

Publications

Li H^{*}, Zhang L^{*}, Li J^{*}, Wu Q^{*}, Qian L, He J, Ni Y, Kovatcheva-Datchary P, Yuan R, Liu S, Shen L, Zhang M, Sheng B, Li P, Kang K, Wu L, Fang Q, Long X, Liu X, Wang X, Li Y, Ye Y, Ye J, Bao Y, Zhao Y, Xu G, Panagiotou G[#], Xu A[#], and Jia W[#] (2024) Resistant starch intake facilitates weight loss in humans by reshaping the gut microbiota. *Nat Metab* 6(3), 578-597.

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