Beta Cell Regeneration: A Novel Strategy for Treating Type 1 Diabetes
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Type 1 diabetes (T1D), which is typically caused by the autoimmune destruction of insulin-producing pancreatic islets, is affecting more than 1 million people in North America according to the National Diabetes Fact Sheet 2011. The key of treating T1D is to replenish the lost beta cells or their products, insulin. Insulin is commonly used for treating T1D. The occasional hypoglycemia post injection and the complications associated with long-term insulin administration may pose potential risks during a life-long insulin administration. Islet transplantation, as an experimental treatment for T1D, has the potential to maintain insulin-independence for a longer duration, but has failed to maintained normoglycemia for more than five years as per many studies [1,2]. During the last two decades, much effort has been made into exploring beta cell regeneration so that the new insulin-producing cells can replenish the lost beta cells caused by autoimmune destruction. Taking advantage of the progresses in protein sciences and regenerative medicine, beta cell regeneration may be promising for treating type 1 diabetes in future.

It is commonly accepted that the total number of islets remains constant in the life time while the size of islet increases with age. Moreover, pancreatic beta cells which have longer life-span do not undergo proliferation frequently [3]. However, many preclinical studies revealed that given proper stimulation some beta cells may regain the potential to proliferate. Proliferation of beta cells is a dynamic process of which the intrinsic pathways have not been thoroughly understood. For example, though pancreatic beta cells which originated from the same ancestor showed no differences in gene expression profile. It was recently revealed that only a small group of specialized beta cells will regain the potential to proliferate under certain conditions [4]. On the other hand, the majority of islets is replication-refractory beta cells which do not react to mitogenic stimulation [5].

Two categories of transcriptional factors or hormones have been used to promote beta cell regeneration. The first category includes factors deeply involved in the development and differentiation of pancreatic beta cells. These factors include cyclin D1/D2, cyclin-dependent kinase 4 (CDK4), glucagon-like peptide-1 (GLP-1), Neurogenin 3 (Ngn3), pancreatic duodenal homeobox-1 (Pdx1) etc [6-9]. However, the molecular pathways bridging these factors were not thoroughly understood. Several factors such as CDK4, GLP-1 and Ngn3 were demonstrated to be effective to increase the mass of pancreatic beta cells in preclinical studies [9,10]. The second category includes other mitogenic factors such as vascular endothelial growth factor (VEGF) and hepatocyte growth factor (HGF). Although these factors were also proven to be helpful in preserving the beta cell mass in T1D animals [11,12], they were absent from the intrinsic pathways governing the development and proliferation of pancreatic beta cells. Therefore, instead of working specifically on pancreatic beta cells, these factors seemed to work predominantly through combination with the native mechanism of beta cell regeneration in a synergistic or additive manner [12,13].

Despite the progress in preclinical studies, two major obstacles need to be overcome to make a real clinical advance. The lack of beta cell specificity and the limited effect of these factors currently known to impact beta cell replication. Taken together, marginal effect of beta cell regeneration is expected if any of these factors are injected intravenously.

Luckily, the recent discovery made by Douglas Melton's group of Harvard University may shed some lights into this area [14]. This group developed a novel insulin resistance mouse model to induce pathologic beta cell replication which typically happened in an early stage of type 2 diabetes. Then, they identified betatrophin, a hormone produced by liver and fat, worked specifically on pancreatic beta cells to promote replication. Most importantly, they demonstrated that the injection of betatrophin expression plasmids via tail vein led to a 17-fold increase in beta cell proliferation in just 8 days compared to the control.

Unlike previous factors, betatrophin showed both specificity and efficacy. Although the mechanisms underlying betatrophin induced beta cell proliferation is unclear, this study opens a new door to eliminate traditional insulin injection. It was reported that Evotec and Johnson & Johnson who have the rights to the molecule planned to turn it into a preclinical type 2 diabetes candidate this year.

Stem cell-based regenerative medicine is another term frequently associated with the beta cell regeneration. Stem cells with proliferation capacity and transdifferentiation potential have been explored in depth to repair the bone defect and heart infarct [15,16]. It was lately discovered that the same mechanism of stem cells, especially bone marrow mesenchymal stem cells can be used to reverse hyperglycemia in T1D animals and promoted the beta cell regeneration [17]. The phenomenon can be explained by two distinct hypotheses: 1) stem cells produce soluble factors to promote the proliferation of preexisting beta cells and/or 2) stem cells can transdifferentiate to replenish the lost beta cells.

The first hypothesis is widely accepted since the gene expression profile of mesenchymal stem cells was revealed [18]. Mesenchymal stem cells produce two types of soluble factors with distinct functions. The immunosuppressive factors such as Prostaglandin E (PGE)-2 and Indoleamine 2, 3-dioxygenase (IDO) prevented the autoimmune destruction of beta cells by reversing the activation of hyper-reactive T cells, while the trophic factors such as VEGF and HGF created an amiable environment for beta cell proliferation [19].

However, for the second hypothesis, it is still unclear whether stem cells can directly transdifferentiate to replenish lost beta cells in vivo. Although several groups have successfully transdifferentiated embryonic stem cells, adult stem cells and induced pluripotent stem cells into insulin-producing cells or even insulin-producing islet-like clusters in vitro, lineage tracing study provided limited support for this hypothesis in vivo [20-22]. Moreover, there are more practical challenges.

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of pushing cell products through the pipeline, such as the lack of standard operational procedures among research groups, the possibility of bacterial and viral contamination, the risk of immunogenicity and tumorigenicity, and the significantly high cost of massive production.

To summarize, expanded knowledge about mechanisms of beta cell proliferation will create new methods to treat T1D. Betatrophin and stem cell therapy seems to be the two possible candidates on the way. For betatrophin, a star molecule to work specifically and potently on pancreatic beta cells, although its mechanism of action is unclear and more preclinical evidences are needed, it is leading the way to the first clinical trials in beta cell regeneration. On the other hand, stem cell therapy which is still under preclinical studies has faced more practical challenges. However, stem cell therapy has demonstrated its potential in many reports as a potent immunosuppressant and trophic mediator, which may provide useful insight to understand the onset of T1D and may prove as valuable assistant in beta cell regeneration.

References