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Biomarkers to identify and isolate senescent cells

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Highlights

- There is no single biomarker which can robustly identify senescent cells.
- Widely used senescent cell biomarkers should be used in combination for accuracy.
- Technologies like tangential flow filtration can be used to isolate senescent cells.
- Other methods, like senolytic viruses, could be used to remove senescent cells.

Abstract

Aging is the main risk factor for many degenerative diseases and declining health. Senescent cells are part of the underlying mechanism for time-dependent tissue dysfunction. These cells can negatively affect neighbouring cells through an altered secretory phenotype: the senescence-associated secretory phenotype (SASP). The SASP induces senescence in healthy cells, promotes tumour formation and progression, and contributes to other age-related diseases such as atherosclerosis, immune-senescence and neurodegeneration. Removal of senescent cells was recently demonstrated to delay age-related degeneration and extend lifespan.

To better understand cell aging and to reap the benefits of senescent cell removal, it is necessary to have a reliable biomarker to identify these cells. Following an introduction to cellular senescence, we discuss several classes of biomarkers in the context of their utility in identifying and/or removing senescent cells from tissues. Although senescence can be induced by a variety of stimuli, senescent cells share some characteristics that enable their identification both in vitro and in vivo. Nevertheless, it may prove difficult to identify a single biomarker capable of distinguishing senescence in all cell types. Therefore, this will not be a comprehensive review of all senescence biomarkers but rather an outlook on technologies and markers that are most suitable to identify and isolate senescent cells.

Keywords: Aging, senescence, biomarkers, cell biology
1. Introduction

1.1. Aging and cellular senescence

Our society is rapidly aging and the incidence of age-related diseases, such as Alzheimer’s, diabetes and cancer is increasing (Christensen et al., 2009). If these trends continue, aging will become a major economic and social burden (Harper, 2014; Kankeu et al., 2013; Wimo et al., 2013). To avert this impending crisis, we must better understand why we age. Aging is a heterogenic process at both the organismal and cellular level. The number of contributing internal and external factors, such as epigenetic changes (Sinclair and Oberdoerffer, 2009) and the environment, make it difficult to categorize and prioritize the importance of each component. Such diversity has given rise to multiple theories regarding the root cause of aging (Harman, 1956; Park and Yeo, 2013; Wei et al., 2001), which sometimes contradict but more often complement one another. The variety of associated causes implies that aging is likely to be multifactorial in nature (Riess and Krüger, 1999; Sheikh et al., 2013).

One known aging factor is cellular senescence. Senescent cells accumulate with age in organisms, albeit at different rates in the various organs (Erusalimsky and Kurz, 2005; Herbig et al., 2006; Jeyapalan et al., 2007; Paradis et al., 2001). Originally, cellular senescence was defined as a loss of replicative capacity (Hayflick, 1965) caused by a progressive shortening of the tandem repeats protecting chromosome ends (telomeres). This eventually leads to chromosomal damage and replicative arrest (Campisi, 1997). Interestingly, cellular senescence can also be induced by stress (Toussaint et al., 2000) and oncogenes (Bartkova et al., 2006), demonstrating that cellular senescence is not only caused by exhaustion of replicative capacity as first thought. Such heterogeneity of cellular senescence, which we will briefly discuss in the next paragraph, has led a field to sometimes unnecessary ambiguity, leaving researchers to disagree what cellular senescence entails (Burton and Faragher, 2015).

For the purposes of this review, we define cellular senescence as a permanent (under physiological conditions) cell cycle arrest that is a result of cellular stress or damage, including but not limited to abnormal activation of oncogenes, telomere shortening and macromolecule accumulation (De Cecco et al., 2011). With this description we deliberately exclude developmental senescence, quiescent cells, and post-mitotic cells.

[Table 1]
1.2. Senescent cells in health and disease

Cellular senescence is thought to have developed as a safeguard to prevent damaged cells from accumulating and either becoming cancerous or causing cancer (Figure 1). However, accumulation of senescent cells in tissues is detrimental to the animal (Herbig et al., 2006; Jeyapalan et al., 2007) as these non-functional cells directly and indirectly damage surrounding cells (Salama et al., 2014). Examples of such damage include occupying niches required by competent cells to function (Lynch, 2004), secreting transforming, inflammatory and otherwise damaging components of the SASP (Campisi and d’Adda di Fagagna, 2007; Coppé et al., 2010, 2008), promoting tumour formation (Leikam et al., 2015; Zacarias-Fluck et al., 2015) and contributing to various age-related diseases such as atherosclerosis (Irvine et al., 2014; Wang and Bennett, 2012).

While long-term accumulation of senescent cells is harmful to the organism, short-term senescence events prevent cancer (Kuilman et al., 2008), guide development (Muñoz-Espín et al., 2013) and improve tissue repair and wound healing (Demaria et al., 2014; Rodier and Campisi, 2011). One proposed mechanism to negate the long-term detrimental effects of senescent cells, while retaining their short-term beneficial functions, is to periodically purge them from the body. Regular elimination of p16-positive senescent cells from functionally wild-type mice slows time-dependent functional decline and extends median lifespan ~30% (Baker et al., 2016, 2011). Moreover, recent studies have demonstrated clearance of senescent cells from wild-type mice using small molecules which target the BCL-2 protein family (Chang et al., 2016; Zhu et al., 2015), lending credence to this approach as a therapeutic strategy. However, no method currently exists to accomplish this in humans, in large part because senescent cells cannot yet be reliably identified in living tissue. In the following sections, we will discuss biomarkers and their utility in identifying or eliminating senescent cells in a living organism.

1.3. Characteristics of useful biomarkers

A biomarker is a biological signature of a condition which enables one to evaluate if the biological system (organism, cell, etc.) possesses that condition or not. Many molecules, such as proteins, nucleic acids, and lipids, can be used as a biomarker. They can be found within the cell, in the adjacent extracellular area, or even systemically in the circulatory system. Importantly, no single marker currently provides an accurate representation of cellular
senescence.

A useful biomarker must display several important features. First, it should be robustly associated with the condition. Although it is likely that (i) context, such as cell type, will be relevant and that (ii) it may not identify all cases of cellular senescence, it is crucial that the presence of the marker strongly correlates with a specific condition. Second, it is essential to know the threshold at which a marker becomes representative of the specific feature. Most proteins are expressed at basal levels in many cells, and simple evaluation of the presence or absence of the protein is not informative. For discrimination purposes, it is imperative to identify a clear threshold value which defines the cellular status. Finally, to be practical, a marker must be quantifiable using current technologies. Even a comprehensive understanding of a marker is not practically helpful unless we are able to monitor its levels or purify cells positive for it.

In this review, we will emphasize markers of cellular senescence that would be practical to assay. Only single-cell markers of senescence will be discussed. Detailed reviews on systemic aging markers have been published previously (Falandry et al., 2013; Pallis et al., 2014).

1.4. The challenge and benefits of finding robust cellular senescence markers

While cellular senescence has been linked to a number of predictable phenotypic traits and representative biomarkers (Table 1), senescent cells are still a heterogeneous population, and this fact significantly complicates a search for a robust senescence biomarker. Senescent cells arising from different stimuli exhibit measurably distinct proteomes (Aan et al., 2013; Dierick et al., 2002; G. J. Aan, 2011). Moreover, gene expression profiles of senescent cells are cell type-specific (Schnabl et al., 2003). Considering recent breakthroughs in our understanding of a need for personalized medicine, it is likely that variation will be seen not only between different senescent cell types but also between individuals. Importantly, potential reversibility of senescent cells has to be taken into account when evaluating what is and what is not cellular senescence. Although the main consensus in the field is to grant senescence a permanent and irreversible status, a number of studies have shown that cell cycle inhibition can be reversed in human senescent fibroblasts by inhibition of p53 and/or p16, two key pathways used to established cellular senescence (Beauséjour et al., 2003; Ide et al., 1983). More recently, it has been shown that inhibition of p38α/β MAPK lowers p16
levels and restores replicative capacity of aged mouse muscle stem cells (Cosgrove et al., 2014). Unfortunately, these studies did not look into other senescence marker in their models. Therefore, it is still not clear if these aged cells are only forced into the cell cycle from p16-dependant growth arrest or if is it a complete or partial reversal of cellular senescence. Interestingly, cellular senescence can be prevented by activating autophagy, either with rapamycin treatment or Atg7 overexpression, in aged satellite mouse and human muscle cells. However, it is not clear from this data if cellular senescence can be reversed (García-Prat et al., 2016). Even if senescence could not be reversed, it is clear that a large senescence biomarkers can be affected by defined molecules and conditions, which is an important consideration when looking for and validating such markers.

As cancer research has demonstrated, having robust, practical markers can enhance the research field in many ways (Henry and Hayes, 2012; Pallis et al., 2014). A reliable readout that identifies senescent cells would facilitate high-throughput screens as well as allow quantification of aging in various animal models. Most importantly, therapeutic techniques would benefit from robust markers to enable removal of senescent cells from tissues (such as blood filtering, Figure 3), stem cell preparations from aged autologous donors (Melk et al., 2009) and identification of replicative exhaustion in stem cell expansions. One example of how such markers could be used is a strategy using selectively-lytic viruses, which are already used in oncology (Elsedawy and Russell, 2013). These are genetically modified viruses which replicate, lyse and kill cells in the presence (or absence) of specific gene products and, therefore, allow selective targeting. This strategy has successfully reached regulatory approvals in China and the USA for cancer treatment (Garber, 2006; Pol et al., 2015). Theoretically, such a therapy could be applied to senescent cell removal as well.

Currently, due to the heterogeneity of cellular senescence and ambiguity of the term (Table 1), there is no known universal biomarker which can selectively but broadly identify senescent cells in different tissues and extracellular environments. In this review, we will discuss a number of studies which have made significant strides towards the identification and validation of cellular senescence biomarkers.

2. **Surface markers & secretion profile**

Surface and external factors are ideal biomarkers, as they can be detected without intracellular delivery of a probe and without harming the cell. Multiple surface markers have
been associated with senescent cells, some more robust than others. Three major groups of such molecules will be covered: secreted factors, plasma membrane (PM)-associated proteins and PM lipid composition.

2.1. Secreted biomarkers

Secreted factors have excellent potential as biomarkers because detection methods can be contact-free, allowing measurement without disturbing the cells. However, secreted markers are intractable to cell isolation until technology is available which can detect the secretion of single-cells during the selection process.

The senescence-associated secretory phenotype (SASP) has been observed in many different cell types (Acosta et al. 2008; Campisi et al. 2011). One study has defined many of the factors which are secreted by aged human fibroblasts and epithelial cells (Campisi et al., 2008). This list of factors includes but is not limited to inflammatory mediators (such as IL-6 and MIP-3a), growth factors (such as HGF, GRO, and IGF-binding proteins), detached cell surface molecules, and extracellular matrix components. Multiple studies have validated SASP and its related factors as robust markers of senescence, which makes them very promising candidates (Acosta et al., 2013; Sandeman et al., 2001).

Interestingly, another secretory profile of senescent cells recently has been identified. The telomere-associated secretory phenotype (TASP) has been associated with replicative senescence (Braig et al., 2014; Jiang et al., 2008). It is not clear if this phenotype should be classified as separate from the SASP phenotype. However, it does provide some new and unique markers to consider. These include cathelin-related antimicrobial peptide (CRAMP), chitinase, stathmin, and EF-1α. Interestingly, one study has shown that CRAMP and chitinase levels highly correlate not only with replicative senescence but also with human aging in general (Lu et al., 2014).

These secretory profiles are useful tools for evaluating general tissue or cell culture senescence. Some recent publications covered various ways of detecting such phenotypes. One such method is the SASP-responsive alkaline phosphatase (SASP-RAP) assay, where activity of secreted alkaline phosphatase is measured by a commercial chemiluminescence kit. SASP-RAP reliably detected senescence in rat renal tubular epithelial cells and mouse embryonic fibroblasts treated with etoposide, a DNA-damaging agent (Gu and Kitamura, 2012). ELISA protocols have also been developed for an array of SASP components,
including IL-6 (Rodier, 2013). Moreover, it also may be possible to detect SASP by monitoring the known effects on surrounding cells. For example, senescent fibroblasts affect epithelial cell proliferation which can, in turn, be quantified as an indirect marker (Parrinello et al., 2005).

Nevertheless, it is important to understand that there are some major limitations inherent to secretory markers. Firstly, the SASP may vary between cell types and between different stages of senescence (Coppé et al., 2010; Maciel-Barón et al., 2016; Rodier et al., 2009). Secondly, current technology does not permit the rapid, high-throughput analysis of single-cell secretory phenotypes necessary to isolate a population on this basis. Although this technological gap currently prevents the use of a secreted marker for selective cell removal, secreted biomarkers are still applicable for general identification of senescent cells and potentially for diagnostic tests aiming to quantify the burden of cellular senescence in an individual.

2.2. Plasma membrane-associated proteins

Another set of surface structures which could be used for identifying and isolating senescent cells are plasma membrane-associated proteins, such as receptors and glycoproteins. As these structures are the main means of interaction between extracellular and intracellular compartments, their composition and functionality is essential for cell homeostasis.

Several independent research groups have shown that plasma membrane (PM) glycoprotein composition is altered in aged cells (Poot et al., 1986; Wu et al., 2009). Glycoproteins are important components and play a role in many functions, including signalling in immune system and hormone responses (Ervasti, 2000; Li et al., 2010). There is a wide range of possible glycoprotein variants, and specific glycoprotein compositions have been linked to a senescence profile. For example, senescent human fibroblasts exhibit higher levels of concanavalin A, fucose, and glucosamine (Blondal et al., 1985).

Expression changes in the concentration of PM-associated proteins could also be used as a senescence biomarker. Althubiti and colleagues have identified 107 PM-associated proteins upregulated in a human bladder cancer cell line induced to senesce by ectopic expression of p16 and p21. Ten of these upregulation events were validated by western blots and immunocytochemistry (Althubiti et al., 2014), while several additional candidates like
ICAM-1 had previously been reported to increase in senescent cells (Schnabl et al., 2003).

Several standard methods exist to measure expression changes in PM proteins, including immunocytochemistry (Obradovic and Jurisic, 2012) and techniques utilizing other protein-specific probes (Liu et al., 2005). Changes in the expression of PM proteins can also be detected with transductionally targeted recombinant viruses, which are able to enter only those cells with the appropriate ligand exposed on the PM. This has been accomplished in cancer cells already and readers are referred to an excellent review by Everts and Curiel for more information (Everts and Curiel, 2004). A similar strategy could be used to target senescent cells. The specific targeting of senescent cells holds promise both as a senescent assay and as a potential therapeutic. Transductionally targeted viruses could be engineered to selectively kill senescent cells or to carry a fluorescence reporter gene for identification (Perez et al., 2013) (Figure 4).

2.3. Plasma membrane lipid composition

The third type of marker which can be used to identify and isolate senescence cells is linked to PM lipid composition (Sud et al., 2007). Although PM lipid-based senescence biomarkers are less established and harder to detect compared to other surface signatures, some promising candidates and techniques have emerged which hold great potential for senescent cell identification and isolation.

Membrane lipid composition and membrane biophysical properties change in senescent cells. Plasma membranes, due to altered lipid composition, become more rigid with age (Fulop et al., 2012a; Momchilova et al., 2014). Moreover, senescent cells increase in size compared to non-senescent analogues (Kim et al., 2015; Rodier and Campisi, 2011). This modifies many essential functions within the cell, such as diffusion, cell size, membrane fusion, chemical and electrical processes, cell elasticity and membrane stiffness (Pontes et al., 2013). Such cellular mechanics have already been used as biomarkers to identify cancer cells, malaria-infected cells and primary cells (Darling et al., 2008; Suresh et al., 2005).

A number of technologies have emerged to detect PM biomechanical and lipid composition differences between cells. For example, tangential flow filtration (Cai et al., 2015) discriminates between cells based on membrane stiffness. Such a method has been successfully used to separate differentiated cells from pluripotent ones (Willoughby et al., 2016). Moreover, a technology which selects cells based on cell mechanics has been used to sort red blood cells based on shape and deformation (Beech et al., 2012), which make this
sub-group of senescence markers even more viable.

Another structure which could be used as a marker is a lipid raft (LR). LRs are small, dynamic structures in the plasma membrane that are high in cholesterol and low in unsaturated phospholipids (Brown, 2006). These structures recruit signalling components to the plasma membrane (Kabouridis, 2006). It has been shown that T cells from young subjects have a different LR distribution and functionality compared to T cells from older individuals, although it is still not clear if these changes are linked to senescent or aged cells in general (Fulop et al., 2012b; Larbi et al., 2006, 2004).

LR dispersal can be measured using CTxB fluorescent molecules that bind to GM1 gangliosides, which are LR markers (Holleran, 2003). Moreover, it is possible to use specific fluorescent LR probes, such that quantifying the number of probes bound could be used as a means of identifying senescent cells (Mikhalyov and Samsonov, 2011). For a full analysis of the most current methods for studying membrane microdomains (including LRs and the signalosome) in intact cells, readers are advised to refer to Lagerholm’s review (Lagerholm et al., 2005).

In summary, membrane lipid composition and its effects on cell biophysical properties represent a promising line of exploration for developing reliable, quantitative senescence markers.

3. Intracellular senescence biomarkers

Many intrinsic changes have been observed in senescent cells. While currently known intrinsic changes are more discrete and better defined compared to surface ones, they are harder to measure in live tissue. Three major groups of potential biomarkers will be covered: DNA-related markers, protective mechanism or damage signatures, and cell cycle genes.

3.1. DNA-related senescence biomarkers

It is still not clear if mutations significantly contribute to aging, but the cumulative number of mutations and the rate at which they accumulate increases with age in model organisms and senescent cells (Kennedy et al., 2012; Sedelnikova et al., 2004). Moreover, accelerated-aging diseases, such as Werner and Cockayne Syndromes, are linked to
malfunctioning DNA repair mechanisms (Rossi et al., 2010). Such findings make it tempting to speculate that DNA quality, metabolism, and maintenance are key mechanisms in the biology of aging. That being said, most of the changes which will be discussed are not exclusive to senescent cells and therefore use as a sole marker would yield a significant proportion of false positives.

Various markers for DNA damage, such as γH2AX, 53BPI foci, Rad17, ATM and MDC1, are commonly used as conditional markers of cellular senescence (Lawless et al. 2010; Wang et al. 2009; Sharma et al. 2012). γH2AX immunocytochemistry has been shown to effectively mark senescent fibroblasts (von Zglinicki et al., 2005). Although these markers are well established, they do not represent cellular senescence directly. γH2AX is a marker for DNA double-strand breaks (DSBs), which is neither necessary for nor exclusive to cellular senescence (Chapman et al., 2012). Similar limitations exist for other DNA damage related biomarkers (Awasthi et al., 2015).

DNA synthesis markers can also serve as negative indicators for cellular senescence. DNA synthesis rate can be analyzed in 5-bromo-2’-deoxyuridine (BrdU) or 5-ethynyl-2’-deoxyuridine (EdU) incorporation assays (Gratzner, 1982; Salic and Mitchison, 2008). Only cells that are undergoing DNA synthesis are able to incorporate transiently available BrdU or EdU nucleosides. The incorporation rate of such synthetic nucleosides is low in senescent cells, which suggests that DNA replication is near-absent (Voutetakis et al., 2015). Another assay for cell proliferation is based on immunostaining for marker Ki-67 (Urruticoechea et al., 2005). However, assays based on DNA synthesis would also identify other non-replicative cells, such as quiescent cells and post-mitotic cells, which make such markers sub-optimal if used alone.

Telomere length can also be used as a biomarker for replicative senescence (Mather et al. 2011; Zietzer & Hillmeister 2014; Bekaert et al. 2005.). However, the shortening of telomeres associated with replicative senescence has yet to be translated into a common assay. There are several ways to measure telomere status in the cell. Measuring telomere length by quantitative PCR is one way (Cawthon 2002). Moreover, protocols have been designed which allow measurement of absolute telomere length (O’Callaghan and Fenech, 2011) or telomere length of a single-cell (Wang et al., 2013). Unfortunately, these PCR-based approaches to telomere measurement require destruction of the cell. Fluorescence in situ hybridization (FISH) can be used to visualize the length of telomeres in fixed but intact cells (O’Sullivan et al., 2005; Ourliac-Garnier and Londoño-Vallejo, 2011), which retains
information such as subcellular localization. Moreover, FISH can be combined with flow cytometry to provide information on telomere length in thousands of cells (Baerlocher et al., 2006; Hultdin et al., 1998).

In addition to the direct measurement of telomere length, another potential aging marker is telomere dysfunction-induced foci (TIF) (Brugat et al., 2010; Sahin et al., 2011). TIF is a term used to describe telomeres which accumulated various DNA damage factors, such as γH2AX, ATM and Mre11 (Badie et al., 2010; Takai et al., 2003). TIF has been shown to increase in baboon fibroblasts that have undergone replicative senescence (Jeyapalan et al., 2007) and also has an effect on the metabolic status of the cell (Sahin et al., 2011). Therefore, it may be possible to identify senescent cells by immunostaining for these factors at telomeres.

Epigenetic changes, including senescence-associated heterochromatic foci (SAHF), can also be used as biomarkers for cellular senescence. Epigenetics largely determine cell-state differences in a genetically identical population, such as within an organism (Sinclair and Oberdoerffer, 2009). Although epigenetic markers vary depending on the environmental context and type of cell (Kosar et al., 2011), epigenetic profiles have been defined for various conditions, including cancer (Dumitrescu, 2012; Mäbert et al., 2014) and aged human cells (Horvath, 2013). A specific heterochromatin profile has been linked to cellular senescence (Kosar et al., 2011; Narita et al., 2003). SAHFs, domains of heterochromatin that contribute to silencing of proliferative genes in senescent cells (Narita, 2007), can be detected by multiple methods, including DAPI staining and immunocytochemistry against SAHF components, such as HP1 (Aird and Zhang, 2013). Another novel assay, called chromatin in vivo assay (CiA), has the potential to be used in high-throughput screening for heterochromatin alterations (Jones, 2012). It may be possible to adapt it to report SAHFs.

In summary, DNA-related signatures are some of the most prominent cellular senescence biomarkers. In addition to the fact that one or more of these markers can robustly be found in most senescent cells, the broad availability of methods for their detection makes this biomarker group extremely potent. Further discussion of how they can be used in combination with other markers to increase specificity will be discussed later in the review.

3.2. Protective mechanism and damage markers

Aging is closely linked to loss of damage repair and/or stress response capabilities, which results in the accumulation of various toxic by-products and other macromolecules (Chen et
al., 2007; Hipkiss, 2006). As a result, enzymatic by-products and activities of various lysosomal and proteosomal enzymes, which are responsible for aggregate clearance, are often altered in aged cells (García-Prat et al., 2016). The ability to detect such changes would help to identify and possibly isolate senescence cells.

Perhaps the most frequently used biomarker for senescent cells is increased activity of β-galactosidase (β-gal) (Bassaneze et al., 2008; Dimri et al., 1995), which reflects increased lysosomal mass (Kurz et al., 2000). Several techniques exist for β-gal detection, including fluorescence-based and cytochemical methods (Debacq-Chainiaux et al., 2009). However, β-gal activity also increases in quiescence and in response to various forms of stress (Yang and Hu, 2005). Thus, β-gal as a sole marker may often yield false-positives and is best used in combination with other markers. A similar marker, α-fucosidase, has recently been reported (Hildebrand et al., 2013). While this marker boasts similar ease of detection, it remains uncertain if it is equally, more, or less specific for senescence than β-gal.

Proteasomes, protein complexes responsible for degrading unneeded or damaged proteins, may also act as useful biomarkers for senescence. For example, 26 S proteasome activity is reduced in human primary senescent fibroblasts (Chondrogianni et al., 2003; Reinheckel et al., 1998; Torres and Perez, 2008), and 26 S proteasome activity has recently been measured using fluorogenic peptide substrates (Georgila et al., 2014). While this assay has not yet been applied to senescent cells, it may serve as a secondary marker in conjunction with other indicators of cellular senescence.

Reactive oxygen species (ROS) can also be used for senescent cell detection. ROS are involved in oxidative stress, signalling and differentiation processes. ROS increase with age and correlate with many age-related cellular changes (Liochev, 2013). Furthermore, elements associated with protection against oxidative stress (Cox2, SOD, Mn and other antioxidants) often show reduced functionality in aged organisms (Espinoza et al., 2008; Paul et al., 2007).

However, oxidative stress, like many other markers mentioned in this review, is only an indirect marker of senescence. It is not necessary for cells to have high ROS activity to become senescent. Furthermore, a low level of ROS, is not only useful but essential for normal cellular functions (Liochev, 2013). Conversely, high ROS levels are not always associated with senescence as they are sometimes linked to differentiation (Paul et al., 2014). This may make interpretation of results problematic if clear thresholds (and their deviation depending on environment) are not determined. Various assays, including commercially available kits, exist to measure oxidative stress markers and levels of intracellular ROS.
Most of these techniques rely on fluorescence detection with either flow cytometry/fluorescence microscopy or chemiluminescence. In summary, markers of cellular damage and protection mechanisms, such as ROS and β-Gal, are well-known signatures of cellular senescence. While not exclusive to cellular senescence, these relatively easy-to-assay markers are frequently employed as companion diagnostics.

### 3.3. Cell cycle genes

Arrest of cell proliferation is a hallmark of a senescent cell, which means that genes responsible for such regulation can potentially be used as markers to identify which cells are senescent.

During growth arrest, various cell cycle inhibitors such as p53 and p21 are activated (Campisi, 2005). p21\textsuperscript{waf} is a cyclin-dependant kinase (CDK) inhibitor which acts by mediating pRB dephosphorylation. Its overexpression promotes cellular senescence as shown by β-gal activity, telomere length and DNA damage monitoring (Huang et al., 2004). Moreover, if p21\textsuperscript{waf} is down-regulated in senescent cells, replicative capacity is restored (Schnabl et al., 2003).

Another molecular alteration that is specific to senescent cells is the phosphorylation (and activation) of p53 at serine 15. This phosphorylation has been observed in normal human fibroblasts that have undergone replicative senescence (Webley et al., 2000), oncogene-induced senescence (Ferbeyre et al., 2000), and other forms of senescence (Ghosh et al., 2008). p53 and the related tumor suppressor p16\textsuperscript{INK4a} are relatively robust markers of senescence as their expression is required to induce the characteristic permanent cell cycle arrest (Campisi and d’Adda di Fagagna, 2007; Krishnamurthy et al., 2004; Vandenberk et al., 2011). However, these genes are not absolute markers of cellular senescence. For example, p53 is also responsible for apoptotic programmed cell death (Amaral et al., 2010). An assay relying solely on p53 levels, therefore, would not be able to discriminate senescent cells from those undergoing apoptosis. Furthermore, mutations may shift p53 activity patterns (Muller and Vousden, 2013). In such cases, p53 mutations, although tumorigenic, may not be linked to cellular senescence. Moreover, a reversible growth arrest state, known as quiescence, also depends on the aforementioned cell cycle inhibitors (Li and Bhatia, 2011). Although such context specificity would make these biomarkers unreliable if used alone, their use in a
4. Summary of most promising senescence biomarkers and combinatory approach

4.1. Synopsis of the most practical markers for identification and isolation of senescent cells.

Some of the cellular senescence biomarkers are more practically viable either because of their relative robustness to accurately select senescent cells or existing technologies for their detection. In the following paragraph, we offer our opinion of the most promising senescence biomarkers.

Firstly, markers for cell cycle genes (p16, p21, p53, Rb) and DNA synthesis (BrdU incorporation) provide a robust way to distinguish dividing cells from non-dividing cells. Considering that cell cycle arrest is a hallmark of cellular senescence, such markers carry great potential. That being said, if used alone, these assays would also identify non-senescent, non-proliferative cells, such as post-mitotic neurons or quiescent stem cells. Other senescence markers should be used in combination to alleviate such off-target effects.

Secondly, due to the non-proliferative and dysfunctional nature of senescent cells, an accumulation of various intracellular junk and a stressed cellular repair mechanism are often observed. This makes a number of pathways a potential source for biomarkers, the most prominent example being β-gal activity in lysosomes. With a number of assays now available to detect such changes, this category of biomarkers may offer several strong candidates for diagnostic assays.

Thirdly, considering recent developments in the ability to quantify cell biomechanical properties (Beech et al., 2012; Darling et al., 2008; Suresh, 2007) and plasma membrane elasticity alterations observed in senescent cells, new technologies could arise that would enable senescent cell identification and isolation. Similar approaches, based on tangential flow filtration, for example, have been used successfully to separate pluripotent cells from heterogeneous populations (Willoughby et al., 2016).

Lastly, senescent cell surface components, such as ICAM-1 and NOTCH3, are very attractive biomarkers due to possibility of screening live cells without damaging them. Moreover, a number of techniques are available to achieve such aims, including immunocytochemistry, transductionally targeted viruses, FACS or magnetic cell sorting. Recent studies have also identified a large number of novel surface markers, which potentially may join the list of commonly used signatures for senescence (Althubiti et al.,
4.2. Using combinations of markers for a more robust and specific assay

To achieve the highest possible specificity of an assay, a combination of the most prominent markers should be used. To date, it is common best practice in the cellular senescence research field to use a panel of markers to validate that cells are senescent (Acosta et al., 2013; Guo et al., 2009). p53, p16, β-gal and telomere length remain the most commonly used markers and their combination should accurately pinpoint senescent cells. Although essential, a four-piece requirement for identification and isolation is practically challenging and cost- and time-ineffective. We propose that a synthetic reporter (or lytic) virus could potentially be used as a single system with combinatorial potential.

Similar strategies have been successfully used in the past for oncolytic viruses where input from multiple sources (genes, receptors etc.) was required for the function of the virus (Barker et al., 2003; Larson et al., 2015; Lee et al., 2010; Singh et al., 2012). For example, one study constructed an adenovirus in which gene E1a was under the control of the hTERT (human telomerase reverse transcriptase) promoter while the viral gene E1b was under the control of the HRE (hypoxia response element) promoter (Wang et al., 2008). Such a system allowed viral replication only in hypoxic cells with active telomerase, which are very common signatures for cancer cells. Although this particular example utilized two different promoters, much more elaborate bio-sensor circuits could be built to allow even greater specificity. Theoretically, most transcriptional signatures can be targeted with these viruses by genetically modifying viral promoters. Moreover, viral coat engineering allows specifying to which surface markers a viral particle can bind and consequently infect a target cell (Verheije and Rottier, 2012).

5. Outlook

In this review, we have covered a number of biological signatures, outlining their advantages and disadvantages as cellular senescence biomarkers. Some of them are widely accepted to mark an aspect of aging (telomeres, ROS etc.), while other molecules, such as lipid rafts, still require further validation as a practical marker.

One of the major limitations of all these markers is the fact that the change is only
relative. For example, ROS is always present in the cells and small amounts are essential for intrinsic signalling. Although an increase in ROS levels is considered to be a marker of oxidative stress and possibly cellular senescence, at what point does the difference become a biomarker? Furthermore, can we robustly classify a measurement that can be considered a definitive indicator of senescence in all circumstances? These questions remain to be answered. Another important aspect is use of biomarker combinations. None of the markers can accurately and robustly detect senescent cells in all given instances. This is why it may be useful to use a combination of markers. Such combinations of complementing markers, such as the ones we have suggested, would offer a diagnostic with better accuracy and robustness.

Although senescent cell markers are widely used in basic research, such signatures may also be useful in the clinic as well. Removal of senescent cells has been shown to have a positive effect on murine health. With increasing knowledge in immuno- and viro-therapies, new opportunities may arise to use senescence biomarkers in senescent cell removal procedures for therapeutic benefit.

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Figure 1 | Senescent cell role in health and disease. (i) Senescent cells accumulate in aged primates (Herbig et al., 2006; Oishi et al., 2014), and (ii) their removal might be impaired due to declining efficiency of the immune system. These accumulating senescent cells can (iii) damage surrounding cells (Biran and Krizhanovsky, 2015) and cause (iv) cancer (Leikam et al., 2015) and (v) other age-related diseases, such as atherosclerosis (Erusalimsky and Kurz, 2005; Fischer et al., 2013; Nishimatsu et al., 2015). Although senescent cells play an important role in (vi) tumor suppression (Tominaga, 2015), (vii) wound healing (Demaria et al., 2014) and development, (viii) periodically removing them does not produce significant side-effects (Baker et al., 2016). Furthermore, (ix) removing senescent cells has been shown to increase health- and life-span in mouse models (Baker et al., 2016, 2011; Chang et al., 2016; Yi Zhu et al., 2015)
Figure 2 | Comparison of replicative senescence (RS) versus stress-induced premature senescence (SIPS). These two different types of senescence have measurably distinct proteomes (text in blue) but share many major senescence phenotypes (text in red). Moreover, it has been shown that different factors are responsible for induction of these phenotypes (Aan et al., 2013; Braig et al., 2014; Chondrogianni and Gonos, 2004; De Cecco et al., 2011; Golde and Miller, 2009).
Figure 3| Schematic for the isolation of senescent cells from blood. This schematic demonstrates the principle of isolating senescent cells from the blood using a dialysis approach. This could be applied to humans to clear the blood of senescent cells, which are known to accelerate aging and cancer in neighbouring cells and to interfere with tissue homeostasis.

Figure 4| Schematic representation of combinatorial senescent cell targeting with a synthetic reporter virus. This is an example of how a virus could be designed to select senescent cells based on a combination of markers. Viruses can be engineered to enter only cells which have a specific surface marker (transductional targeting). Moreover, simple (as demonstrated in the graph) or more elaborate synthetic genetic circuits could be used to allow reporter expression to occur only in cells possessing multiple markers for senescence, such as an active p16 promoter or an increased SA-β-galactosidase activity. Replacing fluorescent proteins with suicide genes, which are functional only when expressed together, would allow generation of senescent-cell-specific lytic viruses. Such a senescent cell removal therapy could be used to remove senescent cells in an effort to increase health- and life-span of patients (Baker et al., 2016, 2011; Chang et al., 2016)
<table>
<thead>
<tr>
<th>Senescent cell phenotypic traits</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permanent cell cycle arrest</td>
<td>Shay et al., 1991</td>
</tr>
<tr>
<td>Persistent DNA damage response (DDR)</td>
<td>Fumagalli et al., 2012</td>
</tr>
<tr>
<td>Senescence-associated heterochromatic foci (SAHFs) and other epigenetic changes</td>
<td>Narita et al., 2003</td>
</tr>
<tr>
<td>Senescence associated secretory phenotype (SASP)</td>
<td>Campisi, 2005</td>
</tr>
<tr>
<td>Altered metabolism including increased lysosomal and proteosomal activity</td>
<td>García-Prat et al., 2016</td>
</tr>
<tr>
<td>Marker category</td>
<td>Senescence marker</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
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<tr>
<td>Secretion and surface</td>
<td>SASP and TASP</td>
</tr>
<tr>
<td></td>
<td>Plasma membrane-associated proteins (concanavalin A, fucose, ICAM-1, DEP1, B2MG, NOTCH3, DCR2)</td>
</tr>
<tr>
<td></td>
<td>Plasma membrane mechanics (solidity, cell size)</td>
</tr>
<tr>
<td></td>
<td>Lipid rafts (LR)</td>
</tr>
<tr>
<td>DNA-associated</td>
<td>DNA damage markers (γH2AX, S3BPI, Rad17, ATR, ATM, MDC1, TIF)</td>
</tr>
<tr>
<td></td>
<td>DNA synthesis</td>
</tr>
<tr>
<td></td>
<td>Telomere length</td>
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<tr>
<td></td>
<td>SAHF</td>
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<tr>
<td>Damage and repair mechanism</td>
<td>Proteasome activity</td>
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<tr>
<td></td>
<td>Lysosomal activity (β-Galactosidase, α-Fucosidase)</td>
</tr>
<tr>
<td></td>
<td>ROS</td>
</tr>
<tr>
<td>Cell-cycle related</td>
<td>p16-pRB axis</td>
</tr>
<tr>
<td></td>
<td>p53-p21 axis</td>
</tr>
<tr>
<td></td>
<td>DEC1 (BHLHB2)</td>
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<td></td>
<td>PPP1A</td>
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</tbody>
</table>
Table 3: A list of techniques which could be applied to remove senescent cells

<table>
<thead>
<tr>
<th>Technique</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood perfusion using surface markers or cell stiffness</td>
<td>Variations of the technique already used in clinic</td>
<td>Only able to remove senescent cells circulating in blood system</td>
</tr>
<tr>
<td>Magnetic sorting using intracellular marker</td>
<td>Established technology exists</td>
<td>No clinical grade MAC sorter is currently available</td>
</tr>
<tr>
<td>Lytic virus</td>
<td>Variations of the technique already used in clinic</td>
<td>Safety concerns of using virotherapy</td>
</tr>
<tr>
<td>Stimulating the immune system</td>
<td>Success and promise in oncotherapy</td>
<td>Lack of robust and unique cell-surface marker for senescent cells</td>
</tr>
<tr>
<td>Chemical compounds</td>
<td>Applicable for therapeutic interventions</td>
<td>Potentially high off-target effects</td>
</tr>
</tbody>
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