Appetite control with relevance to immunometabolism has become critical to the treatment of non-alcoholic fatty liver disease (NAFLD), cardiovascular disease and diabetes [1-4]. The major defect in global chronic disease is autoimmune disease with defective adipose tissue and liver interaction involved with the release of inflammatory cytokines and adipocytokines [5] relevant to toxic immune reactions that involve the pancreas brain, heart, thyroid, kidneys and reproductive organs. Appetite control and autoimmune disease are connected with the identification of Sirtuin 1 (Sirt 1) as the anti-aging gene [1-3] involved in appetite regulation and the prevention of autoimmune disease [4].

Sirt 1 repression is responsible for mitophagy [4] in various cells and connected to autoimmune disease and irreversible programmed cell death in various cells and tissues. Science and medicine and its relevance to genomic medicine now need to consider Sirt 1 gene expression and its plasma Sirt 1 levels with relevance to accelerated immune reactions that trigger acute cardiovascular disease [6]. Sirt 1 is a nicotinamide adenine dinucleotide (NAD +) dependent class III histone deacetylase (HDAC) that targets transcription factors to adapt gene expression to metabolic activity and the deacetylation of nuclear receptors indicate its critical involvement in insulin resistance. In situ hybridization analysis has localized the human Sirt 1 gene to chromosome 10q21.3 [7].

The Sirt 1 gene has now been linked to autoimmune disease and various diseases with deletions, mutations, inversions and aberrations in chromosome 10q21.3 [8-10]. Sirt 1 has a molecular weight (Mol Wt) of 81 kda with Mol Wt variation (81-110 kda) between laboratories. Sirt 1 is now identified as the heat shock gene [11,12] and involved in the deacetylation of heat shock factor 1 (HSF 1) and regulation of heat shock proteins (HSP) and nitric oxide metabolism connected to natural killer cell activity, mitophagy and autoimmune disease connected to chronic disease [13].

Sirt 1 is an acute phase protein involved with neuron proliferation [14] and its regulation of the suprachiasmatic nucleus is involved with control of the circadian rhythm [3]. The circadian rhythm and immune system are closely connected to the immune response [15-17]. Sirt 1 may now be considered to be involved with the circadian rhythm of the immune system and critical to the immune response in various communities. Sirt 1 is important to regulation of HSF1 and T and B cell immune response [18-20] and is involved with B cell antibody regulation and T cell tolerance [21,22]. Sirt 1 is involved in regulation of nuclear receptors peroxisome proliferator-activated receptor (PPAR) and liver X receptor (LXR) with relevance to metabolic homeostasis [3] and may now be connected to the regulation of these nuclear receptors (PPAR/ LXR) in inflammation and immunity [23]. Sirt 1 is now important to the immune mediated control of synaptic plasticity [24,25] and anti-epileptic drugs relevant to the immune system and epileptic stroke [26,27].
diagnostic protein that is critical for mitochondrial apoptosis with early interpretation of global health and chronic disease.

The progression of global chronic disease may now involve the measurement of plasma Sirt 1 for primary autoimmune disease with the critical inter-relationship between immunology tests and other diagnostic tests that involve biochemistry, microbiology, and hematology. Immunization, vaccinations and vaccine programs for various diseases may require specific measurements of plasma Sirt 1 levels. Sirt 1 analysis on blood, plasma and sera with other acute phase proteins (Figure 1) may be important to other immunological tests and collection of plasma/sera sensitive to Sirt 1 stability and associated with appropriate preservative, freeze/thaw conditions and long term storage conditions.

**Figure 1:** Measurements of specific plasma proteins and gene expression assays are required for assessment of mitophagy with connections to various immunological diseases. Analysis of Sirt 1 in the plasma and nucleus will allow early detection of autoimmunity disease as primary induction factor in global chronic disease. Lipidomic assays for immunogenic lipids may be required to determine programmed cell death. Heat shock protein plasma analysis may provide information with relevance to natural killer cell activation in chronic disease. NAFLD and autoimmunity disease are now connected with impaired hepatic drug metabolism associated with drug induced autoimmunity. Bacterial lipopolysaccharides, magnesium and zinc levels should be assessed and relevant to magnesium deficiency induced autoimmune disease.

Bacterial lipopolysaccharides and HSP 60, 70 and 90 need to assayed on samples to avoid misinterpretations with relevance to Sirt 1 repression and mitophagy [4,5,8] in immunological associated chronic diseases. Gene expression assays of nuclear Sirt 1 for assessment of autoimmunity disease may be required [5] and associated with measurements of plasma Sirt 1 levels in various immunological diseases. Sirt 1 is connected to hepatic drug metabolism [28] and Sirt 1 repression may require Sirt 1 plasma analysis to determine drug induced immune hypersensitivity reactions (Figure 1).

Appetite control and nutritional regulation of Sirt 1 has become important to immunotherapy and the clinical treatment of global chronic diseases. Nutritional diets that contain Sirt 1 activators [25,27] have become vital to immunotherapy research to maintain immunometabolism and prevent mitophagy (Figure 1). Lipidomic analysis have identified immunogenic lipids such as ceramide and sphingosine 1 phosphate that induce apoptosis and programmed cell death [29-31]. Appetite control and nutritional intake is essential to prevent the generation of immunogenic lipids, maintain the heat shock gene Sirt 1 [32] with relevance to the generation of immunogenic heat shock proteins (HSP), activation of natural killer cells [5] and nitric oxide induced immunological alterations in humans [13].

Sirt 1 as an immunosenescence gene [13,32] is connected to the senescence of various immune cells and inflammatory responses in aging [33-35]. Appetite and diets that regulate plasma Sirt 1 [36] require plasma analysis of other Sirt 1 regulated protein hormones that are involved in the immune response and glucose homeostasis (Figure 2), such as adiponectin, fibroblast growth factor 21, brain derived neurotrophic factor, neuropeptide Y and apelin [3,37]. Insulin therapy is involved in regulation of the immune system [38,39] but Indian spice therapy (doses) for activation of immune system [40,41] should be carefully assessed with relevance to glucose therapy. Excessive Indian spice intake may interfere with insulin therapy [42] by increased cellular glucose uptake that may induce hyperglycemic mitochondrial apoptosis [42].

**Conclusion**

Human survival and the immune system are now relevant to the global chronic disease epidemic that involves NAFLD, cardiovascular disease and diabetes. Chronic disease assessment require measurement of plasma Sirt 1 levels with relevance to autoimmunity disease and mitophagy. Biotherapy that involves immunotherapy may stabilize mitophagy that is connected to autoimmunity disease and the early induction of global chronic diseases. Appetite control is essential to maintain Sirt 1 levels and to prevent the generation of immunogenic lipids and proteins that induce programmed cell death. Autoimmune disease and immunometabolism defects are now connected to the primary induction of mitophagy with the acceleration of chronic diseases in the developed and developing world.

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References


