Simple Detection Methods for Senescent Cells: Opportunities and Challenges

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Cellular senescence, the irreversible growth arrest of cells from conditional renewal populations combined with a radical shift in their phenotype, is a hallmark of ageing in some mammalian species. In the light of this, interest in the detection of senescent cells in different tissues and different species is increasing. However much of the prior work in this area is heavily slanted towards studies conducted in humans and rodents; and in these species most studies concern primary fibroblasts or cancer cell lines rendered senescent through exposure to a variety of stressors. Complex techniques are now available for the detailed analysis of senescence in these systems. But, rather than focussing on these methods this review instead examines techniques for the simple and reproducible detection of senescent cells. Intended primary for the non-specialist who wishes to quickly detect senescent cells in tissues or species which may lack a significant evidence base on the phenomenon it emphasises the power of the original techniques used to demonstrate the senescence of cells, their interrelationship with other markers and their potential to inform on the senescent state in new species and archival specimens.

Keywords: ageing, senescence, lipofuscin, labelling index, detection

WHAT IS A SENESCENT CELL?

Historical Background

The initiation of the first true cultures of metazoan cells by Alexis Carrel (Carrel, 1913) is a landmark in the history of biology. Carrel was primarily interested in understanding the regulatory factors controlling wound healing. In attempting to do so he established cell culture as a valuable experimental tool for the study of cell morphology and function. Gradual refinement of tissue culture techniques have allowed the culture of progressively more complex cell types and the development of model systems for the study of many fundamental biological processes.

Initially however, studying ageing in vitro appeared impossible. Although it had been proposed that exhaustion of the growth capacity of somatic cells was responsible for organismal ageing (Wesemann, 1889), Carrel's best known experiments appeared to falsify this hypothesis. Carrel initiated cultures of chick fibroblasts which his co-workers claimed to have cultured continuously for over 30 years, much longer than the lifespan of the intact organism (Ebeling, 1942). It followed that the ageing process had to operate at the level of the tissue or organism not the single metazoan cell, which seemed as immortal as its bacterial counterpart. As tissue culture gained in popularity through the 1950s the failure to produce cultures of "immortal" cells was attributed to poor experimental technique. This was not innately implausible given the extraordinary difficulties then involved in carrying out tissue culture experiments that today would be considered trivial. This "poor culture
technique” argument has reappeared regularly as cytologists have pushed the boundaries of tissue culture to the technological limits of their day (Perillo et al., 1989) and will doubtless reoccur in the future, sometimes justifiably.

However, in the early 1960’s (Hayflick and Moorehead, 1961; Hayflick, 1965) Hayflick demonstrated that normal human fibroblasts would only proliferate for a finite number of passages in culture. Latent infectious agents, composition of the medium and depletion of key metabolites were all shown not to be responsible for this failure to grow. Hayflick also observed that fibroblast cultures initiated from embryos grew substantially better than those derived from adults. Combined with the demonstration that normal somatic cells displayed limited lifespans during serial transplantation experiments these observations, led Hayflick to formulate a cellular theory of ageing (Hayflick, 1965; Hayflick, 1979). The original tenets of this hypothesis were that a finite lifespan is an intrinsic property of normal human cells, that cell growth in vitro is somehow related to human ageing and that cultured primary fibroblasts were a useful model system in which to study some aspects of in vivo ageing. Hayflick termed this failure to grow “senescence” a term which, for good or ill, has stuck.

Since Hayflick’s original discovery many different cell types have been shown to behave analogously to human fibroblast populations in vitro. Senescent cells have also been shown by a variety of methods to be present in many different tissues in vivo. This finally silenced the criticism, not uncommon among gerontologists until the 2000s, that senescence was simply a “tissue culture artefact” unworthy of study. Rather, senescence appears to be a ubiquitous characteristic of cells derived from regenerative somatic tissue and a primary causal mechanism of ageing and age-related disease.

However, as the list of cell types showing senescence has grown specialists in different branches of biology have found themselves grappling with questions once restricted to cytologists working predominantly on fibroblasts. How can senescent cells be simply and reproducibly detected both in vitro and in vivo? and to what extent can the same techniques be used in different cell types and different species? To address these questions this article will review the different markers available for the detection of senescent cells with an emphasis on the phenotypic aspects of senescence that originally validated them and how these in turn may inform on the senescent state in other contexts.

**Terminology**

Clarity is essential when discussing the biology and growth dynamics of cells in vitro because the literature uses overlapping, sometimes contradictory, terms for the same cell populations (Hayflick, 1990). Without prejudice to other classification systems, this review will follow the usage of Shall (1987) in which cell cultures can be considered to fall into three types with respect to growth capacity: Primary cells are derived from normal tissue and display a limited lifespan in culture (although in some fields only the initial explant population is referred to as “primary”). “Primaries” are sometimes referred to as mortal cell cultures or as cell strains-the term originally used by Hayflick and favored by this author [although some researchers restrict use of it to subpopulations selected from a culture by cloning, e.g., Freshney (2005)]. Continuous cell lines are by definition composed of cells with an unlimited growth potential (mortal cell populations sometimes being termed finite cell lines). In the Shall typography continuous lines fall into two broad categories; immortal cells are non-tumorigenic, possess unlimited proliferative capacity and at that time were largely derived from rodents (e.g., 3T3 cells). By contrast, transformed cells represented the vast majority of cell lines produced from humans (e.g., HeLa or HepG2 cells). Transformed cell lines have the unlimited lifespan of continuous cell lines but form tumors in nude mice and show a number of other characteristics, such as growth in soft agar, which reflect this tumorigenicity in vitro. The underlying molecular mechanisms controlling growth and giving rise to this typology have become progressively clearer over the years (e.g., Shay et al., 1993; Zou et al., 2009).

Only cell strains spontaneously become senescent in culture, although senescence can be induced in cell lines through a variety of methods (typically, but not exclusively, involving DNA damage). Senescence is distinct from quiescence, a reversible growth arrested state which can be induced by the removal of serum from the medium of primary or immortal cells or through contact inhibition. In systems in which it is experimentally possible to separate the two states (e.g., epidermal keratinocytes grown in low calcium medium) senescence is distinct from terminal differentiation (Norsgaard et al., 1996).

“Senescence” is also regularly used in two distinct ways which can occasion further confusion. Senescence of the entire culture, originally called the “Phase III phenomenon” (Hayflick, 1979) is failure of the culture to proliferate under conditions which had previously allowed sustained cell growth. This growth is generally measured as Population Doublings (PDs) calculated as the number of times the cell population doubles in number during the course of culture Eq. 1.

\[
PD = \log_{10} \left( \frac{\text{Numbers cells harvested}}{\text{Number cells seeded}} \right) - \log_{10} 2
\]

(1)

Primary cell populations typically go through 30–60 cumulative population doublings (CPD) by which time they are overwhelmingly composed of senescent cells (the second sense of the word as used by cytogerontologists). It is worth reiterating that a “senescent” culture is not necessarily free of growth competent cells, although they will be in a minority. This occurs because all that is required for the population to cease to expand and enter an apparently static phase is for the rate of production of newborn cells to be less than or equal to the population death rate (Kalashnik et al., 2000). The highly variable survival times of senescent populations from different cell types reported in the literature (days or weeks in the case of senescent HUVECs compared with months or years in the case of dermal fibroblasts) results in part from this dynamic.

The canonical feature of a senescent cell is its failure to divide in response to a conventional mitotic stimulus but this is also accompanied by radical alterations in cell physiology. Together these constitute the core phenotypes of cellular senescence from
which the validity of all histological markers ultimately derive. Because it is possible to separate the altered phenotype from the cessation of growth in some contexts (e.g., the serial passage of adrenocortical cell strains) the failure of growth and the altered phenotype are considered separately below.

**THE CORE PHENOTYPES OF SENESCENT CELLS**

**The Failure to Grow**

Hayflick and Moorhead (1961) working with 25 independent strains of human fetal fibroblasts made the central finding that primary cultures cease to expand, and defined three distinct stages of growth in vitro. Phase I, the explant culture, was considered to terminate with the formation of the first confluent sheet of cells. Phase II was characterised by vigorous growth requiring repeated subculture. Phase III was a decline phase in which, after approximately 50 CPD, the cells showed cessation of mitosis, the accumulation of cell debris and degeneration of the culture (Hayflick, 1965). These three stages can be observed in a wide variety of different cell types but patience is required since 50 CPD takes roughly a year of continuous passage for fibroblasts (although fibroblast cultures with both greater and lesser proliferative capacities are common).

Hayflick's original model carried two implicit assumptions. Firstly, that the decline in proliferative ability seen during "Phase III" resulted from cell death. Secondly, that the cultures studied were homogeneous populations of cells which were either all growing (in Phase I and II) or all non-growing (Phase III). Simple cell death as a cause of Phase III was quickly excluded by the demonstration that RNA synthesis (measured by incorporation of tritiated uridine) occurred in all population phases (Macieira-Coelho et al., 1966).

Although proliferative homogeneity was disproved a little later it is still widely assumed to be a feature of normal cell populations—perhaps because Hayflick’s original figure showing the three phase model is so regularly reproduced in tissue culture manuals. But Smith and Hayflick (1974) and Smith and Whitney (1980) demonstrated, by isolating and then culturing hundreds of population doubling levels, that primary WI38 and WI26 clones at different mass culture tissue culture plates. These are created by the deposition of arrays (Pontén et al., 1983). This assay is based on the use of custom isolated glial cells using the Pontén mini-cloning technique gradually shifted towards smaller clones as the culture aged. Potentially, re-cloning a clone with a high divisional capacity contain mixtures of clones with very variable growth potentials. Re-cloning a clone with a high divisional capacity produced sub-clones with a range of division potentials that gradually shifted towards smaller clones as the culture aged. Later studies examined the replicative capacity of thousands of isolated glial cells using the Pontén mini-cloning technique (Pontén et al., 1983). This assay is based on the use of custom tissue culture plates. These are created by the deposition of arrays of some hundreds of circular "islands" of palladium (typical areas 45,800 μm² or 18,000 μm²), onto the central area of tissue culture plates that no longer allow cell adhesion (e.g., by being pretreated with agarose) whilst at the same time a ring of palladium is seeded around the outside (the overall effect is somewhat akin to islands of palladium in a lake or inland sea). In contrast to the clone isolation experiments of Smith and Hayflick (1974) or Smith and Whitney (1980) Pontén mini-cloning makes it possible to control for growth artefacts arising from low cell densities by simultaneously plating cells at normal densities on the ring of palladium forming the lake “shore” and cells at clonal densities on the islands (achieved by using a metal spacer to separate the low and high density subpopulations until they have attached). The fraction of non-dividing glial cells in these studies (n = 1760 clones) increases steadily from the earliest to the final passage (~35 population doublings) in a very similar pattern to dermal fibroblasts and represented an early demonstration that senescence could be observed in cells from multiple different tissues.

Labelling studies complemented these single cell analyses. Cristofalo and Sharp (1973) carried out an analysis of the divisional kinetics of embryonic fibroblasts. This required 72 h 3H-thymidine pulse-labeling experiments at every passage throughout the culture lifespan. The fraction of cells which entered S phase was then estimated by autoradiography. They observed that unlabelled (senescent) cells were present in very young cultures, that a few labelled cells were present even in “senescent” cultures and that the fraction of unlabeled cells increased smoothly with serial passage. Additional experiments using WI38 fibroblasts demonstrated that although overall cell cycle length (T½), increased approximately two fold (from ~19 to ~30 h) with serial passage this was insufficient to explain the decline in the labelling index (Grove and Cristofalo, 1977). Thus the fraction of growing cells in the culture declines as the CPD level increases and fibroblast cultures are mixtures of label-excluding, senescent cells and their growing counterparts, the proportions of which alter as the cultures age. If these basics are neglected when studying a primary cell population then the results of any assay that generates an average value from the culture (in practice anything from Western blotting to next generation sequencing) become fundamentally insecure.

It is often unappreciated that such kinetic analyses underpin virtually all the more “sophisticated” techniques sometimes recommended for the identification of senescent cells today (González-Gualda et al. 2021). Although labelling studies have become progressively easier to perform through the replacement of 3H-thymidine first by 5-bromo-2′-deoxyuridine (allowing label incorporating cells to be detected by immunocytochemistry rather than autoradiography) and subsequently by 5-ethyl-2′-deoxyuridine (which simplifies the technique still further through the use of a copper (I) catalyzed “click-chemistry” reaction between the 5-ethynyl group and an azide conjugated dye such as Alexa-594 or Pacific Blue) the basic principles and problems inherent to kinetic analyses are much the same today as they were in the 1970s.

The major problem associated with labelling is the need to ensure that the proportion of cells that take up the label accurately reflects the true culture growth fraction. A short labelling time (e.g., 30–90 min) underestimates this because serially passaged cultures are asynchronous and cells in G1, G2, and M cannot incorporate label. Since the minimum S phase time is 6–8 h in cultures of MRC5 and WI38 fibroblasts regardless of CPD
(Griffiths, 1984) roughly 70% of division competent WI38 cells will not be in S phase at any one time ($T_c = 19–30 \, \text{h}$). Long labels of the type used by Cristofalo and Sharf (1973) avoid this problem but overestimate the growth fraction. This is because growth fractions are calculated as the ratio of nuclei that have incorporated label over the total scored (typically 400 positive or 1,000 total nuclei giving the 95% confidence interval). With 72 h labels, cells in S phase when the label is introduced will go through $G_2$ and M phase during the labelling period and will thus be scored as two separate positive nuclei; a process that could be repeated twice with typical $T_c$ values.

A long label should thus be seen as providing a minimum value for the number of senescent cells present in a population (<5% label incorporating nuclei on a 72 h label is often used as the definition of a fully senescent culture) whilst short labels are useful for determining the decline in a growth fraction during serial passage, particularly if comparing rates between multiple cell populations (Faragher et al. 1993). In theory, $T_c−T_s$ is the perfect labelling time but in practice almost never attempted whereas labelling times of 24 or 48 h are the hardest to interpret but occur in the literature with monotonous regularity (perhaps because they are simply easier to work into routine laboratory schedules). Minor problems with analogue labelling include the frustrating feature that media which contain high levels of thymidine (such as Ham’s F12 or some specialist media with “trade secret” compositions) can cause short labels to fail. BrdU labelling is also incompatible with the simple terminal transferase dUTP nick end labeling (TUNEL) apoptosis assay due to photolysis of any DNA that has incorporated BrdU. Ironically such labelled cells make excellent positive controls in a TUNEL assay.

Given that advantages and disadvantages are inherent in the use of thymidine analogues alternative methods based on the detection of proteins varying in amount and/or conformation through the cell cycle have also been employed to study senescence. Of these antibodies against Topoisomerase II and proliferating cell nuclear antigen (PCNA) have both been used in senescence. Of these antibodies against Topoisomerase II and through the cell cycle have also been employed to study TUNEL assay.

Mechanisms of Senescence: Pathway Components as Specific Markers

In parallel with the study of the kinetics of senescence in primary populations researchers attempted to unravel the mechanistic basis of the process in individual cells. Early cell fusion experiments demonstrated that the phenotype of senescence was common between fibroblast strains from different donors, that it was dominant over growth in synkaryon fusions between growing and senescent cells (Littlefield, 1973) and that senescent nuclei or cytoplasm inhibited DNA synthesis in nuclei derived from young cells when partnered with senescent ones in heterokaryon experiments (Norwood et al. 1990). This work was complemented by the isolation of pairs of daughter cells generated by single mitotic events followed by determination of their proliferative potential. In a significant proportion of these pairs large proliferative differences between the daughters were observed indicating that human fibroblast senescence was controlled by the unequal partitioning of some controlling molecule which, although its nature at that time was unknown, was present at a concentration of less than 100 copies per cell (Jones et al., 1985). Taken together these results were consistent with a model in which senescence was controlled in the nucleus by a few elements and effected by one or more proteins which inhibited the transition of cells from $G_1$-S phase.

As proposed by Olovnikov (1973), the finite elements regulating human fibroblast senescence eventually proved to be telomeres (which progressively shortened with cell division due to the absence of the repair enzyme telomerase). The primary effector protein, initially isolated by expression screening as
senescent cell-derived inhibitor (sdi) was the now well-known cyclin dependent kinase inhibitor p21\textsuperscript{waf} (Noda et al. 1994). This core mechanistic model was validated by the simultaneous demonstrations that the reintroduction of telomerase into a range of human cell types prevented senescence (Bodnar et al. 1998; Wyllie et al. 2000) and that microinjection of blocking antibodies against p21\textsuperscript{waf} (Ma et al. 1999) or its major transcription factor p53 (Gire and Wynford-Thomas, 1998) rescued human fibroblasts from senescence. The observation that senescence in human fibroblasts is associated with activation of the DNA double-strand regulated ataxia-telangiectasia mutated (ATM) signalling pathway leading to the focal accumulation of the repair protein 53BP1 and phosphorylated histone H2AX (γ-H2AX) at telomeres (d’Adda di Fagagna et al. 2003; Gire et al. 2004) extended these studies and raised the possibility that the detection of components of this senescence mechanism could be used specifically to identify senescent cells in vivo.

This approach is attractive in principle. Techniques based on the measurement of telomere length distributions within a population have been used as evidence of cell turnover in vivo and, by inference, the presence of senescent cells in the populations sampled. Using this approach Allsopp et al (1995) demonstrated that human peripheral blood lymphocytes show an average reduction of 2 Kbp of telomere length over 50 years in vivo whereas brain tissue mean telomere length remains constant over the same period. But these methods are complicated by the fact that telomere shortening is not the sole, or even the predominant, mechanism by which mammalian cells enter senescence. Even in humans the telomere-dependent pathway is only used by a subset of cell types. Additional problems of interpretation have emerged as these techniques have grown in popularity.

Sedivy and co-workers (Herbig et al. 2006; Jeyapalan et al. 2007) using a baboon model were among the first to use this approach at the single cell level. Importantly, baboons had recently been shown to display age-associated telomere shortening in leukocytes that closely paralleled that seen in humans (Baerlocher et al. 2003) giving some confidence that telomere-dependent senescence was shared between the two species. An initial study of 30 baboons (age range 5–30 years, 15 males and five females) showed an age-related exponential increase in 53BP foci and γ-H2AX co-localisation with telomeres (so-called telomere dysfunction–induced foci or TIFs). 200–600 fibroblasts were scored from each biopsy with approximately a quarter of the cells positive for these markers, and thus presumably senescent, in the oldest animals. Less than 5% of cells were positive in the youngest members of the cohort.

However, a further study combining the analysis of baboon fibroblasts grown to senescence in vitro with the analysis of skin and muscle biopsies from animals of different ages revealed marked differences in the staining pattern of senescent cells in vitro and presumptively senescent cells in vivo (Jeyapalan et al. 2007). The panel of potential markers used included not only TIF detection but staining for ATM, HIRA, p53, and the cyclin dependent kinase inhibitors p21 and p16. The last was included because it was known to increase markedly in multiple strains of fibroblasts (WI38, IMR-90, MRC5, and NHDF) once senescence was established (Alcorta et al. 1996). HIRA staining was absent from muscle biopsies regardless of donor age but increased from ~20% of fibroblasts in young skin biopsies to over 70% in those derived from 30-year-old baboons. However, the increase in HIRA staining occurred linearly, not exponentially, with age suggesting a distinct aetiology from that giving rise to TIFs. Whilst senescent baboon fibroblasts in vitro were uniformly p21 positive, with multiple γH2AX foci associated with ATM, 53BP1 and telomeres (over 80% of γH2AX foci colocalized with telomeric sequences) only 10% of these cells were p16 positive. But in skin biopsies fibroblasts stained for 53BP1 usually contained only a single focus (70% of 53BP1 positive cells in young animals and 65% in very old animals). In vivo p21 staining in dermal fibroblasts was largely absent but p16 staining was abundant. Such differences between senescent cells in vitro and in vivo should caution against simplistic “if it stains for x it’s a senescent cell” styles of thinking about any specific marker detection assay.

The most intellectually parsimonious hypothesis consistent with the phenotype of senescent baboon fibroblasts in vivo is that researchers are visualising a subset of cells that have been senescent for weeks or months. Stein et al. (1999) reported that whilst p21 progressively accumulates in IMR90 human fibroblasts serially passaged to senescence and is present exclusively in newly senescent cells it subsequently disappears. In contrast p16\textsuperscript{INK4a} is initially absent but increases after IMR90s enter senescence and remains elevated for at least 2 months. Thus, p16\textsuperscript{INK4a} is an excellent potential pathway biomarker for senescent cells in situations where label exclusion or markers of increased cell size (q.v.) cannot readily be used. However some exceptions have been reported (Frescas et al. 2017a).

A particularly important study in this respect is that of Liu et al. (2009) who measured p16 levels in peripheral blood T lymphocytes from human donors (n = 170) aged from 18 to 80. p16\textsuperscript{INK4a} mRNA expression measured using Taqman quantitative RT-PCR showed a highly significant exponential relationship with donor chronological age (Log2[p16] gives R\textsuperscript{2} = 0.4, p < 0.0001 vs donor age). Although a similar relationship was observed at the protein level multiple technical difficulties militated against using this measure. Perhaps predictably p16\textsuperscript{INK4a} levels increased more rapidly with age in smokers compared to non-smokers but provocatively the authors found a blunting of the age-p16 relationship with exercise intensity and duration. The positive association between levels of IL6 (a marker of frailty) and p16 could have indicated an element of "reverse causation" (frail people cannot exercise) however since it is now clear from rodent data that exercise facilitates the clearance of senescent cells by the immune system this can probably be discounted (Schafer et al. 2016).

Even though the measurement of telomere length has been significantly simplified in recent years by the introduction of quantitative-PCR (Montpetit et al. 2014) measurement of p16\textsuperscript{INK4a} message probably offers advantages over this technique for the detection of senescent cells because p16\textsuperscript{INK4a} appears regardless of whether the arrest mechanism is telomere dependent or independent and the message shows a greater...
dynamic range than mean telomere length (~10 fold over 60 years compared to ~2-fold for telomere length over the same period). Nonetheless any quantitative PCR assay remains a relatively costly system to invest in and optimize which begs the question—are there other aspects of the senescent cell phenotype that could be exploited to develop detection systems of equal value but greater ease of use particularly for small scale studies?

The Altered Phenotype Cytological Changes as Markers of Senescence

Senescent cultures of fibroblasts can be clearly distinguished from their growing counterparts by simple light microscopy because the cells are unmistakably bigger. Schneider and Mitus (1976) found that the modal volume of cultured WI38 fibroblasts increased by approximately 40% from early passage to senescence (from 1930 ± 20 μm³ to 2,655 ± 234 μm³). Even larger increases occurred in both RNA (110%) and protein content (~80%). Sherwood et al. (1988) studied IMR90 human fibroblasts using multiparameter flow cytometry and found that mean cell size shifted approximately two-fold between passages 28 and 53. Consistent with earlier work (Cristofalo and Sharf, 1973; Smith and Whitney, 1980) a few such large cells were present in early passage cultures. However senescent cultures were dominated by these large cells whilst their cycling counterparts, even in late-passage cultures, remained relatively small. Similar patterns emerge with other adherent cell types undergoing senescence (e.g., mesenchymal stem cells see Adewoye et al. 2020). Thus, in principle it is possible to identify some types of senescent cells simply by this increase in size coupled with non-invasive markers such as autofluorescence (Poot et al. 1985; Bertolo et al. 2019). But not all cell types show this alteration in cell size at senescence, T cells being the best-known exception (Perillo et al., 1989). Thus, size alone cannot be used as a marker of senescence unless there is prior evidence that hypertrophy accompanies growth arrest.

In senescent fibroblasts this increase in size and protein content is accompanied by a range of ultrastructural changes particularly increased nuclear size, nuclear abnormalities, larger autophagic vacuoles and increased numbers of lysosomes (Lipetz and Cristofalo, 1972; for review see; Stanulis-Praeger, 1987). This latter change is of particular importance for the detection of senescence. Whilst Cristofalo and Kabakjian (1975) were probably the first researchers to report increased activities for the lysosomal enzymes acid phosphatase and β glucuronidase as WI38 cells entered senescence. Campisi and co-workers (Dimri et al., 1995) working with seven different strains of human fibroblasts (including WI-38) modified a histochemical assay for lysosomal β-galactosidase activity to develop probably the most widely used technique for the detection of senescent cells. This colorimetric assay is based on cleavage of the soluble, colourless lactose analogue X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) to form an insoluble blue precipitate (5,5′-dibromo-4,4′-dichloro-indigo). By shifting the pH of the reaction buffer away from the β-galactosidase optimum of pH4 to pH6 the researchers rendered the enzyme less efficient and ensured that only cells with lots of lysosomal β-galactosidase could cleave enough X-gal to generate visible precipitates. Since this effectively limits detectable precipitate to large cells (more formally cells with a high lysosomal mass) which are usually senescent, it is perhaps unsurprising that Dimri et al. (1995) reported a high correlation between cells that were "senescence-associated β-galactosidase" (SAβ-gal) positive and cells that were label excluding on 3H-thymidine long labels.

The original SAβ-gal staining technique is simple and robust, rendering it suitable for use in a variety of settings. Given the range of lysosomal enzymes and substrates available for them this approach is also applicable to a wide range of other enzymes. We found some years ago (Dropova and Faragher, unpublished observations) that when the optimum pH for the enzyme is shifted both dipeptidyl peptidase-4 (CD26) and amino peptidase M show similar “senescence associated” staining patterns to β-galactosidase (unfortunately the coupling chemistry produces a “red on yellow” staining pattern that is harder to score than SAβ-gal). Recently, Hildebrand et al. (2013) showed that the lysosomal enzyme α-fucosidase, is upregulated in cells made senescent through a range of different techniques. Enzyme activity can be visualised using 5-bromo-4-chloro-3-indolyl-α-L-fucopyranoside (X-fuc) in an incubation buffer similar to that for SAβ-gal (but shifted to pH 5.0 not pH6.0). This gives a blue stain similar to X-gal but with the advantage that α-fucosidase induction is somewhat stronger at senescence, particularly in murine cells rendered senescent by drug treatment.

Although the original SAβ-gal histochemical techniques use a paraformaldehyde fixation step a range of fluorescent substrates are also available for β-galactosidase (Kurz et al. 2000; Okamoto et al. 2006; Filho et al. 2018) allowing the visualisation of SAβ-gal in live cells (provided the lysosomes are temporarily alkalinised for example by treatment with bafilomycin A1). One substrate (5-dodecanoylaminofluorescein di-β-D-galactopyranoside) has been successfully used for flow cytometry (Kurz et al. 2000; Noppe et al. 2009) opening up the possibility of flow sorting viable senescent cells. Solution phase variations on these techniques are also available for another style of quantitative read out (Cho and Hwang, 2011).

Users of these techniques should be aware of two potential complications. Firstly, there can be significant delay between entry into senescence and detectable SAβ-gal staining. This was first reported by Thomas et al. (1997) in AGO7086A human mesothelial cells dual stained for Ki67 and SAβ-gal although it was not observed in HUVECs or RPE cells analyzed using identical techniques (Rawes et al., 1997; Kalashnik et al. 2000). Secondly, an increase in the amount of β-galactosidase at senescence is necessary for visualisation. Kurz et al. (2000) using HUVEC showed a 15-fold increase in the amount of enzyme from early passage to senescence; an increase that was paralleled by the increase in lysosomal mass. This does not require an increase in cell size explaining why it is possible, though rarely attempted, to identify senescent T cells by SAβ-gal staining (Gerland et al. 2004; Yang et al. 2018) even though their nucleocytoplasmic ratio is unchanged. However, senescent cells with very short post-mitotic survival times or unaltered lysosome contents will be effectively invisible.
Whilst Stein et al. (1999) had noted that the increase in p16\(^{INK4a}\) roughly parallels the increase in cell size and SAβ-gal in senescing human fibroblasts a more comprehensive temporal analysis was undertaken by Cho and Hwang (2011). Using MCF-7 breast cancer cells rendered senescent by exposure to adriamycin the researchers measured cell cycle-related proteins (including p21 and p53 levels), SAβ-gal activity, cell volume and autofluorescence among other markers. Consistent with earlier studies the levels of p21 and p53 increased within the first 24 h following exposure to the drug whilst cell size increased only for the first 2 days. In contrast, autofluorescence increased 5-fold above baseline levels over 8 days in a roughly linear fashion whilst the number of SAβ-gal positive cells increased in a sigmoidal fashion before stabilising after 6 days. Nonetheless, SAβ-gal activity continued to increase beyond this from an eightfold increase over baseline at day 6 post treatment to a maximum 14-fold increase by day 8. Differences of this scale are in principle detectable simply by immunochemical staining for the enzyme itself although this has rarely been attempted (Joselow et al. 2017).

Thus, although senescent cells from any tissue are potentially detectable by variants of catalytic histochemistry for lysosomal enzyme activity, timing is central to the absolute value of the readout (raising issues of standardisation) and to be visualised cells must 1) have a substantial increase in lysosomal mass irrespective of hypertrophy and 2) have spent sufficient time senescent for enzyme activity to have built up to detectable levels. It should also be borne in mind that any other alterations in cell physiology that meet these criteria will generate a false positive signal. Most famously this can occur in primary fibroblast cultures held confluent or immortal cultures at high cell densities (Severino et al. 2000) and illustrates an important quirk of the routine employment of the technique.

As it stands, the research literature is heavily slanted towards reports of the detection of SAβ-gal in vitro rather than in vivo. This is ironic because the key advantage of the technique is that it gives researchers the capacity to detect senescent cells in tissue samples. When first introduced SAβ-gal was effectively the only assay which allowed senescent cells to be distinguished from their quiescent counterparts in this context. Ironically, in some publications (e.g., Xia et al. 2020) SAβ-gal has been used in vitro to check if a cell type of interest is senescent whilst its senescence in vivo is established using another method (e.g., p16\(^{INK4a}\) staining). It is questionable whether using SAβ-gal staining like this is adding much value to the study.

Unfortunately, reviews of SAβ-gal staining also tend to gloss over the key question of how such stained sections should be scored. This is regrettable, particularly for those using it for the first time because no real consensus has yet emerged around scoring (which raises issues of inter study comparability). A few examples illustrate the range of approaches that have been adopted.

Originally, Dimri et al. (1995) relied on blind scoring of 20 dermal sections from human donors of varying age by a specialist pathologist with the staining frequency presented as a simple scale from minus (indicating no staining) through to “+ + +” (positive cells in all sections of dermis, multiple clusters in all sections of epidermis). Similarly, Paradis et al. (2001) used independent assessment by two pathologists to gauge the localization and number of positive cells in 57 biopsies from normal and abnormal human livers. SAβ-gal staining was classified simply as either “absent” (no or < 10% SAβ-gal positive cells visible) or “present” (> 10% SAβ-gal positive cells visible). Simultaneous review of sections was adopted if the pathologists differed in their opinion.

Kim et al. (2008) took a much more quantitative approach to scoring SAβ-gal in nucleus pulposus chondrocytes within intervertebral disc sections from 25 patients. Here every chondrocyte on the whole section was counted (under ×200 magnification with Nuclear Fast Red counter-stain) and the SAβ-gal positive fraction presented as a percentage. A similarly rigorous histomorphometric approach was used by Gruber et al. (2007) in 57 human disc samples giving an overall incidence of 29.9% (SD ± 24.8, range from 0 to 92%) positive cells.

Berkenkamp et al. (2014) studied the frequency of senescence in mouse renal epithelial cells in vivo using a semi-quantitative approach in which the frequency of SAβ-gal positive cells in 10 random fields of view within representative kidney sections were scored. This was sufficient to yield a statistically significant difference (p < 0.05) between young (3–5 months) or old (18 + month) animals.

In contrast Melk et al. (2003) studying cell senescence in Fischer 344 rat kidneys in vivo quantified SAβ-gal staining simply by imaging kidney sections using Image-Pro Plus Software. A set of slides without the eosin counterstain the group normally employed were photographed, average staining density for the whole section was calculated and the mean staining density of two independent experiments, in arbitrary units, was used as the basis for further calculations. Even this simple approach proved adequate for the demonstration of a significant difference in SAβ-gal staining levels between young (9 months) and old (24 months) rats (0.008 ± 0.003 vs 0.020 ± 0.007 arbitrary units respectively, p < 0.005).

**Metabolic Changes as Markers of Senescence**

Alterations in the amounts and activity of lysosomal enzymes are a subset of the changes that occur in cellular metabolism with senescence. Increased lysosomal size results, at least in part, from dysregulated proteostasis, a key hallmark of ageing. Misfolded proteins are degraded less effectively and form aggregates which, when internalised by lysosomes, contribute to the highly cross-linked and complex materials known collectively as lipofuscin or ceroid (Yin, 1996; Moreno-Garcia et al. 2018). In classical histology these terms are distinct referring to material accumulating within post mitotic or mitotic cell types respectively but colloquially the “ceroid” within senescent cells is often simply called “lipofuscin”. This highly heterogenous “junk” material is responsible for the increased autofluorescence of senescent cells and its presence gives another visualisation option. In essence, rather than staining for the elevated activity of lysosomal breakdown enzymes, simply stain for the “junk” they are trying to break down instead.

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Faragher, Simple Detection of Senescent Cells
This approach can be particularly useful if trying to visualize senescent cells in tissue samples fixed in ways that inactivate lysosomal enzymes (e.g., formalin-fixed paraffin-embedded blocks). As noted earlier immediately following the onset of senescence different markers build up to detectable levels at different rates (Cho et al., 2011) but since the majority of senescent cells in vivo will not usually be immediate entrants this is probably not a serious flaw under normal circumstances. Melk et al. (2003) correlated lipofuscin levels (graded independently by two observers) with SA-β-gal staining in the tubular epithelium of their rat samples and found a highly significant association (p < 0.001) in both young and old animals.

Melk et al. (2003) used a classic histological stain (based on the periodic acid-Schiff reaction) to visualize lipofuscin. Recently Georgakopoulou et al. (2013) used another (Sudan Black B counterstained with Nuclear Fast Red) to co-localise lipofuscin and SA-β-gal in senescent cells. Building on the success of this, but conscious of the difficulties of interpretation of traditional lipofuscin staining, the group undertook the de novo synthesis of a series of Sudan Black B analogues coupled to biotin which they validated in a range of test systems against both proliferation markers (Ki67, BrdU) and SA-β-gal-counterstained with Nuclear Fast Red) to co-localise lipofuscin and a series of Sudan Black B analogues coupled to biotin which they validated in a range of test systems against both proliferation markers (Ki67, BrdU) and SA-β-gal.

This approach allowed the ready visualization of lipofuscin using a peroxidase conjugated anti-biotin antibody and diaminobenzidine (DAB) visualization (although any of a range of anti-biotin detection systems, such as those based on avidin would also probably have worked). In γ-irradiated human fibroblasts in vitro one such analogue (GL13) was used to visualize the kinetics of lipofuscin accumulation giving a timescale broadly similar to that observed by Cho et al. (2011) for autofluorescence. However, all enzyme-based visualisation systems produce staining of variable intensity, so the absolute numbers of senescent cells visualized by this type of lipofuscin detection may differ significantly between researchers depending on simple variables such as the incubation time.

Similarly, Masaldan et al. (2018) building on prior work demonstrating the accumulation of iron within senescing IMR90 fibroblasts and HUVECs (Killelea et al. 2003) studied iron accumulation in mouse embryonic fibroblasts rendered senescent by sublethal γ-irradiation, serial passage or oncogene activation. After 10 days of growth arrest all these types of senescent rodent cells accumulated extremely large amounts of intracellular iron (~15–20 fold compared to growth competent controls). Human fibroblasts and prostate epithelial cells rendered senescent by irradiation or serial passage also accumulated lesser quantities of iron after longer periods of growth arrest (~3.3 to 8.4-fold 21 days post senescence depending upon the cell type and mechanism by which senescence was induced). This accumulation of iron in senescent rodent cells results from the upregulation of the transferrin receptor and a tenfold elevation of ferritin. This in turn was shown to result from impaired lysosome-mediated degradation of this key iron storage protein. Practically, this significant difference in iron handling between the senescent and growing states allowed the group to visualize senescent cells in mouse liver by staining with a rabbit polyclonal antibody against ferritin (in combination with an HRP labelled secondary antibody, DAB visualization and a haematoxylin counterstain). Serial liver sections from young and old animals were stained for SA-β-gal and ferritin respectively and scored by observing four independent fields of view. These findings open up a wide range of potential staining strategies for senescent cells since techniques for the visualization of iron in histological samples are well developed and encompass everything from classical stains such as Perl’s Prussian blue through to quantum dots and can be used in formalin-fixed paraffin-embedded material (Meguro et al., 2007; Duan et al., 2018; van Duijn et al. 2013).

Advanced Glycation End products (AGEs) are often grouped with lipofuscin but unlike the latter can form both extra and intracellularly. They are a heterogeneous group of compounds typically formed by the nonenzymatic (Maillard) reactions of glucose with proteins or lipids. To date limited work has been done to address whether the visualization of AGEs can be used to detect senescent cells but initial studies by Sell et al. (1998) suggest that this should be possible. The authors looked at the accumulation of the AGE pentosidine in strains of reticular and papillary fibroblasts derived from a single donor and passaged to senescence as well as peripheral blood T lymphocytes from 27 donors of varying ages (17–97 years) and states of health. Levels of pentosidine were quantified by HPLC and increased approximately three-fold in both types of fibroblast over the culture replicative lifespan (p < 0.0007) whilst T lymphocytes showed a highly significant increase in pentosidine level with donor age (p < 0.0003). In related work, Kueper et al. (2007) demonstrated that the intermediate filament protein vimentin aggregates in human dermal fibroblasts in vitro due to its modification by pentosidine and carboxymethyllysine (as well as other AGES such as pyrraline and carboxyethyllysine).

More recently Frescas et al. (2017b) conducted a deliberate screen for senescence-specific markers by immunising mice with mouse lung fibroblasts rendered senescent using bleomycin. Human fibroblasts rendered senescent by IR were used as a secondary screen alongside untreated controls. One of the IgMs generated by this approach (9H4) showed approximately a two-fold greater level of staining on senescent cells in this screen. Subsequent Western blotting showed that 9H4 recognises a modified form of vimentin that is presented at the cell surface and then secreted (possibly facilitating the clearance of senescent cells by the innate immune system). The secreted vimentin was modified by a malondialdehyde adduct on cysteine 328. This is provocative because malondialdehyde is an end product of lipid peroxidation that subsequently gives rise to immunologically detectable AGEs (Sajithlal and Gowri Chandrakasan, 1999). Since antibodies to pentosidine and other AGES such as carboxymethyl lysine are commercially available it would seem likely that senescent cells could be identified histochemically using this type of approach.

Vimentin is not the first example of a modified form of a normal protein found at the surface of senescent cells. Porter et al. (1990), Porter et al. (1992) had previously reported the generation of three monoclonal antibodies (SEN-1, SEN-2, and SEN-3) which recognized epitopes on fibronectin only exposed when human fibroblasts become senescent. These antibodies (tragically lost in a laboratory accident shortly thereafter)
communication) could detect senescent human fibroblasts, keratinocytes and mammary epithelial cells in vitro and in vivo in a species-specific manner. If the secretion of modified proteins is a means of signalling for immune clearance then it is likely that there will be significant variations between cell types, species and perhaps individuals as is seen for other components of the Senescence Associated Secretory Phenotype (SASP).

Changes in cellular components as a consequence of dysregulated proteostasis should be conceptually distinguished from those which arise from the transcriptional or post transcriptional regearing that accompanies entry into senescence. These latter changes offer a great many potential markers for the senescent state but can vary widely in the same cell type depending on the inducing stimulus. Nelson et al. (2014) used a microarray-based approach (based on the Affymetric Human Genome U133 Plus 2.0 GeneChip) to compare the transcriptomes of IMR90 fibroblasts rendered senescent either by serial passage or by retroviral infection with H-RAS-V12 (oncogene induced senescence). Compared to proliferation competent controls 5,424 genes were differentially expressed at replicative senescence and 3,188 in OIS. However, there was only moderate overlap between the two states (∼33% of those genes altered by serial passage were altered by OIS but ∼56% of these was also altered by classical replicative senescence). Interestingly although both types of senescence were associated with p16INK4a upregulation the cluster of genes associated with this pathway showed significant differences between the two states (69 out of 118 transcripts commonly downregulated and only 6 out of 31 transcripts commonly upregulated). This has important implications for the use of specific pathway components, as opposed to end points, as pan-specific markers of senescence between different states and tissues.

FUTURE CHALLENGES IN SENESCENT CELL DETECTION

It has been clear for decades that senescence occurs in fibroblast cultures from many mammalian species (Röhme, 1981) but the cell senescence literature remains heavily slanted towards humans and rodents. This is unfortunate because there are several other species which serve as excellent models for human ageing changes or age-related diseases. Given the evidence for the key role played by senescence in ageing the routine detection of senescent cells in these models will become increasingly important but the relevant literature base is currently weak or non-existent.

This is a potential problem because comparative studies of rodent and human fibroblasts show that extrapolation of senescence markers other than label exclusion from one species into another should be done with caution unless there are prior data available. The three models below have been selected to illustrate both the potential gains and the current issues involved in broadening the detection of senescent cells beyond humans and mice.

In the pig, which provides a good model for cardiovascular and intravertebral disc ageing in humans as well as a new model for human progeria (Dorado et al. 2019) there is a small but coherent evidence base for senescent cell detection. Senescent cells can be visualized by the SAβ-gal assay post formaldehyde-fixation (Shi et al. 2018) and serially passaged foetal porcine fibroblasts enter telomere-dependent senescence in a manner that closely resembles that seen in humans. This means that telomere shortening, TIF detection, p16INK4a and p21 WAF can all potentially be used as markers for the senescent state in porcine tissue in vivo (Fukuda et al. 2012; Ji et al. 2012; Donai et al. 2014; Shi et al. 2018). However, Oh et al. (2007) demonstrated that it is possible to produce immortal clones of fibroblasts from both mini-pigs and three-way crossbred meat animals simply by culturing the cells using the 3T3 passage technique. This is not possible using human fibroblasts (McCormick and Maher, 1988) and suggests either that the finding is artefactual or that the molecular pathways regulating entry into senescence are not identical in the two species.

The ageing horse is a particularly valuable model for the types of age-related tendon injuries seen in older humans due to the many similarities between the human Achilles and the equine superficial digital flexor tendon in structure, matrix composition and function. Cultures of human tenocytes from aged and functionally degenerate Achilles show upregulation of p16INK4a protein and a five-fold increase in the SAβGal positive fraction compared to healthy controls (Kohler et al. 2013) a finding that renders the detection of senescent cells in the equine tendon an important goal but studies on senescent cells in equines are limited. Foetal horse kidney cells can be immortalised with SV40 large T antigen (Maeda et al. 2007) indicating p53 and pRb dependent checkpoints and adult equine fibroblasts can be induced to proliferate continuously by the stabilisation of telomere length (Vidale et al. 2012) suggesting equines show telomere dependent senescence. SA-β-gal, p16INK4a protein and γH2AX foci are all detectable in equine chondrocytes following γ irradiation (Copp et al. 2021) suggesting that these would also be useful markers. But as with most other animal models the kinetics of accumulation of these markers post entry into senescence remain unstudied. The type of studies carried out by Cho et al. (2011) are sorely needed.

Selective breeding has created hundreds of dog breeds which vary widely in lifespan presenting an exceptional opportunity to identify pathways associated with ageing. Unfortunately, whilst there is evidence for most of the known hallmarks of ageing in canines (Sándor and Kubinyi, 2019) research on the presence and phenotype of senescent cells in dogs is fragmentary. Whilst breed-specific telomere length has been shown to be a strong predictor of average life span and pathology (Fick et al. 2012) and SA-β-gal has been used to identify senescent canine fibroblasts in vitro there are clearly important differences between human and canine senescence. As with pigs, You et al. (2004) were able to isolate spontaneously immortalized clones from cultures of dog embryonic fibroblasts in vitro. These were shown to have mutations in either p53 or p16INK4a suggesting that these pathways control senescence in canine fibroblasts but appear to be “leaky” compared to humans. A single study suggests that the proinflammatory phenotype of senescent canine dermal fibroblasts may also differ significantly from those of humans.
and rodents (Jimenez et al. 2020). Given the potential value of the model for biogerontology systematic characterisation of canine senescence at the cellular level would clearly be extremely valuable.

Two issues are likely to arise as the detection of senescent cells in different species becomes more commonplace. Firstly, there is a distinct possibility that the relative importance of senescence as an ageing mechanism may differ markedly between them as a result of evolutionary history. Ageing exists as a result of the declining force of natural selection with age and results from two non-exclusive modes of evolutionary gene action. Senescence is thought to have arisen through one of these, antagonistic pleiotropy. This is the selection for alleles or processes that enhance the reproductive success of organisms early in life but which have deleterious effects in the later life course. But, ageing also occurs as a consequence of the inability of natural selection to remove late acting deleterious alleles; a mode of gene action known as mutation accumulation. A wealth of experimental data shows that the relative contributions of antagonistic pleiotropy and mutation accumulation to the evolution of ageing vary widely between species.

One reason for this variation may be that selectively neutral deleterious alleles show frequencies that are influenced by genetic drift and the rate of drift is heavily influenced in turn by effective population size. Unlike most species, humans have undergone both significant genetic bottlenecks and have shifted their survivorship curve from a Type-II population to a Type-I population. This significantly increases drift and thus may well alter the relative importance of cellular senescence as an ageing mechanism between species (Overall and Faragher 2019). Put crudely, humans are evolutionary outliers.

Secondly, different species have very different lifespans which foregrounds the relative extent to which markers of chronological and biological ageing are uncoupled between them. Evidence is emerging from the study of human senescence that some markers better reflect absolute chronological time than cell replication frequency. Maier and colleagues (Waaijer et al. 2012) measured the fraction of senescent cells (via p16INK4a immunostaining) in the dermis and epidermis of a selected sub-group of subjects from the Leiden Longevity Study who were biologically younger than age- and environmentally matched controls (n = 89 per group). The levels of senescent cells correlated closely with the pathological status of the donors whilst age and environment matched controls showed higher levels of senescent cells than the biologically young group. The authors concluded that p16INK4a is a marker of biological time. Further studies using the same experimental design (Waaijer et al. 2016) investigated the relationship between markers of DNA damage (micronuclei, p53BP1 damage foci and telomere associated foci) and health status (n = 40 in each group). There p53BP1 and telomere associated damage foci, but not micronuclei, increased significantly with chronological age but, unlike p16INK4a there was no association between these markers of DNA damage and the health status of the subjects examined. The authors concluded that p53BP1 and telomere associated damage foci effectively measured human chronological age. Studies of this type are likely to be necessary in other species to properly assess the contribution that cell senescence makes to their overall ageing. However, forewarned is at least forearmed and the range of simple detection techniques available suggests that the gaps in our current understanding can be closed much faster in novel systems than was the case when senescence markers were first systematically applied to human systems in the 1970s and 1980s.

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The author confirms being the sole contributor of this work and has approved it for publication.

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Conflict of Interest: The author declares that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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