

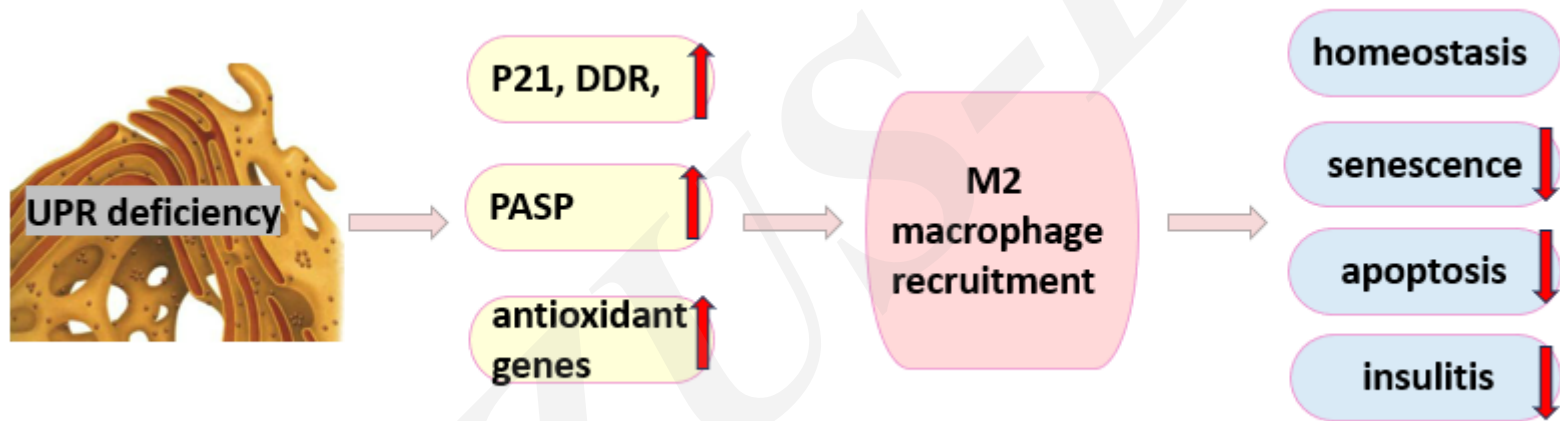
Cite this as: Haipeng CHENG, Zhenwang ZHAO, Dan LIU, Yufei WANG, Min ZHANG. Early senescence of pancreatic β cells induced by unfolded protein response deficiency prevents type 1 diabetes[J]. Journal of Zhejiang University Science B, 2024, 25(9): 796-799.
<http://doi.org/10.1631/jzus.B2400013>

Early senescence of pancreatic β cells induced by unfolded protein response deficiency prevents type 1 diabetes

Key words: Type 1 diabetes (T1D), Senescence, Unfolded protein response (UPR), Pancreatic β , cells

Research Summary

This correspondence revealed an association between pancreatic β -cell UPR deficiency and cellular senescence, providing a novel preventive strategy for T1D.



- UPR deficiency in NOD β cells triggers early senescence.
- p21-mediated secretome induces M2 macrophage recruitment to the islets.
- M2 macrophage promotes immune surveillance, eliminates terminal senesced β cells and reduces β cells apoptosis.

Innovation points

- **Introduction** of various β -cell stresses, dysfunction, and death contributing to T1D pathogenesis.
- **Summary** of the most updated research progress about the role of β -cell ER stress in the development of T1D.
- **Emphasis** of the newly established association between β -cell UPR deficiency and cellular senescence.