

Aging with VitroAge

Traditional keratinocyte cell culture media actively resist the aging process with a variety of anti-aging, protective, and highly-stimulatory ingredients. As a result, they are a poor foundation for in vitro aging studies.

In contrast, the VitroAge medium CnT-AG2 does not contain the concentrated range of anti-aging ingredients found in standard media. Keratinocytes grown in CnT-AG2 are not exposed to these protective factors, and as a result develop a range of age-related changes during several weeks of culture.

Key changes observed in keratinocytes cultured in CnT-AG2 include decreased longevity, progenitor cell activity, DNA repair, protein synthesis, and antioxidant/stress response mechanisms. In parallel a number of cellular functions increase, including protein catabolism, protein carbonylation, and detoxification mechanisms. Aged keratinocytes can also be evaluated using a novel multiple proteomic approach that characterises approximately 100 age-related proteins in parallel. For more information, please visit:

<http://www.cellntec.com/services/cosmetics/standard/aging/vitroage>

Protocol Considerations

Duration: The standard VitroAge protocol involves culturing keratinocytes for a period of 3 weeks in either CnT-Prime (control) medium, CnT-AG2, or CnT-AG2 plus an active ingredient. Cell counts are conducted at the end of each week, to allow evaluation of the comparative proliferation rate of each treatment. Cells may be sampled for evaluation of age-related markers after either 2 or 3 weeks of culture, depending on the particular end-point of interest.

Cells: We recommend the use of early passage keratinocytes isolated from young donors (children or young adult). These keratinocytes should be growing well in CnT-Prime medium prior to transfer to the CnT-AG2 medium, to ensure that the cells will easily tolerate the transfer (transfers between unrelated media can be very challenging).

Passaging: We strongly recommend the use of gentle digestion reagents such as Accutase (#CnT-Accutase-100) when passaging keratinocytes aged in CnT-AG2. These enzymes are far gentler than the aggressive digestion obtained with enzymes such as Trypsin. Accutase does not require a separate reagent to stop the reaction, just dilution prior to centrifugation.

Centrifugation: Keratinocytes grown in CnT-AG2 remain very sticky after detachment for passaging. They adhere rapidly to the walls of the centrifuge tubes, and then do not end up in the cell pellet which significantly reduces growth following re-seeding. For this reason, it is critical that 1% serum albumin is added to the cell solution immediately after detachment, to prevent loss in the centrifuge tube.

Seeding Density: It is suggested to use the standard seeding density of 4'000 cells / cm² during the first passage of culture in CnT-AG2. As the proliferation rate gradually slows in weeks 2 and 3, the seeding density may be increased gradually (for example to 8'000 cells per sq cm), to obtain a larger number of cells for sampling at the end of week 3.

Medium Changes: We recommend 3 changes per week (Mon / Wed / Fri).

For further information about the standard culture protocols we recommend to use with CELLnTEC media, please visit the protocols page: <http://www.cellntec.com/products/resources/protocols>

For other technical questions, please contact our scientific team directly via scientist@cellntec.com