

Rapamycin retards epigenetic ageing of keratinocytes

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Cessation, or even retardation of the ageing process is an appealing notion that has captured the imagination of humans for millennia. Even if it were possible to rejuvenate our bodies or retard the ageing process, how do we measure this?

Age, as we know it, is clearly an inappropriate measurement because it is



based purely on the passing of time, irrespective of biological changes in our body. A promising solution to this challenge is now in sight. In 2013, Steve Horvath (UCLA) described the remarkable mathematical precision by which chemical modifications on DNA called methylation, changes with age. With these methylation profiles, he derived a highly accurate age-predicting algorithm, which is essentially a timeindependent yard-stick to measure age based on biology.

Teaming up with Ken Raj of Public Health England (UK) in 2018, Horvath generated an improved algorithm called the Skin and Blood Clock that is applicable to cultured cells as well as those in the body. By using this epigenetic clock, Raj and Horvath now demonstrate that <u>rapamycin</u>'s retardation of ageing, which has been documented in numerous animal species, also extends to <u>human cells</u>.

While admittedly this is on human cells rather than the body, this finding is nevertheless consistent with the independent observation that a variant of the MLST8 gene, which encodes part of the target of rapamycin, is linked with increased rate of epigenetic ageing in humans. The harmonious convergence between observations from the bench using cultured cells and <u>genome-wide association studies</u> with <u>human tissues</u>, highlight the fidelity of this system, which has been derived and validated in humans, to test the effects of compounds on human ageing. Equally significant is the fact that Rapamycin has been well-documented to prevent arrested cells from becoming senescent—a process known as gero-conversion, which Raj and Horvath have previously demonstrated to be distinct from epigenetic ageing.







Cells from Donors A, B and C were cultured in the continued presence of the indicated compounds. Population doubling at every cell passage was ascertained until replicative senescence, and plotted against time. Credit: Horvath *et al. Aging* (2019). DOI: 10.18632/aging.101976

The Raj research team concluded, " that the life-extending property of rapamycin may be a resultant of its multiple actions which include, but not necessarily limited to suppression of cellular senescence and epigenetic aging, with the possibility of augmentation of cellular proliferative potential."

Hence it would appear that rapamycin possesses the remarkable ability to inhibit not one, but two-independent pathways of ageing. While a twofor-one offer is common in retail, it is exceptionally rare in biology. Just as in retail, offers can come with a catch, and here it is important to note that this effect of rapamycin has yet to be widely tested on other <u>cells</u> types. This caveat notwithstanding, this exciting finding and the available tools and methods described by Raj and Horvath will no doubt encourage the search for even better compounds that may help in promoting healthy human ageing.

More information: Steve Horvath et al, Rapamycin retards epigenetic ageing of keratinocytes independently of its effects on replicative senescence, proliferation and differentiation, *Aging* (2019). <u>DOI:</u> 10.18632/aging.101976

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