**The cellular senescence unification model and telomerase therapy: To treat all age-related diseases**

Steve Liebich

Department of Biomolecular Science and & Chemistry, Clarkson University, 10 Clarkson Ave, Potsdam, NY 02119

Correspondence: Email, liebicsf@clarkson.edu; Phone (315)-262 7005

ORCID: 0000-0002-5379-4532

**Abstract**

Since the discovery of the telomere by Hermann Muller and Barbara McClintock in years 1938-1940, great progress has been made in molecular genetics and in the relatively new field, biogerontology. Almost 40 years have passed since the discovery of telomerase by Carol Greider and Elizabeth Blackburn (1984). The scientific community has linked many naturally occurring ageing mechanisms

in cells to the shortening of telomeres and the lack of telomerase activity in these cells. A great number of mutagens, radiation, and toxic chemicals negatively impact the length of telomeres with their truncation triggering a fatal cascade of events inside the cell which can lead to the state of senescence and eventually to cell death. Even though cellular and bodily ageing is a complex, multilevel, and highly orchestrated natural process happening at the subcellular level of all multicellular organisms, and a single known mechanism is extremely insufficient to explain all observed molecular and morphological changes, a unified cohesive theory of ageing must exist. The cellular senescence unification (CSU) model, combining the free-radical-mitochondrial and telomeric theories, helps establish a strengthened base for future therapies of all age-related disorders. If the CSU model is strong enough to explain and describe most all observed alterations in the ageing cell, a focused and deliberate therapy might be developed and follow its footsteps. This work introduces telomerase therapy as an efficient, highly effective, and clinically favored treatment of most age-related disorders that can be explained by the CSU model. This gene therapy approach is another natural step forward since the first discovery of the telomere 80 years ago.

**Key words:** telomeres, telomerase therapy, cellular senescence, unified theory of ageing

**Introduction**

Each eukaryotic cell division leads to a gradual sequence loss at the chromosomal termini known as telomeres [1]. This so-called end replication problem (ERP) forms the basis of cellular senescence, along with a few other well-established biochemical processes disrupting the cellular homeostasis [2].

Telomeres are composed of repeated oligonucleotide sequences supported by a six-protein complex (shelterin) and the quaternary structure they form *in vivo* [3-5]. Thus, telomeres are the specific genomic protection device, guarding the cell from inexpedient recombination events, degradation through the DNA repair system, and significant genetic material loss [6-9]. In some cell types, a reverse transcriptase known as telomerase, reconstitutes the original length of the shortening chromosomes, saving the chromosomal and genomic stability of the cell [10-14]. In such conditions, the cell undergoes almost an infinite number of mitotic cycles, breaking through the Hayflick phenomenon [15].

Cellular and bodily ageing result from complex, multilevel, and precisely controlled mechanism of decline in effectiveness of physiological processes in the cell and decline in the stability of its genome. Because of this complexity, more than three hundred different or slightly different theories of ageing have been proposed [16]. Many theories overlap, while the others leave gaps too broad to be omitted. New data are very often misleading and contradicting, thus not supporting either of the theories. Therefore, biogerontology needs to find a common denominator for all credible theories, so that one elegant unified theory could explain the entire life-long process of cellular ageing.

All proposed ageing theories are segregated into two main categories: programmed and non-programmed [17]. The first category takes into account all ageing factors genetically inherited and gradually manifested over the span of a lifetime. The second group includes theories emerging from the customarily occurring errors in the genome and accumulated damage in the cell; tear-and-wear is preferably chosen as the *classic* theory considering accumulating damage in the genomic DNA. However, it had been the free-radical and mitochondrial theories that took much of the scientific community’s appraisal in the last century [18-21]. Three decades ago, the telomeric theory of cellular ageing stole the spotlight and since has grown into a well-developed and data-supported biogerontological doctrine, which aims to explain most of the observed hallmarks of cellular senescence [22-25]. These two models, the free-radical-mitochondrial and telomeric theories, have the potential to form a single comprehensive model of ageing explaining the entire complexity of the process. It is referred to here as the cellular senescence unification (CSU) model. For clarity purposes and due to space limitation, only the telomeric theory of ageing will be introduced and described.

Is telomerase, the enzyme of capacity to restore shortened telomeres and stabilize the genome, the definitive answer to the puzzling problem of ageing? Could we take advantage of this finding and apply it to humans, hence aim to improve health, fight age-related diseases (ARDs), and even reverse the ageing machinery in their cells? This article elaborates on the subject and aims to address these two questions with proper and preservative analysis of recent advances in the field of biogerontology.

**The cellular senescence and ageing unification theory**

As Bodnar et al. showed, each human cell which does not express an active hTERT (human telomerase reverse transcriptase) transcript loses its terminal chromosome repeats [11]. If cellular senescence was viewed as a dense system of all cellular and genomic changes occurring over an undefined span of time, relative loss of telomeres would be both one of the causes and effects of the total intracellular changes. By now, telomere loss and lack of telomerase expression in ageing cells is one the most studied and established models in ageing biology. However, it is not, and it must not be treated as *the* cause of gradual cellular dysfunction.

Since the paper of L. Hayflick and P. Moorhead on the limited mitotic capacity of somatic cells, the link between restricted cell doublings and their mortality, telomere shortening and the inevitable replicative senescence phenomenon has become obvious [15]. A large number of publications have been dedicated to the association of telomere shortening with numerous age-related disorders (see below).

A few aspects of cellular senescence must be considered before further discussion. Senescence is not equivalent to the quiescent state of the cell [26]; cell lines like hepatocytes and corneal endothelial cells maintain replicative capability, but do not divide without external stimuli [27, 28]. The all-or-nothing model of cellular senescence, popularized in the past century, has lost its merits over the more consistent and data-supported gradual changes within the cells [22]. Today, the cell senescence model of aging is acknowledged and the postulate that senescence and the non-programmed errors accumulated in genome result from changes in, among others, gene expression and the epigenetic landscape seems to be satisfying and unify other related theories [29].

Due to space limitation and deliberate focus on the telomeric theory, some programmed and non-programmed factors of cellular ageing further explained will be discussed in the light of telomeres shortening, however, those factors reach beyond the telomere biology and affect a plethora of other intracellular sites and processes. Recent research advances in biogerontology show a moderately strong correlation between telomere shortening and other possible processes contributing to cellular ageing, which all cumulatively (and definitely not singlehandedly) lead to the senescent state of the cell, which can be defined as a quiescent (non-mitotic) state of the cell unresponsive to the external stimulation and such cell is found in the G0S stage of the cell cycle [122]. As discussed below, telomere shortening, chromatin destabilization, change in genome dynamics, uneven generation of free radicals, and alteration in gene expression patterning are unequivocal correlated, which may help to draw more definitive conclusions in the future (Figure 1).



**Figure 1**. **Cellular senescence: external and internal factors.** A plethora of mutagens, signaling pathways, cytokines, and oxidative agents influence the rate of telomere shortening in the cell. Environmental stressors (UV light, industrial toxins, carcinogens, and intercalating agents), expressed oncogenes, and chromatin alterations also affect the progression of cellular senescence. Tumor suppressors, including p53, are activated upon those deleterious signals and respond adequately by triggering mitotic arrest, altering cell morphology, secretion of growth factors, cytokines, and apoptotic factors. Telomeres are affected by all aforementioned senescence factors, but the same factors induce their effects through other genomic and non-telomeric pathways. Abbreviations: hSIRT1 (NAD-dependent protein deacetylase sirtuin-1); Chk2 (checkpoint signaling kinase 2). Figure adapted from [51].

After all, senescence is a gradual process occurring in every cell of a multicellular organism, including cells that are mitotically inactive. Senescence should be understood as a complex network of changes in genes expression, DNA damage, ineffective DNA repair mechanisms, misbalance between reactive oxidative species production and scavenging, cell morphology, toxins accumulations, proteomic changes, and finally loss of telomeres. The telomeres field has had over half a century of research and promising findings which in consequence makes it both one of the most studied cellular senescence biomarker and the target for potential gene therapy; the encouraging studies of re-lengthening the shortened telomeres in animals (*in vivo* and *in vitro*) and in humans (*in vitro*) only seal the inevitable importance of the telomeres role in ageing. Even though it is subjective and experimentally unapproachable to test whether a cell is already senescent or “not yet,” the CSU model is supported by our current knowledge, and although not perfect, it is still the best tool we can use to not only explain cellular ageing processes, but also to manipulate them.

Fossel (2012) suggests the relative telomeres length measures as a reliable, clinically practical, and specific biomarker of age-related diseases [30]. Correlation between the telomere corrosion in a chronologically old individual and onset of various age-related diseases is well-known, thus telomeres and their shortening seem to be a reliable source of clinical information and a platform for proper intervention. It must be noted that it is the relative *rate* of telomeres shortening, not just a total loss of their sequences. This gives an insight into the onset of telomere-dependent diseases and their clinical manifestation [31]. This remark has been taken to clinics with commercial enterprises of key telomere researchers: Maria Blasco (*Life Length*) and Calvin Harley & Elizabeth Blackburn (*Telome Health*). These are only the first steps of the telomere research evolving into clinical importance.

However, as long as the relative telomere loss is a strong biomarker in chronologically advanced patients and individuals in greater risk cohorts, the very common telomere length (TL) measure in peripheral blood leukocytes (PBL) seem to be an undesirable method. First, false negative results might affect the diagnosis of a patient still affected by slowly evolving pathological processes. The reason is that leukocyte telomeres might remain relatively stable if a disorder develops in the liver, for instance; moreover, the “old” leukocytes are constantly being replaced by the white blood cells with long telomeres. Second, peripheral leukocytes can be exposed to toxic, stressful or immunological factors that would influence telomere loss in the PBL without any underlying age-related disorder [32]. Multiple studies showed a positive correlation between external factors such as smoking, obesity [33], oxidative damage [34, 35] and past infectious diseases [36-38]. Third, the genomic changes of white blood cells (WBC) are just as important as in other human cells: the age-dependent telomere shortening [39], polymorphisms in the *hTERT* promoter and its regulatory genes [40] (although a Swedish cohort studies of the same single nucleotide polymorphism did not show any correlation [41]), and changes in the epigenetic landscape of the *hTERT* [42]. These findings indicate that the relative PBL telomere loss is a significant biomarker for a number of age-related diseases, but PBLs are not always the right source of telomere attrition information.

Coronary heart disease [43], osteoporosis [44], diabetes [45], and other age-related disorders are correlated with shortened telomeres; the telomere attrition leads to cell senescence as observed in multiple tissues and organs. PBL telomere length (TL) measurements have been applied to a plethora of various age-related disorders, diseases of affluence and immunological disorders, including longitudinal studies of cardiovascular health problems (46), chronic obstructive pulmonary disease [47], familial and sporadic pulmonary fibrosis (48), and hematopoietic malignancies [49]. These data underlie a strong correlation between telomere attrition and the etiopathophysiology of diseases observed in clinical settings.

Human skin fibroblasts were the first telomere-associated senescence model cells, for which the telomerase transfection proved to be liable [11]. In two parallel studies, human keratinocytes and fibroblasts were grown on an immune-compromised mouse and the new skin morphology was assessed for early (20 population doublings) and late (85 population doublings) passages; the late passage cells were further transformed with an external *hTERT* and the re-lengthened telomeres led to skin reconstitution, optimal gene expression and normal filamentous connections observed in the young skin [50].

Almost every tissue type in the human body demonstrates histological and ultracellular changes associated with telomere shortening. For cardiovascular diseases, mice and human models show exceptional correlation between telomere dysfunction (also oxidative stress, proinflammatory molecules activation, etc.) and senescence of vascular endothelial cells leading to development of cardiomyopathy and severe atherosclerosis [51]. The impairment of control mechanisms of stem cell reserves and their differentiation and division in the bone marrow, hugely associated with telomere attrition and immune system age-related changes, is responsible for the pathological decrease in activity in older people. Similar deferment in physiology has been found in glial cells, the cells that divide and proliferate in the brain, whose ultracellular ageing transformation has been presented to be the leading cause of Alzheimer’s disease [52]. These examples indicate the necessity to reform the way how one interprets the pathophysiology of many age-related disorders. The non-divisional pattern of neural or myocardial cells would lead to an inaccurate and misguided conclusion that the telomeric theory could not be taken into account for these cells. However, this inaccuracy stems from the wrong perspective: one must shift focus from the mitotically inactive to the dividing cells found in the proximity of the non-divisional cells. The combined pathophysiologic outputs of the non-dividing, but still homeostatically deteriorating cell and the mitotically active cell, whose telomeres do truncate after each round of mitosis (next to other countless ultracellular ageing processes) draw a more consistent and self-explanatory picture of age-related disorders.

**Control mechanisms of telomere length and telomerase expression**

Senescence is a gradual process that takes place in every living cell of the human body, with cells varying in degrees of senescence. This is a complex and global process incorporating changes in epigenetic status, genomic mutations and breakages, slowed metabolic turnover of the cell, increased levels of oxidative species, failing mitochondria, accumulation of toxic chemical compounds (external and internal), deterioration of signaling pathways, increased turnover of proteins, and increasing area to volume ratio of the cell.

Not all changes lead to the cell’s death though. Senescent cells rarely “die,” they rather move to a quiescent state where they become less or completely mitotically inactive. However, shortened telomeres in somatic cells advance genomic defects, and this destabilization leads to more rapid cellular changes [23, 25]. On the other hand, stem cells are partially protected from the instability by expressing telomerase, an enzyme that extends the missing chromosomal termini. The difference between somatic and stem cells is that the former has an accelerated rate of senescence, while the latter gradually become more senescent [53, 54]. Thereby with old age, stem cells are more senescent than they were in the young organism, but somatic cells are more senescent than the stem cell populations. While most organs store their somatic stem cells in so-called niches, which function as “spare parts” for the ageing organ, the process of replacing aged and/or dead somatic cells with transit cells (dividing and slightly differentiating stem cells) demonstrating self-renewal capability diminishes [54-56].

Telomerase plays an important part in keeping the cell’s homeostasis balanced. Therefore, telomerase expression and activity regulation in the cell is highly significant. The 40-kb *hTERT* locus sits on chromosome 5, only 2 Mb away from the chromosomal end [57]. *hTERT* lacks both TATA and CAAT boxes but is regulated with two canonical E-box sequences [12] (Figure 2). The E-box sequences and numerous CpG islands provide large methylation sites for the *hTERT* locus, which demonstrates high epigenetic regulation [58]. Most human cells do not produce telomerase [12]. The exceptions are stem cells, germ cells, and select white blood cells [59, 60]. Moreover, 85 – 90% of cancer cells express active telomerase [61-63]. It is still unknown why most somatic cells are hypermethylated at the telomerase site and what exact mechanisms participate in gene negative regulation [64]. However, various findings show that some cells are capable of producing telomerase *in vivo*, but these are inactively spliced variants that do not contribute to the telomerase activity [65].

The telomerase-telomere cellular “apparatus” is also controlled by a wide network of proteins. The *trans*-acting regulatory proteins are extremely active at the *hTERT* locus. The most abundant transcription factor (TF) complexes are E-box-binding c-Myc/Max complex, which co-operates with the GC-box far-flung Sp1 TF acting as the activating element [66]; on the other hand, Mad1/Max complex, along with deacetylases, represses the *hTERT* locus [67]. To review the divergent transcription factors involved in the telomerase expression regulatory mechanisms refer to [68-70].



**Figure 2. Telomerase expression mechanism in telomerase-positive cell line.**Telomerase is a ribonucleoprotein complex composed of many subunits, of which telomerase reverse transcriptase (TERT) and telomerase RNA component (TERC) are essential in physiological activity of telomerase. The multicomponent enzymatic complex extends chromosomal termini by synthetizing *de novo* tandem telomeric repeats (5’TTAGGG-3’) at 3’ end of the chromosome. TERC binds the 5’ C-rich strand and its 11-nt sequence serves as the replication template for reverse transcription. Abbreviations: hTERT (human telomerase reverse transcriptase); hTR (human telomerase RNA component). Figure adapted from [115].

Telomeres are supported by another, but not exclusive, set of proteins. The already mentioned shelterin complex, composed of six core proteins, both protects telomeres from exonucleases degradation and provides an interaction matrix for other regulatory factors [71]. The telomeres instability is reflected mainly through the double-strand break (DSB) events. The DSBs induce chromosomal translocations, error-prone recombination events and tumorigenic processes [72]. Two major DNA damage sensing pathways are related to ATM and ATR serine-threonine kinases, which are activated at cell cycle checkpoints, mostly at G0, G1 and S; the DSBs are repaired by two main processes: homologous recombination (HR) and non-homologous recombination end-joining (NHEJ) [73]. A plethora of proteins are used by the cell to protect the genomic stability and the truncated telomeres through the two recombination mechanisms [73, 74].

This complicated and heterogeneous network of signaling pathways and *trans*-acting regulatory elements serves to maintain telomere length and respond to occurring changes at telomeres. Recently, the telomere position effect (TPE), imposed by relative shortening of telomeres and the rate of loss affecting nearby genes through telomeric chromatin looping has attracted much attention [75, 76]. This is because the TPE explains how physical changes of the truncated telomeres affect other gene expression through physical mechanisms, very often affecting genes placed more than 10 Mb from the telomeric region [75]. Additionally, short telomeres themselves may serve as another proximate senescence trigger through DNA repair mechanisms [77]. However, many researchers agree that, apart from the aforementioned mechanisms, the relative length of telomeres and the rate of telomere loss affect the genomic structure at most [30, 53, 78, 79]. The cell responds accordingly to all these changes, and depending on its current biochemical status, becomes more receptive to other deleterious processes inside of it.

**The role of telomerase in cellular pathophysiology**

The negative correlation between telomerase expression and onset of most cancers in the human body has been favored by the scientific community for years [61-63]. The cancerous cells’ *replicative immortality* stressed by Hanahan and Weinberg has been associated with the *hTERT* locus activation, and thus the conclusion withdrawn was simple: telomerase expression promotes tumorigenesis in somatic cells [80].

Interestingly, the remaining cancer types that do not activate the *hTERT* locus still elongate their telomeres through the alternative lengthening of telomeres (ALT) mechanism [81, 82]. This route of telomere elongation takes an advantage of chromosomal instability and strands breaks, which is detrimental to the cell karyotype. A tumorigenic clone line will always sort the cells with longer telomeres compensating the higher proliferative rate [81]; furthermore, the telomere sister chromatic exchange (T-SCE), telomere-repeat arrays exchange resulting from the ALT pathways, yields the unequal sister telomere lengths in different cancerous cell lines [83].

 The misconception is that long telomeres are a biomarker for high cancer risk; this is far from the truth. In fact, in most cancerous cells the telomeres are much shorter than those found in normal cells [84, 85]. Most recent tumor studies find that telomeres in the cancer cells are, through structural analysis, truncated and recombinogenic: dicentric chromosomes, reciprocal translocations and other deleterious karyotypes [86, 87]. Moreover, in most studies correlating the telomere length with concomitant cancer, the telomere length has either been found normal or reduced. In other studies, the telomere length has been found increased with either active or quiescent telomerase [88-91]. In many papers studying the correlation between telomeres length, telomerase expression and cancer cell biology, the histopathological changes in cancer tissue are not reported. Not only is the average length of the cell telomeres measured instead of the shortest telomere length, but these telomere length measurements are taken from the patient’s PBL in malignant diseases not correlated with immune system [30]. These mistakes generate false conclusions and consolidate the misconception.



**Figure 3. Relationship between telomere shortening, increasing genomic instability and varying cancer risk.** With each replicative division the total telomere length decreases, which leads to miscellaneous genomic and intracellular alterations increasing the overall risk of hyperproliferation and tumorigenic mechanisms development. It must be noted that a cell in order to develop cancerous changes must rely on relatively long telomeres for proper genomic stability. If its telomeres are too short, the cell drifts into senescence and eventually dies. Figure created by Michael Fossel, MD, PhD.

There is much evidence on the relation between telomere length maintenance (TLM) and cancer. However, there are no data indicating that cancer is caused by overexpression of telomerase or overly long telomeres. Present studies with somatic cells expressing a recombinant telomerase gene or somatic cells treated with telomerase vectors, and studies on the embryonic pluripotent cells with constitutive telomerase expression, indicate no cancer remarks [26, 92-94]. This can be partially explained by the fact that only the average telomere shortening and its rate, not the total telomere loss, determine the senescence phenotype; thus telomerase expression increases genomic stability by keeping the average telomere loss and re-lengthening constant [95]. Simultaneously, the number of cell divisions increases (uncontrolled growth), which leads to further telomere truncation which must be overcome by a higher rate of telomerase expression (Figure 3). These findings can be supported by showing that knocking down the telomerase gene coincides with tumor growth arrest, but only when the tumor had already grown, not before the tumorigenesis process, not even in a p53-null cell [79].

However, telomerase expression does not serve its canonical function only. Recent findings present multiple non-canonical functions of telomerase, which influence genomic stability, very often resembling the ones found in the ALT mechanisms [96]. All in all, both the telomerase- and ALT-based telomere length maintenance pathways stabilize the already destabilized genome of the cancerous cell. Thereby, it is not about a maximum telomere length that can keep the cancer cell from apoptosis, but about keeping telomere length above the critical threshold, different for each organism. This also explains why tumor cells in mice upregulate mTERT expression, even though the mural telomeres (approx. 50 kb) are much longer than in humans (approx. 15 kb in a young human cell) [78]. This notion does not hold true for some individuals, however, if their cellular environment manifests a variety of mutations (genome) and preconditioned disorderliness in biochemical pathways and DNA repair mechanisms (cell); although in these cases the telomerase expression introduces new level of complexity to the unbalanced system rather than becoming the leading cause of tumorigenesis. The studies in mice corroborate the observations that, contrary to the general assumptions, telomerase does not increase the risk of tumor development, but also protects the cells from such [97, 98].

**The beginnings of telomere-targeted therapies**

The number of publications on telomerase and its different biological effects has immensely increased since the dawn of the 2000s [99]. Since the discovery of telomerase, its impact diverged into two schools of thought. The first school is “pro-TERT,” favoring its regenerative function at shortened telomeres, extending the cell’s population doubling, and restoring mitotic capacity. The other school is mainly “anti-TERT”, which extends to applying knowledge of telomeres, telomerase, and their investment in the cell’s homeostatic profile in treating miscellaneous cancerous diseases. In fact, the anti-telomerase therapies have been widely exploited in cancer research in the last three decades.

Anti-telomerase molecules have been chosen by selected biotechnology companies as potential drugs, mostly targeted against cancerous tissues [100-102]. One such pharmaceutical, N3’-P5’ *thio*-phosphoramidate (NPS) oligonucleotide (GRN163) acting as a telomerase antagonist, although primarily shown as a competitive and efficient cancer drug introduced by Geron Corporation, had turned out to deliver no expected results. Interestingly, the anti-telomerase therapies seem to impair the findings of the opposite school, i.e. upregulating telomerase levels in the cell for therapeutic reasoning. This only showcases that the situation is far more complicated than the initial findings had indicated.

It is still inconclusive and more research must be done to sincerely state that telomerase itself does not increase the risk of cancer development, but there are more than plain premises helping to conclude the germinal findings about telomerase’s role in cancer [103]. It must be highlighted again that it is optimal telomere length that must be present in a cancerous cell in order to satisfy the unstoppable growth. If the telomeres were too long, they would provide the cell with incontestable genomic stability, thus precluding cancer development [104]. On the other hand, if the average length of the cancerous cell’s telomeres was too small, the cell would fall into the senescence route and eventually die through apoptosis [105, 106]. Therefore, we should expect different clinical results depending on the dosage of anti-telomerase therapy, duration, and the activity of anti-telomerase agents. Perhaps, this sort of cancer treatment would yield positive clinical effects in a short run, but the long-term consequences seen in the patient are difficult to predict, especially if to consider the complexity of the tumorigenesis process and the still uncertain role of telomeres and telomerase in its development.

Thereby, instead of focusing sole attention on cancer in the light of telomerase, we should consider the telomerase therapeutic methods that could potentially help confine age-related and other degenerative diseases. Some agree that telomere extension through the reverse transcription of telomerase complex outweighs the risk of cancer promotion [107].

Transient transfection of somatic cells with telomerase transcript, full or truncated, has shown that 1) a causation relationship exists between the length of telomeres and the expressed levels of telomerase, and 2) extrinsic delivery of telomerase to the cells restores their function, but does not change their phenotype [26, 108, 109]. The first proof was presented by Bodnar’s laboratory in 1998, when human retinal pigment cells and foreskin fibroblasts were transfected with the hTERT-SV40 construct, and established the laboratory immortal cell lines [11]. Bodnar’s work initiated a cascade of future experiments using extrinsic telomerase catalytic subunit as the cells’ ‘transformer’, establishing immortalized and phenotypically regular transformed cells [110-112]. The groundbreaking experiment presented by DePinho’s lab using a special mural TERT transcript engineered with a knock-in TERT-estrogen receptor (mTERT-ER) allele controlled by an extrinsic estrogen receptor modulator, 4-hydroxytamoxifen (4-OHT) showed how reversing the systemic degenerative phenotypes and restoring telomere length in mice with various telomere dysfunctions can be achieved by external, dosage-dependent telomerase expression regulation [92]. Other studies with somatic cells successfully transfected with TERT subunit are reviewed in [107].

From time to time, different therapeutic variations concerning telomerase are reinforced by other biomolecules. In one such study, telomerase therapy was supported by ectopic Bcl-2 transfection [113]. Telomerase as a biomolecule in tissue engineering and regenerative medicine (TERM) has already been strongly suggested [114, 115]. Moreover, it is commonly agreed that transient, ectopic delivery of telomerase under constant physiological control is the only acceptable way for a therapeutic approach. This model, aided by viral vector technology, had been well-tested in the past [50, 116, 117]. The same model, but modified and extended, works perfectly fine with adeno-associated virus (AAV) as a viral vector for ectopic TERT delivery, which does not cause tumor transformation or any genomic instabilities within the transfected cells [93]. Those advancements show that we are ready to embrace experimental data coming from telomere research and apply this knowledge to clinics, where patients could benefit from billions of dollars spent on telomerase, cancer, and gene therapy research.

**The future of ageing: Telomerase therapy** **aims.**

Telomerase therapy is rooted in the molecular foundations of cellular pathology. As such, it might target multiple potential diseases directly associated with genomic instability and/or TLM disruption. Every age-related disorder with an etiology correlated with telomere shortening, supported by experimental and clinical data, can become a potential target for therapy. Therapy may not only be used to cure age-related diseases, but it also can be applied to prevent their occurrence. The premise is clear: if we can suppress genomic instability and strengthen cellular homeostasis, the risks of disease onset and its progression might be reduced.

Previously in the text, reversing cellular senescence was mentioned. This is an important aspect of every biogerontological therapy which must be addressed with high prudence. From a biomedical standpoint, we are currently not able to *reverse* molecular ageing in terms of the simple understanding: one cannot reverse the biological clock as this would stand against the second law of thermodynamics. However, one could help suppress the somatic damage to the genome and prevent further deleterious changes happening at the molecular level. These actions will reverse the molecular hallmarks of cellular senescence, giving an individual a better quality of living.

Moreover, as it was already discussed, a strong correlation between cancerous transformation and the telomerase apparatus control exists. If the total loss of telomeric repeats in a given set of chromosomes dictates further mechanisms being activated and deactivated during rigorous checkpoints of the cell cycle, preventing the cell from telomere truncation would yield broadly positive clinical outcomes for cancer. More experimental data is needed to draw proper conclusions, and it is still too early to declare any final statement on the correlation between cancer development and telomere length control, but based on the data we have at the moment we can conclude that there is a promising platform for telomerase therapy targeting cancerous changes at the molecular level. In such terms, telomerase-based gene therapy could become a splendid showcase of the potential of telomerase: to cure age-related disorders and to protect from cancer with one single platform.

**Evaluating telomerase therapy**.

Human clinical studies must be completed before we can state that the therapy is successful and rational. Predictions set a new course of developing gene therapy for age-related diseases, where a biopsychosocial model of disorder must be evaluated for the full understanding of potential telomerase therapy. Telocyte, an American biotechnology company, is one of the first commercial entities focused on producing an effective telomerase-based gene therapy platform to cure Alzheimer’s disease and other age-related diseases. As discussed thoroughly by Fossel in his recent publication on the role of telomeres in developing Alzheimer’s disease and other dementias, even the ARDs of the brain, with the main functional cells, neurons, mitotically inactive and fully differentiated, can still be explained on the basis of gradually shortening telomeres by taking into account the remaining 80% of the brain’s glial cells which are capable of mitotic division [52]. Because glial cells are supportive cells, their morphological and physiological deterioration affects the neural cells more than it had been predicted [118, 119]. If the main upstream cause of age-associated dementias are the senescing glial (supportive) cells, whose deteriorated state stems from genomic instability triggered mainly by shortening telomeres, in this case the progressive functional failure of neurons and the occurrence of the hallmarks of Alzheimer’s disease (tau tangles and β-amyloid plaques) can be fully understood, pathophysiologically, through the biology of telomeres. Therefore, Telocyte’s prototypic treatment system is based on extracellular telomerase transcript delivery. It will be done through the AAV-9 vector injected into the spinal fluid and acting transiently on the glial cells. If successful, this gene therapy approach builds a bigger therapeutic platform which will serve to eventually cure most age-related disorders.

The biology of ageing is a crossroad for miscellaneous fields of medicine and biology, where one must not speak about one cause, one effect or a simplified correlational model. Using novel proteomics of an isolated chromatin segments (PICh) technique, it has been found that three cell lines manifesting two types of TLM, through telomerase and ALT, were associated with ~400 different proteins, sharing 98 proteins in common [118]. Many of those proteins could be associated with different genomic loci, cell cycle stages and chromatin looping effects, but this approximation showcases the great abundance of regulatory elements involved directly or indirectly in the processes of telomere length regulation (TRF1, POT1, Apollo), DNA damage mechanisms (Rad50, BLM, ERCC1, PARP1), chromosome end protection (Ku 70, Ku 80, ATM, ATR, WRN), and many others [119]. Large numbers of proteins on the cross-talk of different paths must be counted as well, which only adds on to the complexity of the biology of telomeres.

**Conclusions**

Should a therapy based solely on telomerase be trusted? This question requires a definitive answer based only on data; no speculations have room here. However, we are certain that telomerase therapy might become a gold standard for all age-related diseases in the next 10 years. Recent findings (cited elsewhere in the text) showcase undoubtedly the positive role of telomerase in “old” and senescent cells, which retrieve their biological functions and add to the pool of “young” and healthy cells of the tissue. The exact role of telomeres and their length control in cancer development is still debatable, but further steps in clinical research will finally resolve this dilemma.

There is yet much to say in ageing research. New discoveries consistently broaden our understanding of telomere biology and demonstrate its complexity, which is understandable in terms of its highly significant role in homeostasis. It must be noted that, even though many signaling pathways and metabolic processes are involved, the main role of telomeres is to protect the chromosomal ends from degradation and NHEJ, which destabilizes the genome and disrupts cellular physiology. If this process can be stopped, reversed, and stabilized in the metabolic environment of the cell, we should do it. We must push the progress of our own findings and apply them in real-world situations, where patients are the final recipients of what once was started in the laboratory.

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**Author’s contributions**

Steve Liebich is the sole author of the article.

**Conflicts of interest**

The author declares no conflict of interest.

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