Paleolithic-Style Diet and Coronary Artery Disease: The Tissue is the Issue?

Eri Toda, Takahashi Toru, R.B. Singh, Shaan E. Alam, Fabien De Meester, Agnieszka Wilczynska and Douglas Wilson
1Department of Cardiology, Tokai University Hachioji Hospital, Tokyo, Japan
2Department of Nutrition, Graduate School of Human Environment Science, Fukuoka Women’s University, Japan
3The Tsim Tsoum Institute, Krakow, Poland
4University College of Medical Sciences, New Delhi, India
5Fellow Wolfson Research Institute, School of Medicine Pharmacy and Health, Durham University, UK

Abstract: Approach: While the Western diet is proatherogenic, Paleolithic-style diet may be protective against Cardiovascular Diseases (CVDs). Results: Western diet is characterized with energy dense, refined, foods with a high glycemic index (e.g., refined starches; biscuits and bread) and unhealthy lipids (e.g., trans fats, saturated fat, omega-6 rich oils) poor in omega-3 fatty acids, phytochemicals and fibre. These diets are known to predispose inflammation and the epidemic of Non-Communicable Diseases (NCDs). CVD, diabetes mellitus, obesity, cancer and depression, are associated with increased production of Thromboxane A2 (TXA2), leucotrienes, prostacyclin, interleukins-1 and -6, tumor necrosis factor-alpha and C-reactive proteins in the tissues. Increased dietary intake of omega-6 fatty acids is known to enhance all these biomarkers which have adverse pro-inflammatory effects resulting in to CVDs. Functional food approaches including consumption of a Mediterranean diet rich in fruits, vegetables, nuts, canola oil, olive oil characterized with low omega-6/omega-3 ratio in the diet, as well as physical activity and meditation can modulate inflammation as well as body-mind interactions and may be protective against risk of CVD and all-cause mortality. Conclusion: Inflammation appears to be an important unifying hypothesis. In the absence of inflammation in the tissues, total cholesterol and other lipids may have neutral effects in the arterial tissues and myocardium. Therefore, the tissue is possibly the main issue for treatment.

Key words: Non-Communicable Diseases (NCDs), Cardiovascular Disease (CVD), Alpha-Linolenic Acid (ALA), Coronary Artery Disease (CAD), Saturated Fatty Acids (SFA)

INTRODUCTION

The focus of this review is based on the following premises and evidence, viz. (a) apparently non-significant evolutionary change in human genetic composition, that shaped human needs, over several recent millennia, (b) the dietary advice given by ancient Eastern physicians for the prevention of atherosclerosis and heart attacks, (c) the generally adverse dramatic change in dietary composition in modern times, presumably affecting gene regulatory mechanisms, giving rise to cardiovascular and related diseases that is supported by epidemiological and clinical trial evidence, (d) that the brain-body (liver, heart and gut axes), i.e., mind-body interactions play an important role in the genesis of proinflammatory responses (Takahashi et al., 2012) that a higher blood omega-6/omega-3 fatty acid ratio may adversely change membrane structures and associated signalling mechanisms. The cumulative evidence behind this review suggests that it is the latter point which merits further, perhaps novel, research on membranes structures in relevant end-organs. All these issues are discussed below.

The diets of Homo sapiens was characterized with natural foods; fruits, vegetables, green leaves, seeds, eggs, fish, meat from running animals and honey during the Paleolithic period (Eaton et al., 1988; 1998). These foods were also available to pre-agricultural humans which shaped modern human’s genetic nutritional
requirement (Simopoulos, 2003; 2009; Meester, 2008; 2009). Cereal grains (refined) and vegetable oils that are rich in omega-6 fatty acids and trans fats and low in amino acids are relatively recent addition to the human dietary patterns that represent dramatic departure from those foods and nutrients to which we are adapted (Simopoulos, 2003; 2009; Meester, 2008). Early man also had enormous physical activity without any mental stress. The role of fatty acids, essential and non-essential amino acids, antioxidants and vitamins in the prevention and pathogenesis of Cardiovascular Diseases (CVD), type 2 diabetes and insulin resistance are well known (Esposito and Glugliano, 2006; Katcher et al., 2008; Meester, 2009; Fung et al., 2009). There is evidence that the Mediterranean diet which has similarity with Paleolithic diet can influence brain function related to gut-liver-brain-heart axis indicating its influence on mind-body connection (Singh et al., 2008; Wang et al., 2008; Singh et al., 2011a; Wilson et al., 2011).

There have been marked changes in the food supply with the development of agriculture about 10,000 years ago from now. However, only a non-significant change in our genes occurred during the past 10 centuries, due to the presence of omega-3 fatty acids, amino acids, vitamins and antioxidants in the diet. However, now humans appear to live in a nutritional environment which completely differs from that for which our genetic constitution was selected (Eaton et al., 1988, 1998; Simopoulos 2003; Meester, 2008; 2009). Food consumption patterns have changed significantly during the last 100-160 years, causing increased intake of Saturated Fatty Acids (SFA), trans fat, refined carbohydrates and linoleic acid and decrease in omega-3 fatty acids, from grain-fed cattle, tamed at farm houses, rather than meat from running animals. In South East Asia, 60% of the mortality occurs due to Non-Communicable Diseases (NCDs) where diet and lifestyle may be responsible for these deaths (Dans et al., 2011). Apart from hyperlipidemia and hyperglycemia, oxidative stress and inflammation appear to be important mechanisms in the pathogenesis and prevention of diet related NCDs (Singh et al., 1992a; Esposito et al., 2004; Vogel, 2006; Singh et al., 2009).

Adverse effects of the diet were known to Indians from the ancient times, which are evident from the following verse from an ancient scripture Bhagwatgeeta (Bhagavad-Gita, 1998). “Foods which are bitter, acid, salted, burnt, fried and pungent, give rise to pain, mental stress and diseases” (3100 BC). Charaka (600 BC), a great physician of India, knew about the role of diet and lifestyle in the pathogenesis of heart attack, which would be clear from the following verse. “Heart attack is born by the intake of fatty meals, overeating, excess of sleep, lack of exercise and anxiety”, Charaka Sutra, 600BC.

Sushruta (600 BCE) (Bhishagratna, 2006) who was a surgeon from Vishwamitra family, gave a more clear description of atherosclerosis or madroga; “Excess intake of fatty foods and lack of exercise causes obesity and narrowing of the channels taking blood to the heart. It is useful to use googol, triphala and silajit in the treatment”. These herbs are known to have high content of antioxidant flavonoids, vitamins and minerals as well as fibres.

About 2000 years ago (1st century), Confucius (Hsu, 1974), the Chinese philosopher taught his students, “the higher the quality of foods, the better and never rely upon the delicacy of cooking”. Thus a dietary guideline based on experience, observation and thinking was given as: “cereals the basic, fruits the subsidiary, meat the beneficial and vegetable the supplementary”. Therefore, according to WHO experts (Anon, 1990), the concept of eating, a diet high in animal foods and preference for meat and greasy foods was shaped in China.

However, possibly the meat was rich in omega-3 fatty acids without any trans fat and w-6 fat and the total fat intake remained within desirable limits and was not excessive as in the West. Therefore, this advice by the Chinese thinker appears to be profound.

**Food and nutrient intake during Paleolithic period:**

The food and nutrient intake among hunter-gatherers and among Western and Asian populations show marked reduction in the consumption of omega-3 fatty acids, vitamins, antioxidants and amino acids and significant increase in the intakes of carbohydrates, (mainly refined), fat (saturated, trans fat and linoleic acid) and salt compared to the Paleolithic period (Tables 1-5). The protein or amino acid intake was 2.5 fold greater (33 vs. 13%) in the Paleolithic diet compared to modern diet (Table 3). Approximately 10,000 years ago, prior to the Agricultural Revolution, our diet was based on an enormous variety of wild plants. However, today about 17% of plant species provide 90% of the world’s food supply which is mainly contributed by grains. Wheat, corn and rice account for three fourths of the world’s grain production on which humans are dependent for food supply. Grains are high in omega-6 fatty acids and carbohydrates and low in omega-3 fatty acids and antioxidants compared to leafy green vegetables. Eaton and co-workers (Eaton et al., 1988; 1998) have estimated higher intakes for protein, calcium, potassium.
and ascorbic acid and a lower intake of sodium in the diet of the late Paleolithic period than the current diets in the developed and developing countries. Green leafy vegetables are also rich sources of antioxidants, magnesium, omega-3 fatty acids and carotenoids. These appear to be high in the Paleolithic diet. The proteins which were from both animal (running animals and eggs) and plant sources (green leaves and seeds), should have been composed of both essential and non-essential amino acids (Table 1 and 3).

Fatty acids in the diet and development of CVDs and diabetes: There has been an enormous increase in omega-6 fatty acid (about 30 g day$^{-1}$) in the diet due to the production of oils from vegetable seeds such as corn, sunflower, saiflower, soybean and cotton. Increased intake of meat has resulted in greater intake of arachidonic acid (0.2-1.0 mg day$^{-1}$), whereas the consumption of Alpha-Linolenic Acid (ALA) has decreased (about 0.55 g day$^{-1}$) and the amounts of Eicosapentaenoic Acid (EPA) and Docosahexaenoic Acid (DHA) are 48 and 72 mg day$^{-1}$ respectively (Table 4-6). A relative and absolute decrease in omega-3 fatty acids has led to an imbalance and increase in the ratio of omega-6/omega-3 fatty acids to up to 50 in South Asia and other developing countries, consuming vegetable seed oils (corn, soyabean, saiflower, sunflower, cotton) (Lorgeril et al., 1994; Harper and Jacobson, 2005; Gal et al., 2008; Solfi et al., 2010). Saturated Fatty Acids (SFA) and Trans Fatty Acids (TFA) elevate, PUFA decrease and mono-unsaturated fats MUFAs have beneficial effects on total and low Density Lipoprotein Cholesterol (LDL) as well as on HDL cholesterol. Omega-6 PUFA and TFA also decrease HDL cholesterol and increase insulin resistance, free radical stress and inflammation, which may enhance atherosclerosis (Lorgeril et al., 1994; Singh et al., 2002; Harper and Jacobson, 2005). Increased intake of total fat, TFA, SFA and omega-6 fatty acids and refined carbohydrates, may cause insulin resistance resulting in metabolic syndrome (Meester, 2008; Singh et al. 2008). Decreased intake of MUFA (Singh et al., 1996; 1999; 2000; Aratti et al., 2004), omega-3 fatty acids (Singh et al., 2011c; Pella et al., 2003; Simopoulos 2003; Kartikey et al., 2010), fibre and phytochemicals, may enhance the metabolic syndrome, leading to CVDs and other chronic diseases (Lorgeril et al., 1999; Kang et al., 2004; Iso et al., 2006; Meester, 2009; Eaton, 2009).

### Table 1: Food and nutrient intake among hunter-gatherer and Western populations

| Food and nutrient | Hunter-gatherer | Western population | Asians |
|-------------------|-----------------|--------------------|--------|
| Energy density    | Low             | High               | Low    |
| Protein           | High            | Low–moderate       | Low    |
| Animal            | High            | Moderate           | High   |
| Vegetable         | Very low        | Low–moderate       | Low    |
| Carbohydrate      | Low–moderate    | Moderate–rapidly absorbed | High–slow |
| High (>30g)       |                |                    |        |
| Fibre             | Low             | Low (<15g)         | Low    |
| Fat               | Low             | High               | Low    |
| Animal            | Very low        | High               | High   |
| Vegetable         | High(2.3g/day)  | Low (0.2g/day)     | 0.5–0.85g |
| Total omega-3     | Low 2.4         | High 15–20         | 25–50  |
| Ratio omega-6:omega-3 | High            | Low                | moderate |
| Vitamin and minerals |
| Linoleic acid     | 4.28            |                    |        |
| Alpha-linoleic acid | 11.4            |                    |        |
| Animal            | 4.56            |                    |        |
| Linoleic acids    | 1.21            |                    |        |
| Total             | 8.84            |                    |        |
| Linoleic acid     | 12.6            |                    |        |
| Alpha linoleic acid | 1.81            |                    |        |
| Long chain omega-3 fatty acids |
| Eicosapentaenoic acid(omega-3)(EPA) | 0.39 | |
| Docosatetraenoic acid(omega-6)(DTA) | 0.12 | |
| Docosapentaenoic acid(omega-3)(DPA) | 0.42 | |
| Docosahexaenoic acid(omega-3)(DHA) | 0.27 | |
| Total long chain omega-3 fatty acids | 1.20 | |
| Ratios of omega-6:omega-3 | 0.70 | |
| Linoleic acid/alpha linoleic acid+ DTA/EPA+DPA+DHA | 1.79 | |
| Total omega-6/omega-3 | 0.77 | |

### Note: Modified from Eaton et al. (1998) ref 4 and Singh et al. (2011c), Ref 6

### Table 2: Estimated fatty acid consumption in the late Paleolithic period

| Sources | Plants |
|---------|--------|
| Linoleic acid | 4.28 |
| Alpha-linoleic acid | 11.4 |
| Animal | 4.56 |
| Linoleic acids | 1.21 |
| Total | 8.84 |
| Linoleic acid | 12.6 |
| Alpha linoleic acid | 1.81 |
| Long chain omega-3 fatty acids |
| Eicosapentaenoic acid(omega-3)(EPA) | 0.39 |
| Docosatetraenoic acid(omega-6)(DTA) | 0.12 |
| Docosapentaenoic acid(omega-3)(DPA) | 0.42 |
| Docosahexaenoic acid(omega-3)(DHA) | 0.27 |
| Total long chain omega-3 fatty acids | 1.20 |
| Ratios of omega-6:omega-3 | 0.70 |
| Linoleic acid/alpha linoleic acid+ DTA/EPA+DPA+DHA | 1.79 |
| Total omega-6/omega-3 | 0.77 |

### Note: Modified from Eaton et al. (1998) ref 4 and Singh et al. (2011b), Ref 6

### Table 3: Nutrient composition in the late Paleolithic and current recommendations

| Nutrient per day | Late Paleolithic | Current recommendation |
|------------------|------------------|-----------------------|
| Total dietary energy |                |                      |
| Protein | 33               | 12                   |
| Carbohydrate | 46               | 58                   |
| Fat | 21               | 30                   |
| Alcohol | 0                | moderate alcohol     |
| P/S ratio | 1.41             | 1                    |
| Cholesterol, mg | 520              | 300                  |
| Fibre, g | 100–150          | 30–60                |
| Sodium, mg | 690              | 1100–3300            |
| Calcium, mg | 1500–2000        | 800–1600             |
| Ascorbic acid, mg | 440             | 60                   |

### Note: Modified from Eaton et al., 1988 their references 1, 2 and Singh et al., 2011c, their references 1,2,46
Paleolithic-style diet on cardiovascular functions and NCDs. Several of these studies were conducted when thrombolysis and angioplasty were not freely available for the treatment of Acute Coronary Syndrome (ACS) (Singh et al., 1992b; 1992c). Therefore it is not possible to conduct such studies again to demonstrate the role of diet in the prevention of ACS, although the present authors (Singh et al., 1995; 2011b; 2011c) and many other agencies now advise increased intake of fruits and vegetables for the prevention of CVDs (Anon 1988; 1990; 1991). A recent study reported that one micro RNA in particular, MIR 168a, which is highly enriched in rice, was found to inhibit a protein that helps remove Low-Density Lipoprotein (LDL) from the blood, suggesting that microRNAs can influence gene expression across kingdoms (Zhang et al., 2011). Diet can also provide beneficial or adverse effects by its influence on gut-brain-liver axis leading to increase or decline in CVDs (Gal et al., 2008; Wang et al., 2008). These gut-brain-liver studies and others (Pella et al., 2003; Simopoulos, 1999); Kang et al 2004; Meester, 2009; Eaton, 2009; Singh et al., 2010; Kartikey et al., 2010). There is an experimental study which showed that increased availability of omega-6 fatty acid in the tissues, enhances the formation of omega-3 fatty acids by the animals due to extraordinary capability of the animals to fight the adverse effects of omega-6 fatty acids (Kang et al., 2004).These studies, further emphasize that it is the tissue which is the main issue rather than associated risk factors for CVDs and other NCDs.

Epidemiological studies: Epidemiological studies indicate that a prudent dietary pattern characterized by fruit, vegetable, legume and whole grain intake appears to be protective. The protective effects of prudent dietary patterns appear to be due to a low omega-6/omega-3 ratio of such diets because Western diet and South Asian diets have high omega-6/omega-3 ratio of 20-50 as shown in recent studies (Pella et al., 2003; Meester, 2009; Eaton, 2009; Singh et al., 2010; Kartikey et al., 2011d). The INTERHEART study, involving participants from 52 countries (Iqbal et al., 2008) examined the relationship between dietary patterns and risk of Acute Coronary Syndrome (ACS). Consistent with previous studies in single within-population cohort studies, the authors found an inverse association between the prudent pattern score and risk of ACS and a significant positive association between the Western pattern score and increased risk of ACS. No association of Oriental diet with risk of ACS was reported. A dietary risk score based on 7 food items on the food-frequency questionnaire (meat, salty snacks, fried foods, fruits, green leafy vegetables, cooked vegetables and other raw vegetables) was constructed by the authors. The investigators found that a higher score, indicating a poor diet was strongly associated
with ACS risk and the subjects in the highest quartile of the score had nearly a 2-fold increased risk, even after adjustment for established coronary risk factors. On the basis of an arbitrary cut point of the score (top 3 quartiles versus the bottom quartile), the investigators estimated that 30% of Myocardial Infarction (MI) cases could be explained by unhealthy diets worldwide. The INTERHEART study is the first large study to quantify eating patterns in all geographic regions of the world. It provides evidence that despite different food habits in various populations, reproducible patterns can be found in diverse regions of the world. These findings are important because there has been a concern that dietary patterns derived through a data-driven approach such as Principal Components Analysis may be highly unstable and non-reproducible because of very different eating habits in different populations.

In a large, prospective, observational study (Heidemann et al., 2008) involving 72, 113 female nurses who were free of Coronary Artery Disease (CAD), stroke, diabetes and cancer, Factor Analysis identified 2 dietary patterns from data collected using serial food frequency questionnaires. One pattern, called prudent, was characterized by a high consumption of vegetables, fruit, legumes, fish, poultry and whole grains. The other pattern, called Western, corresponded to a high consumption of red meat, processed meat, refined grains, french fries, sweets and desserts. Individuals were classified by their level of adherence to both the prudent diet and the Western diet. After baseline data collection in 1984, follow-up lasted 18 years, during which time 6011 deaths occurred (3319 [52%] as a result of cancer; 1154 (19%) resulting from Cardiovascular Disease (CVD); and 1718 (29%) resulting from other causes).

There was a 17% lower risk of total mortality among those who were most adherent to the prudent diet (highest versus lowest quintile of adherence), a 28% lower risk of CVD mortality and 30% lower mortality from non-CVD, non-cancer causes. Cancer was not associated with the inverse prudent dietary pattern. A comparison of the highest and lowest quintiles of adherence showed that consumption of the Western diet was associated with increased total mortality (21%), CVD mortality (22%), cancer mortality (16%) and mortality from non-CVD, non-cancer causes (31%). Hence, except for cancer, risk relationships for the prudent and Western dietary patterns appear to be the inverse of each other. Mortality thus was increased as adherence to the prudent diet decreased and adherence to the Western diet increased. In one cross-sectional survey of 6940 subjects, above 25 years of age, fruit, vegetable and legume intake were inversely associated with risk of pre-hypertension and hypertension in five Indian cities (Singh et al., 2011c).

A meta-analysis of cohort studies, quantitatively assessed the relation between fruit and vegetable intake and incidence of CAD which reported Relative Risks (RRs) and corresponding 95% Confidence Interval (CI) of CAD with respect to frequency of fruit and vegetable intake (He et al., 2007). A total of 278 459 subjects (9143 CAD events) were included, with a median follow-up of 11 years. The individuals who had less than 3 servings/day of fruit and vegetables, the pooled RR of CAD was 0.93 (95% CI: 0.86-1.00, p = 0.06) whereas those with more than 5 servings/day, the RR was 0.83 (0.77-0.89, p = 0.0001). This meta-analysis of prospective cohort studies demonstrated that increased consumption of fruit and vegetables, less than 3 to more than 5 servings/day is related to a 17% reduction, whereas increased intake to 3-5 servings/day is associated with a smaller and borderline significant reduction in CAD risk. These results provide evidence supporting that 5 or more servings per day of fruit and vegetables, are needed to protect from CVD.

| Foods and nutrients                  | Paleolithic-style diet group (n = 204) | Standard diet group (n = 202) |
|--------------------------------------|---------------------------------------|-------------------------------|
|                                      | 4-7 days                              | After 1 year                  | 4-7 days                              | After 1 year                  |
| Fruits and vegetables (g. Day⁻¹)     | 508.4(28.66)**                        | 575(91.4)**                   | 254.4(17.2)                           | 220.3(19.6)                   |
| Potato, radish,                      | 60.5(6.8)                             | 115(12.7)**                   | 72.0(12.5)                            | 155.6(32.5)                   |
| Legumes and pulses (g. Day⁻¹)        | 80.5(6.6)**                           | 95.0(8.9)**                   | 52.5(4.6)                             | 45.6(5.6)                     |
| Almonds and walnuts (g. Day⁻¹)       | 82.4(5.7)**                           | 75.5(5.2)**                   | -                                     | -                            |
| Fish (g. Day⁻¹)                      | 52.5(6.5)**                           | 22.4(4.1)**                   | 20.2(3.1)                             | 10.5(3.5)                     |
| Chicken (g. Day⁻¹)                   | -                                     | 10.2(3.2)*                    | 76.2(6.5)                             | 66.5(10.5)                    |
| Mustard or soybean oil              | 18.4(3.9)*                            | 31.5(5.5)**                   | 10.5(2.3)                             | 6.8(2.8)                      |
| Butter or clarified butter (g. Day⁻¹)| 2.5(0.6)**                            | 3.30(0.71)**                  | 10.5(2.6)                             | 12.6(3.5)                     |
| Skim milk (ml day⁻¹)                 | 161.2(12.0)                           | 152(14.5)*                    | 150.2(8.0)                            | 165.5(16.1)                   |
| Wheat chapatti                       | 5.5(1.6)**                            | 30.6(5.5)                     | 50.6(6.6)                             | 55.6(7.8)                     |
| Bread, biscuits (g. Day⁻¹)           | 10.6(2.2)*                            | 25.5(6.2)**                   | 230.6(20.1)                           | 212.2(18.1)                   |
| Rice and wheat cereals (g. Day⁻¹)    | 25.6(2.4)                             | 30.6(5.5)                     | 30.2(3.1)                             | 35.6(4.8)                     |
| Honey or raisins (g. Day⁻¹)          | 2.6(0.8)                              | 5.5(1.2)                      | -                                     | -                            |
| Sugar (g. Day⁻¹)                     | 16.4(3.7)*                            | 12.6(3.4)*                    | 25.5(5.4)                             | 30.5(7.6)                     |
| Total Adherence score (%)            | 65.2(17.2)                            | 63.9(14.8)                    | 123.0(30.0)                           | 71.0(30.0)                    |
| Total foods                          | 1184.6(254)                           | 983.4(213)                    | 862(204)                              | 862(204)                     |

Note: P values for mean (standard deviation) were obtained by comparison of intervention and control groups after 1 week and after 1 year, * = p<0.05, ** = p<0.01, Singh et al., 2012 their reference 40

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Dietary-patterning analysis has been increasingly used recently, as an alternative method to traditional single-nutrient analysis because it can assess cumulative effects of the overall nutrients in the diet. Habitual intake patterns are typically quantified by statistical methods such as Factor or Cluster Analysis or diet-quality indexes based on prevailing dietary recommendations or traditional diets e.g., the Mediterranean diet (Renaud et al., 1995; Lorgeril et al., 1994; Sofi et al., 2010), the Japanese diet (Iso et al., 2006) and the Indo-Mediterranean diet (Singh et al., 2002; Pella et al., 2003). Principal Components Analysis is commonly used to define dietary patterns using food consumption information to identify common underlying dimensions (factors or patterns) of food intake. The method aggregates specific food items based on the degree to which these food items are correlated with each other. A summary score for each pattern is then derived and can be used to examine relationships between various eating patterns and outcomes of interest such as CAD, diabetes mellitus, stroke and other chronic diseases. Earlier validation studies found that 2 major patterns (the prudent and Western patterns) identified through Principal Components Analysis of food consumption data assessed by food frequency questionnaires were reproducible over time and correlated reasonably well with the patterns identified from diet records. The consistent association observed between the Western or unhealthy dietary pattern (high in animal products, salty snacks, refined starches and sugar and fried foods and low in fruits and vegetables) and ACS risk in different regions of the world from the INTERHEART study and other studies as well as in our study, provide consistent evidence of the adverse effects of globalization on human nutrition and chronic disease risk. However, this evidence is indirect because these studies did not specifically assess the impact of global trade and marketing on food consumption patterns across different countries (Renaud et al., 1995; Knoops et al., 2004; He et al., 2007; Heidemann et al. 2008; Hu et al. 2008; Iqbal et al., 2008; Singh et al., 2011c). Despite this weakness, most recent studies suggest that the current trend of dietary convergence toward a typical Western diet characterized by high omega-6/omega-3 ratio of fatty acids is likely to play a role in the globalization of obesity, CVD, diabetes and cancer.

**Intervention trials on low omega-6/omega-3 fatty acid ratio Paleolithic-style diet and mortality:** Cohort studies provide an association of diet with risk of CVDs and deaths. However, randomized, controlled intervention trials are necessary to provide a scientific proof that diet has a role in the prevention of CVDs (Singh et al., 2002; Gal et al., 2008; Sofi et al., 2010). Intervention trials, using the whole diet approach so far produced are also in line with this epidemiological evidence. The effect of Paleolithic-style diet was examined in patients (n = 204 intervention group, n = 202 control group) with acute coronary syndromes, which showed significant decline in total cardiac events as well as in total mortality after 6 weeks and the benefit continued after one year (Singh et al., 1992b; 1992c). Further follow up for 2 years in this study (Singh et al., 2012) is different from the published work, because its emphasis is on the Paleolithic dietary patterns and ALA content of the diet to be responsible for the significant greater survival in the intervention group compared to control group (Table 6 and 7). Dietary patterns before entry to the study showed a higher omega-6/omega-3 ratio of 3.25 in the diets of both the groups. Intervention group A was advised a Paleolithic style diet with omega-6/omega-3 fatty acid ratio of 4.3 compared to standard diet group with ratio of 20 (Table 6 and 7). After a follow up of 2 years, total mortality was significantly declined in the Paleolithic style diet group compared to control group as shown in Fig. 1 for the intervention group compared to control group (85.3% vs. 74.8%, p<0.001). The mortality was lowest among subjects with omega-6/omega-3 ratio of less than 10 which showed graded increase with increase in the fatty acid ratio in both the groups as shown in Fig. 2.

In the Lyon diet heart study 605 patients who had a myocardial infarction were randomly assigned to a ‘Mediterranean-style’ diet or a control diet resembling the American Heart Association Step I diet.
Table 8: Total cardiovascular events in the Paleolithic-style diet and control group

| Data                              | Indo-Mediterranean diet (n = 499) | Standard Diet (n = 501) |
|-----------------------------------|-----------------------------------|-------------------------|
| Non fatal myocardial infarction   | 21 (4.2%)*                        | 43(8.6%)                |
| Fatal myocardial infarction       | 12 (2.4%)                         | 17(3.4%)                |
| Sudden cardiac death              | 6(1.2%)                           | 16(3.2%)                |
| Total cardiac events              | 39(7.8%)**                        | 76(15.2%)               |
| Stroke                            | 7(1.4%)                           | 13(2.6%)                |
| Stroke death                      | 2(0.4%)                           | 3(0.6%)                 |
| Total cardiovascular events       | 48(9.6%)**                        | 92(18.3%)               |
| Total deaths                      | 24(4.8%)                          | 38(8 %)                 |

Note: Values are number (%), *= p<0.01, **= p<0.001, Singh et al. (2002), their references 35, 50.

The Mediterranean diet model supplied 30% of energy from fats and <10% of energy from saturated fatty acids, whereas the intake of 18:3 (n-3) (α-linolenic acid) provided >0.6% of energy. After a mean follow-up of 27 months, the risk of new acute myocardial infarction and episodes of unstable angina was reduced by about 70% by the Mediterranean diet. Moreover, total mortality was also reduced by 70%. Long-term follow up for 4 years also showed that the beneficial effects of diet were continued.

Singh et al., 2002 tested an ‘Indo-Mediterranean diet’ in 1000 patients in India, with existing coronary disease or at high risk for coronary disease. Half of the patients (n = 499 vs. 501) were administered a diet rich in fruits, vegetables, whole grains, walnuts, mustard and soy bean oil as a source for omega-3 fat and the rest (501) patients were advised to take prudent diet advised by the National Cholesterol Education Program step 1 diet in 1988 (Anon 1988). At the end of 2 year follow up, the Paleolithic-style diet group consumed significantly more fruits, vegetables and legumes than did the control group (537±127 vs. 231±19 g day⁻¹, p<0.001) as well as more mustard and soy bean oil (31±6.5 vs. 15.2±5.5 g day⁻¹). The mean intake of ALA was over two fold greater in the Paleolithic-style diet group compared to control group. (1.8±0.4 vs. 0.8±0.2 g day⁻¹, p<0.001). The omega-6/omega-3 ratio of fatty acids was slightly higher at baseline in the intervention group than in the control group (39±12 vs. 34±10) yet both these values are extremely high, reflecting a diet with a very high omega-6 content yet low omega-3 (Pella et al., 2003). At the end of two years follow up, this ratio showed a marked decline in the intervention group, which was greater than that observed in the control group consuming control diet (9.1±12 vs. 21±10, p<0.001). The study endpoints were: significant decline in the total cardiac events, sudden cardiac death and non-fatal infarction in the intervention group compared to the control group (Table 8).

Esposito et al., 2004 randomized 180 patients (99 men, 81 women) with metabolic syndrome to a Mediterranean style diet, characterized with whole grains, vegetables, fruits, nuts and olive oil vs. a cardiac-prudent diet with fat intake <30%. After a follow up of 2 years, subjects in the intervention diet showed greater weight loss, had lower C-reactive protein and proinflammatory cytokine levels, had less insulin resistance, as well as lower total cholesterol and triglycerides and higher HDL cholesterol. The prevalence of metabolic syndrome was reduced to one half. The Japan Public Health Centre based study (Iso et al., 2006) showed that eating more omega-3 fatty acids by increased intake of fish was associated with significant reduction in cardiovascular disease and cardiac mortality. The diet and re-infarction trial (Burr et al., 1989) showed that modest intake of fish, 2 servings per week, can cause significant a decrease in total mortality and cardiovascular mortality.
Since no benefit was observed in non-fatal infarction, the authors concluded that omega-3 fatty acids may have prevented ventricular fibrillation by altering cardiomyocyte cell membrane phospholipids. There is experimental evidence indicating that the very long chain omega-3 fatty acids in fish oil and fatty fish have an important effect on the pathogenesis of arrhythmias in the setting of myocardial ischemia and reperfusion both in-vivo and in-vitro (McLennan et al., 1988). There is additional evidence from other studies indicating the role of omega-6/omega-3 ratio of fatty acids in the pathogenesis of Non-communicable diseases (Endres et al., 1989; Kumar et al., 1992; Simopoulos, 1994; 2006; 2008; Appel, 2008).

There is clear evidence that dietary changes or omega-3 fatty acid supplementation and decrease in omega-6 fatty acid intake may induce a marked alteration in the omega-6/omega-3 fatty acid ratio in tissue. For example, a value of omega-6/omega-3 ratio of 1-2:1 has been shown to be the ratio found in the traditional diet of Cretans and it was this study that led to the concept of a Mediterranean diet and its relation to a Paleolithic-style diet discussed in this review. Consequently, the myocardial and arterial tissue may, through conditioning by the nutrients of a Paleolithic-style diet, be protected against the otherwise proinflammatory risk of a generalized Western diet.

Several cohort studies and intervention trials suggest that increased consumption of functional foods like fruits, green leafy vegetables, nuts and legumes decreases the risk of CVD morbidity and mortality. Potentially protective contents of these foods include ALA, EPA, DHA, folate, magnesium, calcium, potassium, fibre, vitamin E, carotenoids, arginine, cysteine, oleate and favorable lysine to arginine and methionine to arginine ratios. These nutrients are important in the cardiomyocyte and arterial cell functions which are determinant of CVDs. The lysine to arginine ratio is potentially involved in atherogenesis and methionine to arginine ratio is important for endothelial function because arginine is the precursor of NO which protects the endothelium and methionine is the precursor of homocysteine that is known to cause endothelial damage. Folic acid is protective because it antagonizes the homocysteines.

Calcium/magnesium ratio indicating high magnesium is also protective to cardiomyocyte and endothelial cells. These findings indicate that it is the tissue which is the main issue in the pathogenesis and prevention of CVDs. If the tissue concentration of omega-6/omega-3 ratio is 1:1 along with other nutrients in proper ratio, high LDL cholesterol (unoxidised) would be neutral without any evidence of endothelial dysfunction. This concept has become more relevant because microRNA have been demonstrated in the blood and milk after feeding of rice to experimental animals (Zhang et al., 2011). Therefore, for prevention of CVDs, eat 400 g day$^{-1}$ of fruits, vegetables and nuts and another 400 g day$^{-1}$ of legumes and other whole grains along with 30-50g of canola oil+ olive oil to protect our tissues which is the major issue.

**CONCLUSION**

In summary, this review suggests that the prevention of CVDs can be substantially achieved by consuming a notional Paleolithic or Mediterranean-style diet such as the daily consumption of suitable proportions of fruit, vegetables and nuts, legumes and canola and olive oil. Furthermore, it has focused on the need for more research on membrane and signaling pathways in end-organ tissue and the brain.

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