving hyperhomocysteinemia but also important for the recoupling of the BH4 cofactor to the eNOS enzyme for the production of eNO.

The lowering of homocysteine is an indirect effect of folic acid aiding in the decreased production of ADMA due to the effect of Hcy on DDAH. Folic acid given to patients with T2DM have an improvement of endothelial relaxation before Hcy has time to be lowered. It works almost immediately before the Hcy is lowered [28,29].

When you look at the three arms of importance for the eNOS reaction there are two possible changes in the treatment paradigm (in addition to lowering the substrates of the A-FLIGHT toxicities (table 6) and the RAAS acronym [8]) for these patients who suffer from endothelial dysfunction associated with atheroscleropathy, IR, MS, PD, and T2DM.

First, we need to take a global risk reduction approach (by treating each risk factor in the A-FLIGHT toxicities acronym in order to decrease the clinical substrates of lipids, hypertension, and glucotoxicity without elevating endog-

enous insulin or amylin) in order to decrease the progression of the disease. Second, we should now include in the treatment paradigm the use of folic acid and arginine supplementation. T2DM is now considered to be a coronary risk equivalent and therefore all patients with T2DM should be on folate supplementation, as well as, the non-diabetic atherosclerotic patient.

Those patients with persistent endothelial cell dysfunction after the standard goals have been reached should consider arginine supplementation and possibly should be considered even sooner in the treatment paradigm if there is a history or evidence of symptomatic peripheral arterial disease, congestive heart failure and / or diastolic dysfunction [17,30,31].

Folic acid - 5MTHF

This water soluble B vitamin has been recently gaining considerable attention due to its positive role in cardio-vascular disease, neurodegenerative disorders, neural tube defects, and cancer. Folic acid plays an important role in the remethylation of homocysteine and is thus capable of lowering elevated levels of homocysteine (figure 3) [32].

Deficiency or impairment of folate metabolism is associated with hyperhomocysteinemia (hHcy), hypomethylation (the decreased one carbon unit transfer to purines and pyrimidines for DNA repair and biosynthesis), DNA damage, and impaired cell proliferation, malignancies, and impaired eNO production [33–37]. 5-MTHF is an electron donor, a hydrogen donor, and a methyl donor.

Each of these attributes confers a unique role to its function. In regards to the eNOS reaction it supplies both hydrogen and electrons to BH2 and BH3 to restore BH4 (the necessary cofactor for the eNOS reaction). This attribute allows the eNOS reaction to re-couple and restores the capability of this reaction to be once again a net producer of eNO and restores endothelial dependent vasodilatation in T2DM and hyperlipidemia [38]. Recently, the combination of folic acid, vitamin B12, and vitamin B6 was able to demonstrate a reduction of in-stent restenosis and recurrent events in up to one year follow up in the Swiss heart study [39–41].

The eNOS hypothesis

Is T2DM a vascular disease with a dysfunctional eNOS enzyme reaction, resulting in endothelial cell dysfunction and associated diminished eNO production? Is atheroscleropathy an early manifestation and hyperglycemia a later manifestation?

The following quote is pertinent to this discussion.

"Diabetes must take its place alongside the other major risk factors as important causes of cardiovascular disease. In fact, from the point of view of cardiovascular medicine, it may be appropriate to say: "diabetes is a cardiovascular disease" [42].

Implications of the hypothesis

Could it be that one of the polygenic causes of T2DM involve the eNOS enzyme? Such as one or more gene polymorphisms resulting in a defective but functioning eNOS enzyme. This would certainly help to explain the earlier discussed atheroscleropathy associated with T2DM.

This gene polymorphism could interact with the environment to produce an even more defective eNOS enzyme with increasing endothelial cell dysfunction. Smoking, obesity, hyperlipidemia, hypertension, sedentary life style are all known risk factors and all too often they cluster creating the metabolic syndrome with a markedly elevated tension of redox stress not only in the islet but also within the intima of the arterial vessel wall. A single amino acid substitution in the primary structure could have a considerable effect on the structure and function of this omnipotent enzyme and its reaction producing eNO.

Here is an example where a defective gene could be made much more impotent if there were additional stresses placed on it through a stressor molecule such as ADMA. This endogenously produced competitive inhibitor of Larginine (as well as an endogenous inhibitor of the eNOS reaction) would cause the eNOS reaction to uncouple and become a net producer of superoxide even before the glucose becomes elevated, thus stressing the islet and the intima before hyperglycemia is detected. This is why we should consider stress testing (by utilizing the 75 gram oral 2 hour post prandial glucose tolerance test) the patient in order to detect impaired glucose tolerance (stage III table 9) in order to reduce the oxidative and postprandial reductive stress to these organs in order to spare them from the devastating natural progressive history of T2DM and atheroscleropathy. Exercise induced skeletal muscle glucose transport (GLUT 4) is eNOS dependent [43-45]. If the production of eNO were defective due to gene polymorphism and environmental interaction there would be increasing peripheral insulin resistance. This is just what Duplain and Scherrer were able to demonstrate in their eNOS -/- model: Insulin resistance, hypertension, and hyperlipidemia.

In humans with gene polymorphism the eNOS enzyme may be capable to withstand the many years of redox stress before the defect in eNOS becomes evident. However, if the stressors were of sufficient magnitude such as seen in the adolescent youth with sedentary life style plus high carbohydrate, high fat western diet (over nutrition)

the manifestation of type 2 diabetes may become evident before the youth has reached the more normal middle age in life. This is just what we are experiencing in the USA. Not only are the adolescent youth developing T2DM at an alarming rate, but also, the age of onset of T2DM in adults is becoming younger and younger. It is entirely possible that these abnormal gene polymorphisms could be very old and have never caused a problem until there was sufficient changes in the environment (the thrifty gene hypothesis: gene – environmental interaction).

Peripheral insulin resistance would certainly place an undo burden on the pancreas with elevated levels of endogenous insulin and amylin with a chronically activated RAAS at the local organ-tissue levels.

The additional heightened redox stress and the A-FLIGHT toxicities associated with insulin resistance and metabolic syndrome would only aggravate the underlying eNOS polymorphisms by placing an undo burden on the islet and the intima.

Conclusion

"YOU CAN'T SEE THE FOREST FOR THE TREES"

If we approach T2DM as initially being a vascular disease for many years with glucotoxicity being a later manifestation, then as clinicians and researchers we may be more likely to view this disease in a global perspective.

The Steno-2 study by Gaede P et al. [46,47] just published, revealed a significant reduction in cardiovascular and microvascular events of nearly 50%. Their intensified intervention against multiple risk factors in patients with type 2 diabetes was quite impressive.

Therefore, in T2DM we should consider looking at the multiple toxicities and how they interact with each other to form "the whole disease" and return metabolic and redox homeostasis to the organism.

"We must protect the forests (the whole disease) for our children, grandchildren and children yet to be born. We must protect the forests for those who can't speak for themselves such as the birds, animals, fish and trees (and our patients)." --- Chief Edward Moody, Qwatsinas, Nuxalk Nation.

Competing interests

None declared.

Authors' Contributions

MH and ST collaborated equally on this article. MH wrote and edited this article. ST collaborated on writing and editing this article.

Abbreviations

Accelerated atherosclerosis (atheroscleropathy), type 2 diabetes mellitus (T2DM), nitric oxide (NO), endothelial nitric oxide (eNO), nitric oxide synthase (NOS), neural NOS-1 (nNOS), inducible NOS-2 (iNOS), endothelial NOS-3 (eNOS), superoxide (O2'), tetrahydrobiopterin (BH4), asymmetrical dimethylarginine (ADMA), dimethylarginine dimethylaminohydrolose (DDAH), protein arginine N-methyltransferases (PRMTs), nicotinamide dinucleotide phosphate reduced (NAD(P)H), glutamic $298 \rightarrow \text{aspartyl gene polymorphism (Glu298} \rightarrow \text{Asp GP}),$ perioxynitrite [ONOO'], coronary artery disease (CAD), insulin resistance (IR), metabolic syndrome (MS), prediabetes (PD), homocysteine (Hcy), hyperhomocysteinemia (hHcy), renin-angiotensin-aldosterone-system (RAAS), multiple metabolic toxicities of IR, MR, PD, and T2DM producing reactive oxygen species (A-FLIGHT acronym), reactive oxygen species (ROS), 5-methyl tetrahydrofolate (5MTHF), folic acid (5MTHF), glucose transporter 4 (GLUT 4), gene knockout (-/-).

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