Type 1 Diabetes Versus Type 2 Diabetes/Metabolic Syndrome, Opposite Extremes of an Immune Spectrum Disorder Induced by Vaccines

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Abstract: There is an epidemic in children of type 2 diabetes and metabolic syndrome including individual diseases that form the components of metabolic syndrome. The epidemic resembles the epidemic of type 1 diabetes in children which has been linked to immunization. The epidemic of obesity in US children has a statistically significant positive correlation with the number of vaccine doses recommended. There is a similar trend with both hypertension and metabolic syndrome. The incidence of type 2 diabetes in Japanese children decreased significantly following the discontinuation of the BCG vaccine, a vaccine which is associated with an increased risk of type 1 diabetes. This paper describes two aberrant responses to immunization. At one extreme immunization leads to progressive autoimmune diseases including type 1 diabetes. A second response to immunization, and an opposite extreme to autoimmunity, is for the body to suppress the immune system through increased cortisol activity and other counter measures leading to type 2 diabetes and metabolic syndrome. Some vaccine recipients may have a mixed response, falling between the extremes, such as an incomplete autoimmune disorder or an intermittent autoimmune disorder. The propensity to develop a particular response relates to race. Japanese children produce large amounts of cortisol following immunization and have lower risk of type 1 diabetes but higher risk of type 2 diabetes than White children. Analysis using Austin Bradford-Hill criteria for causation support a causal relation between immunization and metabolic syndrome. Additional studies are needed to further characterize this risk.

Keywords: Vaccines, type 1 diabetes, type 2 diabetes, metabolic syndrome.

I. BACKGROUND

There is an epidemic of several diseases in human children and adults including hypertension, obesity, hyperlipidemia, low high density lipoprotein [HDL] cholesterol, microalbuminurea, and insulin resistance [1, 2]. This syndrome has been collectively classified as metabolic syndrome [3] and is closely associated with type 2 diabetes [4] and other health problems including death [5]. Many have blamed poor diet [6] and lack of exercise for the epidemics of type 2 diabetes and metabolic syndrome. Diet and exercise have been touted as the cure for metabolic syndrome but have not been very effective [7] and have not stopped the epidemic to date. The poor diet and exercise theory does not explain the obesity epidemic in children under 6 month of age who don't drink many sodas, don't eat a lot of fried potatoes and have never been very active. Recent data from a Massachusetts health maintenance organization [HMO] shows a 73% increase in overweight infants under 6 months of age from 1980 to 2001 [8].

Several investigators have proposed that metabolic syndrome is an inflammatory condition or the result of increased cortisol production, a hormone that suppresses inflammatory conditions. The epidemic of metabolic syndrome in children mirrors an epidemic of type 1 diabetes in children, which has been linked to a class of immune stimulants, vaccines [9-12]. A proposed mechanism of vaccine induced metabolic syndrome is presented. Epidemiological evidence supporting an association between immunization and metabolic syndrome is presented. Epidemiological and experimental data are reviewed supporting a racial basis for determining whether an individual is more likely to develop an autoimmune disease like type 1 diabetes or metabolic disease as a complication of immunization.

II. PROPOSED MECHANISM OF VACCINE INDUCED METABOLIC SYNDROME

A. Autoimmunity and Metabolic Syndrome are Opposing Ends of an Immune Spectrum Disorder

The proposed mechanism of immunization induced metabolic syndrome is by an intrinsic neuroendocrine feedback loop to suppress an immune system chronically activated by immunization. It is well accepted that in all organ systems there are homeostatic mechanisms to regulate their activity. Autoimmune diseases are conditions that result from an over active immune system. Likewise one would expect that there would be one or more diseases that arise when the body attempts to suppress what it interprets as an over active immune system. Cortisol is a hormone that suppresses the immune system and can prevent autoimmunity. Hypersecretion of cortisol however can lead to Cushingoid Syndrome which closely resembles metabolic syndrome. Vaccines have been shown to cause autoimmune diseases including type 1 diabetes [9-12] and other chronic inflamma-
B. Inflammation as the Cause of Metabolic Syndrome

It has been proposed that metabolic syndrome is an inflammatory condition [22]. This belief is supported by studies that have shown that inflammation preceeds metabolic syndrome [23, 24]. A study on Finnish middle aged men [23] found men with elevated C-reactive protein (CRP) concentrations had higher age-adjusted risk of developing metabolic syndrome. A study of men and women in Mexico [24] found women with elevated CRP in the highest tertile had an increased relative risk of developing metabolic syndrome.

Some have suggested that metabolic syndrome causes inflammation [25]. While metabolic syndrome may cause inflammation it is more likely that inflammation initially causes metabolic syndrome. Inflammation has been associated with the development of components of metabolic syndrome, independent of the presence of characteristics of metabolic syndrome [23, 24]. Glucose intolerance/type 2 diabetes [26] and hypertension [27, 28] are both independently associated with inflammation. There is additional evidence that once metabolic syndrome begins it causes more inflammation [25] which in turn makes the disease worse. Adipocytes and the accompanying macrophages appear to make inflammatory mediators. While there is an association between obesity and inflammation [29], obesity can exist without inflammation as demonstrated by obese individuals with a healthy metabolic profile [30]. The later evidence supports the view that in many inflammation precedes the development of metabolic syndrome.

Metabolic syndrome has a remarkable similarity to mild Cushingoid Syndrome [31, 32] and several have suggested that metabolic syndrome is caused by increased cortisol activity [33-35]. There is data showing increased cortisol levels associated with metabolic syndrome [36]. There is evidence that increased peripheral activation of cortisol secondary to increased enzymatic activity of 11-beta hydroxysteroid dehydrogenase type 1 contributes to the development of metabolic syndrome [37, 38]. Some have also suggested that metabolic syndrome is in part due to increased cellular uptake of cortisol [39]. Excessive amounts of exogenous glucocorticoids are known to cause hypertension, obesity, hyperlipidemia, and glucose intolerance. The effect is dose dependent. There are many similarities between excessive cortisol activity and metabolic syndrome. Excessive cortisol activity is associated with metabolic disturbances including increased glucose levels, obesity and hyperlipidemia [40] just like in metabolic syndrome. Excessive cortisol activity is also associated with increased cardiovascular events [41, 42] just like metabolic syndrome.

There is biological evidence that specific lymphokines released during inflammation can cause the release of cortisol and cause the biological changes that occur in metabolic syndrome. It has been hypothesized [43] that the metabolic syndrome like responses to lymphokines provide a short term survival advantage helping the host survive noxious events. Hyperlipidemia for example may help the body clear fat soluble toxins. A problem arises in certain individuals when inflammation becomes chronic and the changes lead to metabolic syndrome. In these cases changes that are a survival advantage acutely are chronically an hazard.

C. Vaccines as an Inducer of Metabolic Syndrome

Vaccines have been shown to stimulate the immune system in the short term causing the release of cytokines that can increase cortisol activity. The acellular diphtheria tetanus pertussis vaccine has been reported to cause the release of IL-6 [44]. The Diphtheria-Tetanus-Polio-Typhim vaccine stimulated IL-6 production [45]. The Diphtheria-Tetanus-whole cell Pertussis but not the Diphtheria-Tetanus-acellular Pertussis vaccine elicited increased IL-6 at 2 days post immunization [46]. The influenza vaccine stimulated release of IL-6 and IL-10 [47]. The influenza and pneumococcal vaccine caused rises in CRP [48]. Researchers in France have linked aluminum adjuvants in vaccines to an inflammatory condition called myofascitis [13, 49]. Several papers have shown that immunization of children can increase cortisol levels at least in the short term [14-21].

Cytokine production, particularly IL-6, increases with age [50-52] and this can explain the increase in metabolic syndrome with age. Both IL-1 [53, 54] and IL-6 [55-57] enhance cortisol release and thus have the potential to cause metabolic syndrome. IL-6 has been associated with the development of metabolic syndrome [58, 59]. In addition IL-6 has been directly associated with the development of diabetes [26], insulin resistance [60] and altered lipid levels [61-63].

D. Resetting the Hypothalamus

A second mechanism by which immunization may lead to the induction of metabolic syndrome is through the resetting of the hypothalamus. Immunization in the first year of life may affect the onset of metabolic syndrome by resetting of the hypothalamus, creating more cortisol release. Two well characterized examples support this hypothesis. It has been shown that the hypothalamus is reset in children who undergo stress in utero. These children produce higher cortisol release and hence have increased symptoms resembling metabolic syndrome [64-66]. A second example of resetting the hypothalamus is in children which are born with low birth weight and are at increased risk of developing metabolic syndrome [67].

III. EPIDEMIOLOGY OF VACCINE INDUCED TYPE 2 DIABETES AND METABOLIC SYNDROME

Epidemiological data support the proposed mechanism of vaccine induced type 2 diabetes and metabolic syndrome.

A. Epidemic of Type 2 Diabetes and Metabolic Syndrome Resembles the Epidemic of Type 1 Diabetes

One line of support for the proposed mechanism of vaccine induced metabolic syndrome is that the epidemic of metabolic syndrome and type 2 diabetes resembles the epidemic of type 1 diabetes. The role of vaccines in causing the epidemic of type 1 diabetes is supported by data from a prospective clinical trial, animal toxicity data as well as epidemiological data [9-12]. There is an epidemic of metabolic syndrome and its components in children living in the US [1, 2] and other countries including the UK and Australia [68,
Data on the prevalence of metabolic syndrome, obesity, and hypertension in US children has been published covering a period of at least 10 years. Obesity in US children aged 4 to 12 years old increased on average of 3.23-5.85% per year, depending on race, from 1986 to 1998 in the National Longitudinal Survey of Youths [70]. The prevalence of overweight children increased on average of 1.41% to 3.60% per year, depending on race, in the same age group. Similar rises were seen in children age 0-19 years old in the NHANES study [71, 72]. Between 1988 and 2000 the prevalence of obesity rose 4.4% per year in children age 12-19 years old, 3.4% year in children age 6-11 years old, and 4.2% per year in children age 2-5 years old. Blood pressure also rose in children and adolescents between 1988 to 2000 [73] [p<0.001]. Metabolic syndrome increased on average of 4% a year in adolescents aged 12-19 years old according to US NHANES data from 1988-1992 to 1999-2000 [2] [p<0.001]. In comparison Type 1 diabetes in children increased 2.3% per year in the US from 1978-2004 [74].

The findings do not appear to be limited to the US. Data from Finnish children [75] shows the prevalence of obesity in children age 12-18 in the years 1977 to 1999 increased from 1.1% to 2.7% in boys [relative risk 2.45] and from 0.4

Table 1. Correlation Between Vaccine Doses and Obesity, Metabolic Syndrome and Diastolic Blood Pressure

<table>
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<th>YEAR</th>
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P<0.05

The number of doses of vaccines recommended for children in the US increased from 24 to 45 in the years 1978 to 2006. The rise in the number of doses is statistically associated with the increase in obesity in US children age 2-19. A similar but not statistically significant association is also seen in the US with metabolic syndrome and diastolic blood pressure in children.
to 1.4% in girls [relative risk 3.5]; on average a 5% rise per year. The prevalence of overweight increased from 7.2% to 16.7% in boys and 4.0% to 9.8% in girls, on average a 4% rise per year. During this time the combined measles mumps rubella and hemophilus vaccines were added to the immunization schedule as well as a more potent pertussis vaccine [10]. This increase in obesity is very similar to the 3.4% per year increase in type 1 diabetes from 1965 to 1996 [76] which has been attributed to an increase in number of vaccines [10, 11].

B. Positive Correlation Between the Number of Doses of Vaccine Administered and the Prevalence of Obesity

There is an association between the number of vaccine doses recommended for US children and rises in obesity and metabolic syndrome. The number of pediatric vaccines universally recommended in the US by the Centers for Disease Control and Prevention [CDC] has increased from seven to thirteen and the number of recommended doses from 24 to 45. There was a statistically significant correlation between the number of doses of vaccine given and obesity in children age 2-5, 6-11 and 12-19 (Table 1). Statistics were performed using the program Statistica, Stat Soft, 1993. The Pearson Product-Moment Correlation module was used to determine the statistical association between number of vaccines doses recommended with obesity. Data on US rates of metabolic syndrome and hypertension in children exists for only two periods of time, two data points, so statistical correlations can not be calculated. However there was a similar trend with both of these outcomes and both outcomes showed a statistically significant rises during the study period. There was a similar trend in Finland with a positive correlation between the number of vaccine doses given and an increased prevalence of obesity, just like in the US [10, 11].

C. Decline in Type 2 Diabetes Occurred Following Discontinuation of the Tuberculosis Vaccine [BCG]

The rates of Type 2 diabetes and metabolic syndrome is increasing throughout the world [77]. However, there was a statistically significant drop in the incidence of Type 2 diabetes in Tokyo Japan in elementary and junior high school students [78, 79]. The drop occurred in 2003 after the discontinuation of routine BCG vaccination of elementary and junior high school students [78, 79]. The drop occurred in 2003 after the discontinuation of routine BCG vaccination of elementary and junior high school students in Japan [80]. BCG vaccination has previously been associated with an increased risk of type 1 diabetes [9].

D. Racial Predisposition for Development of Type 1 or Type 2 Diabetes and the Association with Cortisol Secretion Following Immunization

The proposed mechanism that some respond to immunization by developing autoimmune disease while others respond to immunization by developing metabolic syndrome is supported by racial differences in the rates of type 1 and type 2 diabetes and cortisol secretion following immunization. Type 1 diabetes is an autoimmune disease while type 2 diabetes is related to insulin resistance, a component of metabolic syndrome. Type 2 diabetes is associated with obesity, another component of metabolic syndrome, while type 1 diabetes is not. White children including adolescents tend to have an high absolute incidence of type 1 diabetes and a high ratios of the incidence of type 1/type 2 diabetes compared children of Japanese or Chinese decent. The later children have higher incidences of type 2 diabetes and, consequently, a lower ratio of the incidence of type 1 to type 2 diabetes [78, 81-83].

The racial differences in the incidence of type 1 and type 2 diabetes can be explained by cortisol release following immunization. Japanese children have a higher cortisol responses to immunization than White children [84]. The differences in cortisol responses of Whites and Japanese appears to extend into adulthood as well [85]. Because they release more cortisol following immunization, Japanese children would be expected to have a lower rate of autoimmune diseases including type 1 diabetes as well as a lower ratio of type 1/type 2 diabetes compared to Caucasians. Elevated cortisol secretion can cause metabolic syndrome, as described above, but cortisol is an immune suppressant and prevents autoimmune disease. Decreases in cortisol following adrenalectomy leads to increased rates of type 1 diabetes in mice [86] and experimental autoimmune diseases [87, 88].

IV. AUSTIN BRADFORD-HILL CRITERIA FOR Establishing Causation

Austin Bradford-Hill’s criteria [89] are used to show whether an association is likely to be an causative relationship. He listed nine factors and all nine are met in the association between immunization and metabolic syndrome (Table 2). Austin Bradford-Hill analysis in part relies on bioplausibility. Based on the existing knowledge it is proposed that vaccines induce inflammation which increases cortisol or cortisol like activity leading to a Cushingoid like state which is metabolic syndrome.

V. CLINICAL IMPLICATIONS FOR Reversibility of the Metabolic Syndrome

Current studies are being performed to try to reverse or prevent metabolic syndrome by decreasing cortisol activity. One approach is to inhibit the enzyme 11-beta hydroxysteroid dehydrogenase type 1, an enzyme which increases the activity of cortisol in the peripheral tissue. Evidence from animal models of autoimmune disease suggests that suppression of cortisol, while likely to suppress metabolic syndrome, will increase the risk of autoimmune diseases. Decreasing cortisol production in rodents by adrenalectomy, for example, greatly increases the risk of diabetes in [86]. A safe and effective method of suppressing metabolic syndrome will require the reduction of inflammation, the precursor to both autoimmune disease and metabolic syndrome. Ideally one will want to reduce exposure to vaccines that lead to chronic inflammation. Once exposure to inflammatory mediators have been initiated, one can attempt to suppress the inflammation. There is evidence that certain anti-inflammatory agents will prevent development of metabolic syndrome. An anti-inflammatory drug AGI-1067 in its Phase III study called ARISE, showed a 64 percent reduction in patients developing diabetes in the group reviving AGI-1067 as compared to prospective controls.

VI. FUTURE STUDIES

Vaccines continue to be approved based on small studies with only short term follow up. Theses studies are inade-
extreme of the disorder is the development of progressive autoimmune diseases such as type 1 diabetes. A second response to immunization, and an opposite extreme to autoimmunity, is for the body to suppress the immune system through increased cortisol activity and other counter measures leading to type 2 diabetes and metabolic syndrome. Some vaccine recipients may have a mixed response, falling between the extremes, such as an incomplete autoimmune disorder or an intermittent autoimmune disorder. The propensity to develop a particular response relates to race. The propensity to develop a particular response with racial differences. Japanese children produce larger amounts of cortisol following immunization and have a lower risk of type 1 diabetes but higher risk of type 2 diabetes than white children. Additional studies are needed to further characterize this risk.

**ACKNOWLEDGEMENT**

The author is president and share holder of Classen Immunotherapies. Classen Immunotherapies holds many patents and patent applications related to testing vaccines for their risk of chronic diseases including type 1 and type 2 diabetes.

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