Medical cost savings in Sakado City and worldwide achieved by preventing disease by folic acid fortification

Yasuo Kagawa¹, Mami Hiraoka², Mitsuyo Kageyama³, Yoshiko Kontai⁴, Mayumi Yurimoto⁵, Chiharu Nishijima¹, and Kaori Sakamoto¹
¹Department of Medical Chemistry, Kagawa Nutrition University, Sakado City, ²Shukutoku University, College of Nursing and Nutrition, School of Nutrition, Chiba City, ³Yamanashi Gakuin University Faculty of Health and Nutrition, Kofu City, ⁴University of Niigata Prefecture Faculty of Human Life Studies Department of Health and Nutrition, Niigata City, and ⁵Yakult Honsha Co., Ltd., Tokyo, Japan

ABSTRACT The introduction of mandatory fortification of grains with folate in 1998 in the United States resulted in 767 fewer spina bifida cases annually and a cost saving of $603 million per year. However, far more significant medical cost savings result from preventing common diseases, including myocardial infarction, stroke, dementia and osteoporosis. A cost-effectiveness analysis showed a gain of 266 649 quality-adjusted life-years and $3.6 billion saved annually, mainly due to the reduction of cardiac infarction. The recommended folate intake in Japan is 240 µg/day whereas it is 400 µg/day internationally. Our Sakado Folate Project targeted individuals with genetic polymorphism of methylentetrahydrofolate reductase or with hyperhomocysteinemia. Using, for example, folate-fortified rice, resulted in an increase in serum folate and a decrease in serum homocysteine in the participants, and reduced medical costs were achieved by decreasing myocardial infarction, stroke, dementia and fracture. Due to the small population of Sakado City (approximately 101 000) and small number of births (693) in 2015, a decrease in spina bifida could not be confirmed but there was a significant decrease in the number of very low birthweight infants. The genome notification of subjects was effective in motivating intake of folate, but the increase in serum folate (from 17.4 to 22.5 nmol/L, 129%) was less than that observed following compulsory folic acid fortification of cereals in the USA (from 12.1 to 30.2 nmol/L, 149.6%). Mandatory folic acid fortification is cheap in decreasing medical costs and is thus recommended in Japan.

Key Words: folate, fortification, genetic polymorphism, medical cost

INTRODUCTION

The authors reviewed the medical literature reporting key studies on folic acid fortification, policy recommendations from several foreign organizations, and our own Sakado Folate Project, and drew conclusions regarding the decrease in the incidence of disease and the medical cost savings after the folate intake of the population is increased. Folate refers to a collection of “folates” that is not chemically well-characterized and includes other members of the pteroylglutamate family. These family members are differentiated by the reduced state of the pteridine ring, a one carbon substitution at the N5 and/or N10 positions (formyl, methyl, methylene, and methenyl), and the length of the γ-glutamyls. Polyglutamyl 5-methyltetrahydrofolate species are the most abundant naturally occurring folate in vegetables, and 5-formyltetrahydrofolate species are the second most abundant (Wang et al. 2010). The bioavailability of synthetic folic acid (monoglutamyl pteridine) is approximately 70% better than that of folate (mainly polyglutamyl pteridines) from dietary sources and thus the term dietary folate equivalent (DFE) is used to describe the effects of different folate molecular species (Joint FAO/WHO Expert Consultation on Human Vitamin and Mineral Requirements 1998). Folate prevents megaloblastic anemia (Chanarin 1987) and many other diseases because it is essential to the synthesis, repair (Lindahl 2013), and epigenetic regulation (Janke et al. 2015) of DNA. Folate is therefore the link between diet, metabolism, and epigenetics (Janke et al. 2015). Thus, the mechanism by which folate prevents many diseases, such as neural tube defects (NTD), has been studied at the molecular level (Ichi et al. 2012). NTDs are the second most common group of serious birth defects. Each year, 300 000 to 400 000 infants worldwide are born with NTDs. NTDs result from failure of the neural tube to close properly approximately 28 days post-conception. Two of the most common NTDs are spina bifida and anencephaly. One of the earliest recommendations on folate intake was the Recommendations of the Clinical Teratology Committee of the Canadian College of Medical Geneticists, which stated that primary prevention of NTDs is preferable to treatment or to prenatal detection and abortion (Van Allen et al. 1993). The use of biomarkers such as homocysteine to detect folate deficiency has been described in detail by Bailey et al. (2015). There is now conclusive evidence that most NTDs can be prevented by the ingestion of folic acid near the time of conception (Joint FAO/WHO Expert Consultation on Human Vitamin and Mineral Requirements 1998). Thus, mandatory folic acid fortification of cereal products has been implemented in 82 countries. Mandatory fortification has led to medical cost savings due to the prevention not only of NTDs (Grosse et al. 2016), but also of many other diseases described in the next section (Bentley et al. 2009).

DISEASES PREVENTED BY FOLIC ACID FORTIFICATION

Megaloblastic anemia (Chanarin 1987) and NTDs (Grosse et al. 2016) are well-known conditions caused by inadequate folate intake, but folate is implicated in many other diseases. For example,
moderate hyperhomocysteinemia induced by low folate status is an independent risk factor for cardiovascular disease (CVD) (Ishihara et al. 2008; Bentley et al. 2009), stroke (Yang et al. 2006; Huo et al. 2015; Li et al. 2016b), colon cancer (Bentley et al. 2009), dementia (Jernérén et al. 2015), fracture (Gjesdal et al. 2007), kidney diseases (Massy 2003) and depression (Bottiglieri 2005; Hiraoka et al. 2014). These diseases can be prevented or alleviated by increased folate intake, as shown by many epidemiological and interventional studies. The meta-analysis of 30 randomized controlled trials involving 82,334 participants indicated a 10% lower risk of stroke and a 4% lower risk of overall CVD with folic acid supplementation (Li et al. 2016b). Folic acid supplementation provided a greater benefit for preventing CVD in participants with low plasma folate levels and without preexisting CVD, and resulted in larger decreases in homocysteine levels (Li et al. 2016b). Folate is required by the brain for the synthesis of monoamine neurotransmitters, and numerous observational studies have suggested an association between folate deficiency and depression (Bottiglieri 2005). Folate intake below the Recommended Daily Allowance at study baseline was associated with increased risk of the incidence of Mild Cognitive Impairment (MCI)/probable dementia (hazard ratio 2.0, 95% CI 1.3 to 2.9) after controlling for multiple confounders (Agnew-Blais et al. 2015). Plasma homocysteine concentration, if only moderately elevated, is an independent risk factor for CVD and stroke (Joint FAO/WHO Expert Consultation on Human Vitamin and Mineral Requirements 1998). Increased risk has been associated with values higher than 11 mmol/L, which is well within what is generally considered to be the normal range (5–15 mmol/L) of plasma homocysteine levels (Joint FAO/WHO Expert Consultation on Human Vitamin and Mineral Requirements 1998). Randomized control tests on the prevention of CVD and dementia with folate often result in negative data. However, these diseases develop very slowly, and during a test that lasts only a few years, endothelial cells or neurons are either already damaged, or the disease has progressed insufficiently for necrosis to have developed.

GENETIC POLYMORPHISMS OF FOLATE METABOLISM

Serum folate and total homocysteine (tHcy) levels are influenced by both folate intake and genetic polymorphisms. There are several single nucleotide polymorphisms (SNPs) that affect folate metabolism, including the 5, 10-methylenetetrahydrofolate reductase (MTHFR) gene C677T. The presence of the C677T mutation in the MTHFR gene (rs1801133) has been regarded as a genetic risk factor for coronary artery diseases and neural tube defects (Li et al. 2016b). The frequency of the T allele of MTHFR is 36.9%, 41.7%, 58.6% and 8.1%, in Japan, the USA, Mexico and Indonesia, respectively (Sadewa et al. 2002). The authors also tested the following high frequency SNPs affecting folate and homocysteine metabolism (Hiraoka et al. 2004): MTHFR A1298C, CBS844ins68 (cystathionine-β-synthase), MS A2756G (methionine synthase) (Cai et al. 2014), MTRR A66G (methionine synthase reductase) (Zhi et al. 2016), and RFC-1 G80A (reduced folate carrier-1) (Li et al. 2016a). CBS844ins68 is the 8th exon insertion mutation and comprises 68 base pairs (Cai et al. 2014). However, these SNPs, except for MTHFR C677T, did not significantly affect serum Hcy, folate, vitamin B6 and vitamin B12 levels (Hiraoka et al. 2004). The frequencies of mutant homozygotes were lower in MTHFR A1298C (1.3%), MS A2756G (5.3%) and MTRR A66G (8.5%) than in MTHFR C677T (17.3%) and RFC-1 A80G (18.6%), and thus may not affect the overall incidence of disease in large populations. In contrast, the TT genotype of MTHFR C677T showed significantly lower serum folate and higher serum Hcy than the CC and CT genotypes (Hiraoka et al. 2004). The TT genotype of the C677T polymorphism in MTHFR has been associated with CVD (Morita et al. 1998), dementia (Kageyama et al. 2008), osteoporosis (Gjesdal et al. 2007), depression (Bottiglieri 2005) and kidney diseases (Sakamoto et al. 2015). In addition, there are many risk SNPs of CVD, metabolic syndrome, and other chronic diseases (Kagawa 2012; Kagawa et al. 2012) that affect the MTHFR genotypes and must be considered for disease prevention and the reduction of medical costs.

Information obtained from the above studies on SNPs indicates that folate intake sufficient to reduce plasma Hcy to a minimum level of less than 7.0 μmol/L is required to reduce disease risk (Joint FAO/WHO Expert Consultation on Human Vitamin and Mineral Requirements 1998). In Japan, however, a recommendation for pregnant women to take a folate supplement was made by the Ministry of Health, Labor and Welfare only in 2000. The possible benefit of lowering plasma homocysteine through increased folate intake can be proven only by an intervention trial with folic acid supplementation in large populations. Using plasma homocysteine as a biomarker for folate adequacy can only be done on an individual basis after the possibility of a genetic mutation or an inadequate dietary intake can be proven only by an intervention trial with folic acid supplementation in large populations. The prevention of CVD and the medical cost savings expected from folic acid supplementation was also estimated (Homburger 1998). Bayesian statistical and decision-analytic techniques were used to estimate the cost–benefit and sample size of a placebo-controlled trial of folate targeted to US citizens aged 35 to 84 years with elevated serum homocysteine levels. The main end point is event-free survival (i.e., survival without new ischemic heart disease or stroke) at 5 years. Because the screening and other costs associated with this study were small compared with the consequences of stroke, ischemic heart disease, or death, the increase in 5-year event-free survival with folate that should compel the use of

MEDICAL COST SAVINGS RESULTING FROM FOLIC ACID FORTIFICATION

Folic acid supplementation of the expectant mother’s diet should result in fewer NTDs among infants and ancillary savings in medical costs (Van Allen et al. 1993). A minimum dosage of folic acid of 0.8 mg/day and a maximum dosage of 5.0 mg/day was recommended (Van Allen et al. 1993), along with a well-balanced, nutritious diet for all women who are at increased risk of having offspring with NTDs and who are planning a pregnancy. The results from economic evaluations demonstrate that folic acid fortification of food and preconception folic acid consumption are cost-effective ways to reduce the incidence and prevalence of NTDs (Yi et al. 2011). The annual direct medical cost per patient was estimated to be €42,943 ($51,574 in 2003) for NTD and between €11,728 ($11,061 in 1993) to €54,270 ($65,177 in 2003) for spina bifida in the USA (Yi et al. 2011). The prevention of CVD and the medical cost savings expected from folic acid supplementation was also estimated (Homburger 1998). Bayesian statistical and decision-analytic techniques were used to estimate the cost–benefit and sample size of a placebo-controlled trial of folate targeted to US citizens aged 35 to 84 years with elevated serum homocysteine levels. The main end point is event-free survival (i.e., survival without new ischemic heart disease or stroke) at 5 years. Because the screening and other costs associated with this study were small compared with the consequences of stroke, ischemic heart disease, or death, the increase in 5-year event-free survival with folate that should compel the use of

© 2017 The Authors. Congenital Anomalies published by John Wiley & Sons Australia, Ltd on behalf of Japanese Teratology Society.
Cost savings by folic acid fortification

The event-free survival rate for the placebo and folate groups were assumed to lie, with 95% probability, between 83.7% and 87.7%, and 82.9% and 90.7%, respectively. Such a trial would provide an expected societal cost–benefit savings exceeding $11 billion within 15 years (Hornberger 1998).

In 1998, compulsory folic acid fortification was started in the US and Canada, and the FDA mandated that all enriched cereal grain products be fortified with folic acid at a level of 140 μg/100 g of cereal product (Ganji and Kafai 2006). The exact human test using 13C- and 14C double labeled folate fortified cereals established this fortification level (Pfeiffer et al. 1997). This fortification mandate is estimated to have reduced the annual number of US live birth spina bifida cases by 767 (Grosse et al. 2016). The present value of mean direct lifetime cost per infant with spina bifida is estimated to be $791 900, excluding caregiving costs. Folic acid fortification is estimated to reduce the present value of total direct costs for each year’s birth cohort by $603 million above the cost of fortification (Grosse et al. 2016). The present value of mean direct lifetime costs of spina bifida by 767 (Grosse et al. 2016). The present value of mean direct lifetime costs of spina bifida per newborn born alive in the US was $9 080 (Nagao et al. 2004). The expected societal cost savings by folic acid fortification for a

OUTLINE OF THE SAKADO FOLATE PROJECT

The Japanese RDA is insufficient for persons with polymorphism in genes involved in folate metabolism and in aged persons with decreased folate bioavailability. The average folate intake in 2014 was 284 μg/day, which is higher than the Japanese RDA of 240 μg/day for adults (Ministry of Health, Labor and Welfare 2014). However, the folate intake of women aged 20–39 years was only 243–253 μg/day in 2014, which is much less than the RDA of 480 μg/day for pregnant women (Ministry of Health, Labor and Welfare 2014). There are three reports on the folate intake of Japanese pregnant women: 289 ± 150 μg/day (Mito et al. 2007), 336 ± 161 μg/day (Nakano and Ishii 2004) and 283 μg/day (Kondo et al. 2013, standard deviation not clear, the value of 2011). These values are much less than the RDA of 480 μg/day. The folate intake of 50% of the poor pregnant women was less than 260 μg/day (Onda R, reported in the Japanese Society of Nutrition and Dietetics, 2015). The folate intake higher than these levels among Japanese was quite effective in preventing cardiovascular diseases was established by Japan Collaborative Cohort Study on a total of 23 119 men and 35 611 women, age 40 to 79 years (Cui et al. 2010). Increased dietary folate intake from < 272 μg/day to > 536 μg/day resulted in 51% reduction of mortality for men from heart failure (P < 0.01) and that from ischemic heart diseases (down to 72%, P < 0.05), and for women ischemic heart diseases (down to 56% P < 0.01). These inverse associations did not change materially after adjustment for cardiovascular risk factors, so, folate intake of 243–253 μg/day by general Japanese is not enough to prevent cardiovascular diseases and thus more folate intake will proportionally reduce the medical costs.

Moreover, the bioavailability of folate is low in persons with genetic polymorphism (Hirao et al. 2004). About 15% of Japanese with the TT genotype of MTHFR C677T require 400 μg/day of folate to increase the serum folate level to that of CC and CT genotypes. Folate deficiency among persons with dementia was confirmed by meta-analysis of 31 studies (Lopes da Silva et al. 2014) and of Japanese dementia studies (Kageyama et al. 2008). The risk of brain infarction is 3.4-fold higher in persons with the TT type compared to persons with the CC type (Morita et al. 1998). The randomized controlled double blind test was performed to confirm the exact folate requirement of Japanese (Hirao et al. 2014).

To prevent the above-described diseases caused by folate deficiency, Sakado City in Japan made an agreement with Kagawa Nutrition University called the “Sakado Folate Project” in 2006. The Project offered participants a lecture, genotyping, blood analysis, nutrition surveys, and guidance according to the collected data (Fig. 1). The lecture was similar to that given in the Biomarkers of Nutrition for Development (BOND) project and was designed to provide evidence-based advice to anyone with an interest in the role of folate, vitamin B12, and other micronutrients in health (Bailey et al. 2015). We provided people with an overview of available biomarkers (serum folate and homocysteine concentrations) and the interpretation of these biomarkers across a range of clinical and population-based uses. We also explained the genetic polymorphisms involved in folate deficiency, and obtained written informed consent from participants in accordance with the instructions of the Declaration of Helsinki. The study procedures were approved by the Kagawa Nutrition University, Human Subjects and Genome Ethics Committee. There were 1101 participants (male: n = 345, female: n = 756) aged 61 ± 10 years. All participants completed a dietary intake analysis by means of a validated, brief self-administered diet history questionnaire (DHQ-L) (Sasaki 2004) containing questions about the consumption frequency of 56 foods and beverages commonly consumed by the Japanese population. Blood chemistry analysis included serum folate and tHcy concentration. The serum was isolated and stored at −80°C until analysis. Serum folate and vitamin B12 concentrations were measured at an external laboratory (SRL, Inc., Tokyo, Japan) using an Access 2

© 2017 The Authors. Congenital Anomalies published by John Wiley & Sons Australia, Ltd on behalf of Japanese Teratology Society.
chemiluminescence enzyme immunoassay (Beckman Coulter, Inc., Carlsbad, CA). Serum tHcy concentration was determined by enzyme assay using an Alfressa Auto Hcy kit (Alfressa Pharma, Inc., Osaka, Japan) (Hiraoka et al. 2014). In order to genotype a large number of blood samples rapidly and at low cost, we developed an automated genotyping machine using a bead array in capillary tubes (Kagawa et al. 2010) and used this approach for MTHFR and other risk SNPs (Kagawa et al., 2012).

One month after obtaining the blood samples, the genotype was announced by the attending medical doctor to subjects who had agreed to know their genotypes, and nutritional and exercise guidance was provided by a registered dietitian. To supply enough folate easily, we developed “folate fortified rice” containing 26.7 mg folate, 187 mg thiamin, 66.7 mg vitamin B₆, and 320 μg vitamin B₁₂ per 100 g rice in collaboration with House-wellness Foods Company. We also developed a folate fortified bread called Sakado Folate Bread (folic acid 340 ± 21 μg/100 g (215 ± 14.7 μg/slice/64.0 g bread) and succeeded in improving folate status and depressive states (Hiraoka et al. 2014). Commercially available Sakado Folate Bread contains 160 μg folic acid/slice.

Serum folate concentrations were significantly increased and serum tHcy concentrations decreased at 4 months after the onset of this intervention by the Project (Fig. 2). The horizontal lines in panel A (serum folate), B (serum homocysteine), C (dietary folate intake) and D (dietary green-yellow vegetables) indicate the standard values (7 ng/mL), (6 μg/mL), (300 μg/day) and (200 μg/day), respectively. In individuals with the TT genotype, these changes were particularly marked and intake of ‘green vegetables’, which are rich in folate, was increased in the TT genotype participants. After the instructional Project, 86% of the subjects consumed folic acid fortified food, of which 42% consumed folate-fortified rice every day.

The most important aspects of the Project were the health education of about 101 000 citizens due to the efforts of volunteers and wider consumption of folate-fortified food, especially folate-fortified rice. The above-mentioned BOND project (Bailey et al. 2015) was designed to easily explain the role of folate and vitamin B₁₂ to the general public by volunteer nutritionists. In Sakado Folate Project, we also obtained help of 125 nutrition advisors (nutrition mates) who both visited people’s neighborhoods and worked in numerous facilities called “Terakoya”. BOND project was similar in their guidance principle, but the implementation of Sakado Folate Project using nutritional advisors, “Terakoya” and other measures were quite different from those of BOND project. According to the official report prepared by Sakado City on the nutritional behavior of its citizens, after the start of this Project 80% of the population knew about the importance of folic acid, 90% tried to consume more vegetables, and 73% wanted to obtain health advice. The population of Sakado City was only about 101 000 in 2015 and there were 693 births in 2015; consequently, a decrease in neural tube defects could not be confirmed. However, the birth rate of very low birth weight infants (body weight < 1500 g) decreased from 7.8 (1998–2002) to 6.1 (2006–2010) per thousand births after the Project. The total fertility rate in the first period was 1.13 and second period was 1.20 (Ministry of Internal Affairs. Statics Bureau of Japanese Government, 1990a,b).

There are five reasons for the overall success of the Sakado Folate Project.

1. Effective organization of volunteers recommended by the government to promote a healthy diet.
2. The invention of a rapid and inexpensive genetic polymorphism analyzer.
3. The involvement of a well-trained registered dietician in the nutritional survey who was available throughout the project to give advice.
4. The cooperation of the local government and the mayor of Sakado City.
5. The development of two folate-fortified foods: Sakado Folate Rice and Sakado Folate Bread.

**MEDICAL COST SAVINGS RESULTING FROM THE SAKADO FOLATE PROJECT**

We verified the effect of personalized nutrition intervention on the folate status of participants; this, together with genotype notification, was effective in motivating individuals to change their dietary habits and improve their folate intake (Fig. 3). Thus, the subjects with the
most risky TT genotype showed the highest increase in serum folate and the largest decrease in serum homocysteine (Figs 2, 3).

Both the prevalence of disease and the medical costs of subjects participating in the Project, and of the general Sakado population, were reduced. A nutrition survey conducted by the Saitama Prefecture Department of Health in 2011 revealed that the prevalence (% of adult population) of obesity, hypertension, type 2 diabetes and dyslipidemia diagnosed by physicians was significantly lower than that in the males of three other cities (Fig. 4). The data on Figure 4 are based on adult population in 2011. The male population showed significantly lower prevalence of these diseases than the other three cities ($P = 0.04$), but female population did not reach significant level. The average ages of males and females in Sakado city in 2011 were 42.4 and 44.4, respectively, while those of three cities (Yoshikawa,
Asaka and Okegawa cities) on average were younger; 41.4 and 43.1 years old (Ministry of Internal Affairs. Statics Bureau of Japanese Government, 1990a,b).

This was reflected in the relative decrease in medical costs in Sakado City immediately after the start of the Project in 2006 (Fig. 5). The relative decrease of medical costs from 2003–2005 to 2006–2008 in Sakado City plotted in Figure 5 is statistically significant ($P = 0.0459$, using Wilcoxon rank sum test). The actual medical costs of National Health Insurance per person was ¥303,900 in 2006–2007 compared with those of ¥290,399 in 2001–2005 in Sakado City. The increment ¥13,501 was the smallest among all cities in Saitama Prefecture. The abscissa in Figure 5 is the year and the ordinate indicates the medical costs for each of the four cities being compared relative to the average total medical costs in Japan (1.00). A few years may be necessary for this Project to expect a reduction in the significant amount of the medical expenditure. However, in the large population of Sakado city, similar to United States as soon as the serum folate elevation, rapid decrease in stroke took place within a year (Yang et al. 2006), because the increase of serum folate and decrease of serum homocysteine takes place only within a few weeks (Hiraoka et al. 2004, 2014). Moreover, rapid preventive effects of folate on cardiac infarction was reported in detail (Moens et al. 2008). In addition, the subjects of the Project in 2006 and 2007 were mainly local health and nutrition leaders to teach citizens rapidly. Since there were no rapid demographic change in Sakado city (Ministry of Internal Affairs. Statics Bureau of Japanese Government, 1990a,b), the rapid reduction in the medical expense is attributed to the Project. We should show the values to the year 2014 or 2015 in Figure 5, however, unfortunately, there was a large reform of the medical insurance system during 2008, which is Comprehensive Reform of Social Security: Advanced Elderly Medical Service System (Medical care for the elderly over 75 years old) (Ministry of Health, Labor and Welfare 1990), so we have to discontinue the graph at 2008. Although the total medical cost were changed, in 2014, that of Sakado city (average age: male 42.8, female 44.7) was 95.7% of that of all cities in Saitama (average age: male 42.3, female 44.3), and 92.5% of that of neighboring Kawagoe city (average age: male 42.6, female 44.7).

The average medical costs saved per person per month (during May and October before the project from 2003 to 2005 minus those after the project from 2006 to 2008) after the Project for cardiac infarct, cerebrovascular disease, dementia and osteoporosis were ¥364–297 = –¥67, ¥420–¥409 = –¥11, ¥85–¥47 = –¥38 and ¥230–¥222 = –¥8, respectively. These data were based on the automatic computer output of Japanese Health Care Insurance System of Sakado city, and the medical costs were analyzed by experts of medical epidemiology. However, the results for average citizens shown in Figures 4 and 5 do not represent the direct effects of folate intake because we did not measure the serum folate levels of all citizens.

The relationship between the costs and the changes in serum folate level was used to estimate the medical cost savings (Bentley et al. 2009). The proportional association between the changes in serum/plasma folate and the intake of folate in the diet based on information from several studies reporting these two parameters were calculated using a linear equation that describes their association (Quinlivan and Gregory 2007): change in serum folate (mg/L) = [intake of folic acid as DFE (μg/day) $\times 0.0145$] $+ 0.132$.
(Dary 2009). The formula considers the equivalence of folic acid in terms of DFE based on the approximately 70% higher bioavailability of monoglutamyl pteridine over dietary sources of folate. Table 1 summarizes the changes in serum folate of three genotypes of MTHFR before and after the Project, with the intake of folate being expressed in terms of synthetic folate (monoglutamyl pteridine). The serum folate concentration in the 1101 subjects increased from 7.68 ng/mL to 9.92 ng/mL, a change of 2.24 ng/mL. However, the subjects notified of carrying the TT genotype (n = 191) showed serum folate concentration increased 178% from 6.87 (17.4 nmol/L) to 12.22 (22.5 nmol/L). The calculated medical costs saved per person was ¥1,669 for a total of ¥1,784,042 per year (Sakado City Local Government 2015) according to the relationship between folate intake and medical cost reported (Bentley et al. 2009), if the relationship proposed by Bentley is applicable to the people of Sakado.

However, these estimated medical cost savings were limited to coronary infarction, colon cancer and spina bifida (Bentley 2009), and therefore we measured the medical cost savings of the subjects participating in the Project (Fig. 6). To do this, we compared the average medical costs of Sakado citizens (expected costs: ¥10 276 to ¥12 196; average: ¥11 076 per person per month) with those of participants in the Project. The average medical costs of the participants fluctuated greatly, between ¥4518 to ¥13 793, (average: ¥12 196; average: ¥11 076 per person per month) with those of Sakado citizens (expected costs: ¥10 276 and therefore we measured the medical cost savings of the subjects participating in the Project. The average medical costs of the Sakado Folate Project. The abscissa shows the year and the ordinate indicates the medical costs per person per month in yen. Black line: expected costs of citizens in Sakado City, Gray line: average of actual costs of the subjects participating in the Project, Light gray line: median of the actual costs of the subjects participating in the Project.

In summary, folic acid fortification was effective in preventing not only NTDs but also other diseases, including CVD, dementia, kidney diseases and osteoporosis, irrespective of genetic polymorphisms such as the MTHFR TT genotype. The medical costs saved were dependent on the disease involved, the projects implemented in each country, and the economic evaluation methods used. The genotyping was effective in high risk approach with increased motivation (Figs 2, 3). Genome notification played an important role in the Sakado Folate Project and was effective in motivating folate intake, especially in participants with the TT genotype, yet the increase in serum folate (129%) was less than that observed following compulsory folic acid fortification in the USA (149.6%). Evaluation showed that compulsory folic acid fortification is more cost-effective than the genotype-based educational approach used in the Sakado Folate Project. In conclusion, compulsory folic acid fortification is the best method to prevent disease and reduce medical costs.

CONCLUSION

ACKNOWLEDGMENTS

This study was supported by Grants-in-Aid for Scientific Research from the Ministry of Education, Science, and Culture of Japan, JSPS

© 2017 The Authors. Congenital Anomalies published by John Wiley & Sons Australia, Ltd on behalf of Japanese Teratology Society.
DISCLOSURES

None.

REFERENCES

Agnew-Blais JC, Wassernth-Smoller S, Kang JH et al. 2015. Folate, vitamin B-6, and vitamin B-12 intake and mild cognitive impairment and probable dementia in the Women’s Health Initiative Memory Study. J Acad Nutr Diet 115 (2):231–241.

Bailey LB, Stover PJ, McNulty H et al. 2015. Biomarkers of nutrition for development-folate review. J Nutr 145 (7):1636S–1680S.

Bottiglieri T. 2005. Homocysteine and folate metabolism in depression. Prog Neuropsychopharmacol Biol Psychiatry 29:1103–1112.

Cai W, Yin L, Yang F, Zhang L, Cheng J. 2014. Association between Hcy levels and the CBS844ins68 and MTHFR C677T polymorphisms with essential hypertension. Biomed Rep 2 (6):861–868.

Chamarin I. 1987. Megaloblastic anemia, cobalamin, and folate. J Clin Pathol 40 (9):978–984.

Cui R, Iso H, Date C, Kikuchi S, Tamakoshi A, Japan Collaborative Cohort Study Group. 2010. Dietary folate and vitamin B9 and B12 intake in relation to mortality from cardiovascular diseases: Japanese collaborative cohort study. Stroke 41 (6):1285–1289.

Dary O. 2009. Nutritional interpretation of folic acid interventions. Nutr Rev 67 (4):235–244.

Fanji V, Kafai R. 2006. Trends in serum folate, RBC folate, and circulating total homocysteine concentrations in the United States: analysis of data from National Health and Nutrition Examination Surveys, 1988–1994, 1999–2000, and 2001–2002. J Nutr 136:153–158.

Gjesdal CG, Vollset SE, Ueland PM, Refsum H, Meyer HE, Tell GS. 2007. Total homocysteine concentrations in the United States: analysis of data from the NHANES III surveys 1988–1994 and 1999–2000. J Bone Miner Res 22 (5):747–756.

Grosse SD, Berry RJ, Tilford MJ, Kucik JE, Waitzman NJ. 2016. Retrospective assessment of cost savings from prevention: Folic acid fortification and spina bifida in the U.S. Am J Prev Med 50 (S1):S74–S80.

Hiraoka M, Fukushima A, Sakuma K et al. 2014. Folic acid supplementation reduced depression score in young Japanese women with depression-susceptible genetic polymorphisms. Nutr Health 1 (2):1–10.

Hiraoka M, Kagawa Y, Hiraoka M, Miyashita-Hatano Y et al. 2010. Automated single nucleotide polymorphism typing using bead array in capillary tube. J Biosci Bioeng 111:505–508.

Kagawa Y, Yanagisawa Y, Inoue M et al. 2012. Single nucleotide polymorphism of thrythlon genes for energy metabolism: evolutionary origins and prospects for intervention to prevent obesity-related diseases. Biochem Biophys Res Commun 295:207–222.

Kageyama M, Hiraoka M, Kagawa Y. 2008. Relationship between genetic polymorphism, serum folate and homocysteine in Alzheimer’s Disease. Asia Pac J Public Health 20 (Suppl):111–117.

Lindahl T. 2013. My journey to DNA repair. Genomics Proteomics Bioinformatics 11 (1):2–7.

Lopes da Silva S, Vellas B, Elefants S et al. 2014. Plasma nutrient status of patients with Alzheimer’s disease: Systematic review and meta-analysis. Alzheimers Dement 10 (4):485–502.

Massy ZA. 2003. Potential strategies to normalize the levels of homocysteine in chronic renal failure patients. Kidney Int Suppl 2003 (84):S134–S136.

Ministry of Health, Labor and Welfare. 2014. The National Health and Nutrition Survey Japan, 2014 (partly in English). www.mhlw.go.jp/bunya/kenkou/eiyou/h24-houkoku.html. Accessed in 2016.

Ministry of Health, Labor and Welfare. 1990–2014. Health and Medical Services. Health Care Insurance System. http://www.mhlw.go.jp/english/ wp/wp-hw9d/02e.pdf. Accessed in 2015.

Ministry of Internal Affairs. Statics Bureau of Japanese Government 1990a–2015. Average age of all cities and towns in Saitama Prefecture. https://www.pref.saitama.lg.jp/a0701/hokentoukei/gotoku.html. Accessed in 2016.

Ministry of Internal Affairs. Statics Bureau of Japanese Government 2000b–2015. Statistics of Child birth and fertility rates in Saitama Prefecture 1990–2014. https://www.pref.saitama.lg.jp/a0701i/hokentoukei/gotoku.html. Accessed in 2016.

Mito N, Takimoto H, Umegaki K et al. 2007. Folate intakes and folate gene polymorphism, heme methyltransferase gene 80G/A polymorphism and methotrexate efficacy or methotrexate-related toxicity in rheumatoid arthritis: A meta-analysis. Int Immunopharmacol 38:8–15.

Moens AL, Champion HC, Claeys MJ et al. 2008. High-dose folate acid pretreatment blunts cardiac dysfunction during ischemia coupled to maintenance of high-energy phosphates and reduces postreperfusion injury. Circulation 117 (14):1810–1819.

© 2017 The Authors. Congenital Anomalies published by John Wiley & Sons Australia, Ltd on behalf of Japanese Teratology Society.
Sadewa AH, Sunarti, Sutomo R et al. 2002. The C677T mutation in the methylene tetrahydrofolate reductase gene among the Indonesian Javanese population. Kobe J Med Sci 48 (5–6):137–144.
Saitama Prefecture Local Government. 2011. Health and Nutritional Survey Data 2011. https://www.pref.saitama.lg.jp/a0704/data/documents/505961.pdf. Accessed in 2012.
Sakado City Local Government. 2015. Report on the Sakado Folate Project to Saitama Prefecture.
Sasaki S. 2004. Development and evaluation of dietary assessment methods using biomarkers and diet history questionnaires for individuals. Tokyo, Japan: Ministry of Health, Labor and Welfare (in Japanese).
Van Allen MI, Fraser FC, Dallaire L et al. 1993. Recommendations on the use of folic acid supplementation to prevent the recurrence of neural tube defects. Clinical Teratology Committee, Canadian College of Medical Geneticists. CMAJ 149 (9):1239–1243.
Wang C, Riedl KM, Schwartz SJ. 2010. A liquid chromatography-tandem mass spectrometric method for quantitative determination of native 5-methylenetetrahydrofolate and its polyglutamyl derivatives in raw vegetables. J Chromatogr B Analyt Technol Biomed Life Sci 878 (29):2949–2958.
Yang Q, Botto LD, Erickson JD et al. 2006. Improvement in stroke mortality in Canada and the United States, 1990 to 2002. Circulation 113 (10):1335–1343.
Yi Y, Lindemann M, Colligs A, Snowball C. 2011. Economic burden of neural tube defects and impact of prevention with folic acid: a literature review. Eur J Pediatr 170 (11):1391–1400.
Zhi X, Yang B, Fan S et al. 2016. Gender-specific interactions of MTHFR C677T and MTRR A66G polymorphisms with overweight/obesity on serum lipid levels in a Chinese Han population. Lipids Health Dis 15 (1):185.