ER stress as a trigger for beta-cell dysfunction and autoimmunity in type 1 diabetes

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Et al.
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Type 1 diabetes is an autoimmune disease characterized by the destruction of pancreatic β-cells and an absolute deficiency of insulin. Patients with type 1 diabetes are insulin dependent for life and require multiple daily insulin injections or the use of an insulin pump. It has been considered that β-cell dysfunction and death in type 1 diabetes results from a combination of inflammation, autoimmunity, β-cell stress, and insulin resistance (1-5). Clinical and experimental evidence has indicated that defects in β-cell function precede the massive death of β-cells by severe infiltration of T cells into the islets and the clinical onset of type 1 diabetes (6-9). However, the mechanisms involved in β-cell dysfunction before the onset of clinical type 1 diabetes are unclear. In this issue of Diabetes, Tersey et al. (10) add a new dimension to the progression of type 1 diabetes by demonstrating that endoplasmic reticulum (ER) stress in β-cells precedes the clinical onset of type 1 diabetes.

ER STRESS AS A TRIGGER FOR β-CELL DYSFUNCTION IN TYPE 1 DIABETES

The ER performs a number of important cellular tasks, including protein folding, calcium regulation, redox regulation, and life or death decisions (11,12). Within the β-cell, insulin production and secretion depend on the processing capacity of the ER network. Thus, the ER of the β-cell must maintain homeostatic balance in order to efficiently produce insulin and maintain viability. Perturbations to the ER by genetic or environmental factors can disrupt ER homeostasis and lead to the development of ER stress. Unresolved ER stress has deleterious effects on pancreatic β-cell function and survival, and it can influence the progression of diabetes (11,13). Using NOD mice, a well-established model of type 1 diabetes, Tersey et al. (10) demonstrated activation of ER stress pathways in β-cells prior to the onset of type 1 diabetes. Isolated islets from prediabetic NOD mice displayed age-dependent increases in expression of ER stress markers, morphologic alteration in ER structure by electron microscopy, and activation of the nuclear factor (NF)-κB pathway, which is known to be linked to ER stress. Tersey et al. also showed that MIN6 β-cells treated with a mixture of proinflammatory cytokines, a condition that mimics the immunological microenvironment of type 1 diabetes, displayed evidence of polyribosomal runoff, a finding consistent with ER stress-mediated blockage of translational initiation. Recent clinical and genetic evidence indicates that acquired or inherited ER dysfunction can lead to β-cell death in Wolfram syndrome, a rare genetic form of diabetes, as well as β-cell death in both type 1 and type 2 diabetes. While the role of ER stress in the pathogenesis of type 2 diabetes is well established (14,15), the involvement of this pathway in type 1 diabetes remained elusive. The results of Tersey et al. clearly indicate that ER stress is an important pathogenic component of β-cell dysfunction before the onset of type 1 diabetes.

TARGETING ER STRESS FOR PREVENTING β-CELL DYSFUNCTION AND AUTOIMMUNITY IN TYPE 1 DIABETES

As always, new findings raise new questions. The most intriguing question might be, what are the consequences of ER stress-mediated β-cell dysfunction in type 1 diabetes? The authors (10) provide compelling evidence that the NF-κB pathway is activated in ER-stressed β-cells prior to the onset of type 1 diabetes. NF-κB has been shown to play an important role in β-cell death during progression of type 1 diabetes. Thus, ER-stressed β-cells in prediabetic NOD mice could eventually die through NF-κB signaling and lead to frank diabetes. Another interesting possibility is that ER stress in β-cells may trigger autoimmunity and the severe infiltration of T cells into the islets (Fig. 1). Type 1 diabetes is an autoimmune disease that evolves over years in genetically susceptible individuals who are exposed to unknown environmental triggers. Importantly, all major β-cell autoantigens, including insulin, GAD65, IA-2, ZnT-8, and chromogranin A, traffic through the ER (16). ER dysfunction may cause aberrant changes in the folding and posttranslational modifications of these proteins, leading to the production of neo-self antigens. Neo-self antigens produced from ER-stressed β-cells may trigger autoimmunity in individuals with a susceptible genetic background. In either case, now it is possible to determine subjects with high risk for type 1 diabetes with greater precision using biomarkers for ER-stressed β-cells. The fact that ER function is altered in type 1 diabetes suggests the possibility of using secreted molecules from ER-stressed β-cells as early biomarkers of type 1 diabetes progression. The data by Tersey et al. (10) also suggest that chemical or biological compounds that maintain ER homeostasis could be used for therapeutic purposes. To date, there are no effective therapies targeting the ER for preventing type 1 diabetes. A novel therapeutic strategy that aims to target the common molecular processes that are altered in ER-stressed β-cells may therefore offer promise. Because preventing type 1 diabetes before β-cell function decreases below critical levels may prove to be less challenging than curing established type 1 diabetes (17), further studies on this topic are particularly
important for the development of novel prevention, diagnostic, and therapeutic strategies for type 1 diabetes.

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