Cellular Senescence by the Epigenetic Regulators Inhibitor of Growth

Thanakorn Pungsrinont and Aria Baniahmad

Keywords: ING; Senescence; Cancer; Epigenetics; Aging; PHD; Evolution

Abstract

The epigenetic regulatory tumor suppressor, INHibitor of Growth 1 (ING1), obtained more focus since it has been suggested as one of the aging-related candidate genes among healthy elderly individuals. ING1 belongs to the ING family proteins characterized by a plant homeodomain (PHD), which is important for recognizing and binding to histone marks, thus allowing ING to regulate genes expression through histone modification and chromatin changes. Interestingly, the PHD of ING proteins is highly conserved among species between mammals, insects and plants. The ING factors regulate the program of cellular senescence and DNA repair, which are suggested to have a protective role in inhibiting cancer cells proliferation. Here, we provide an insight into the functional role of ING factors in development and tumor cells.

Introduction

Epigenetic and genetic factors are suggested to be involved in the aging process [1-3]. Indeed, aging research on various model organisms like Caenorhabditis elegans or Drosophila melanogaster improved our understanding of genomic, epigenetic and proteomic aspects regarding the lifespan of these organisms [2-5]. Specific set of genes or genetic loci that are related to longevity and aging are being analyzed in these model systems. Genetic and epigenetic factors appear to have significant influence also on human longevity, since the heritability of human lifespan was estimated in a range of 20-30% in many studies [6-10]. Unlike genetics, epigenetics refers to "functional changes of the genome without changing the DNA sequence". This includes chromatin changes and remodeling, which in general is triggered by factors that promote or remove histone modifications and regulate exchange of histone variants [11,12]. However, the underlying mechanisms linking epigenetics to aging are poorly understood. One reason is the fact that aging is associated with a variety of human disorders, which includes cancer [13].

Interestingly, the gene encoding the epigenetic regulator tumor suppressor, inhibitor of growth 1 (ING1) has been suggested to be one of the aging-related candidate genes among 47 healthy individuals at the age of 85 years or older [14]. Within this cohort, no aging-related diseases such as cancer, cardiovascular disease, pulmonary disease, diabetes, or Alzheimer disease have been diagnosed.

The ING tumor suppressors are localized in the nucleus and directly associated with chromatin regulation and control of gene expression [15,16]. ING factors control various cellular pathways which include cell cycle control, DNA repair and two tumor protective pathways: apoptosis and cellular senescence that both seems to be important pathways for tumor suppression.

In general, the Hayflick limit suggests a limited cell division potential of primary cells that is also termed replicative lifespan [17]. Cells having approached the replicative lifespan are metabolically active, exhibit a changed morphology, and are also termed to be cellular senescent. Cellular senescence occurs naturally in vivo during normal development and is involved in embryonic patterning [18]. Also during tumorigensesis, the pre-malignant tumors exhibit high levels of senescent cells in vivo [19]. During further steps of tumor progression, the level of senescent cells declines, suggesting that malignant tumor cells evade from the cellular senescence pathway and from the other anti-tumor pathway, apoptosis. It is therefore suggested that during tumorigensesis, tumor cells escape from the cellular senescence pathway and undergo selection to evolve and develop into malignant cancer.

Interestingly, cellular senescence is a cellular pathway that is characterized by an irreversible cell cycle arrest that is mostly induced by replicative lifespan or cellular stress and therefore is suggested to act against cancer malignancy [19,20]. However, cellular senescence can be either detrimental or beneficial, depending on the physiological context and situation. Cellular senescence can disrupt normal tissue structures and functions, but on the other hand, cellular senescence is an effective mechanism to suppress cancer cells proliferation [19-21]. Notably, in primary human cells the ectopic expression of either ING1 or ING2 alone is sufficient to inhibit cell proliferation by inducing cellular senescence as an underlying mechanism [22,23].

Among the ING family members, the human ING1 and ING2 are two closely related proteins that share high identity and homology in amino acid sequence [24], indicating that they exhibit similar tumor suppressive functions. The most widely expressed isoform of ING1 and ING2 are ING1b and ING2a, respectively. Most studies support their role as tumor suppressors as their expression is often found to be decreased or lost in many human tumors [25-29]. The loss or...
reduction of ING1 and/or ING2 expression could be the result of misregulation of transcription factors or gene inactivation mechanisms, and since their loss often occurs at an early stage of tumor development [30-32], it suggests that decreasing ING1 or ING2 expression in pre-malignant tumors contributes at an early stage to malignant tumorigenesis.

To analyze their functional role in tumor cells, many overexpression studies revealed that either ING1 or ING2 expression result in tumor growth inhibition [22,23,33-34] and accordingly knockdown (KD) or knockout (KO) result in an enhanced tumor proliferation [33,35-37]. These studies provide an insight to understand the functions of ING1 and ING2 to regulate cell growth. The molecular pathways are discussed below.

**ING proteins as epigenetic regulatory tumor suppressors induce cellular senescence through their PHD**

A tumor suppressor gene encodes for a protein that suppresses tumor growth. The ING family of genes and the corresponding encoded proteins were originally identified in 1996 [33], and later characterized as candidates for tumor suppressors because they are involved in many processes such as cell growth, apoptosis, cellular senescence, migration, and DNA repair [32,38]. The ING proteins are characterized by a well-conserved carboxyl-terminal region that contains a plant homeodomain (PHD) [24]. The histone binding and modification is an interesting ability of ING proteins. The PHD selectively binds preferentially to trimethylated lysine 4 of the histone 3 (H3K4me3), which is present in nucleosomes of transcriptional active genes at promoters and downstream of transcription start sites [39,40], thus linking the ING proteins to epigenetic regulation [16].

Although the PHD domain of both ING1 and ING2 binds to the activating mark of the histone modification (H3K4me3), it was a surprise that ING1 and ING2 interact with a histone deacetylase (HDAC) complex. The amino-terminus of ING1 and ING2 directly interacts with the mSin3a/HDAC1-2/SAP30 complex that leads to gene silencing [16,39,41]. This suggests that ING tumor suppressors, if recruited, may counteract and inhibit some active gene loci. For example, it has been described that H3K4me3 is required for ING2 binding at the cyclin D1 promoter and that cyclin D1 expression is transcriptionally inactivated by the mSin3a/HDAC1 complex [42]. Cyclin D1 expression is controlled by E2F factors that are regulated by the retinoblastoma protein (pRb). Thus, these findings link the ING protein to the induction of cellular senescence through inactivation of pRb by reducing the activity of cyclin D1-cyclin-dependent kinase 4 (CDK4) complex (Figure 1).

Interestingly, ING1 and ING2 are also complexed with histone methyltransferase (HMT) [39,43]. The HMT activity can methylate both histones H1 and H3 at the amino-terminal residues. The ING2-associated HMT activity seems to methylate mono- and dimethylation of histone H3 at lysine 4 [43]. This finding suggests that ING factors recognize and modify histone marks with the PHD region that is required for chromatin association to active chromatin sites.

Interestingly, ING1 and ING2 also interact with the transcriptional coactivator p300, which has an intrinsic histone acetyltransferase (HAT) activity [23,44]. This interaction leads to epigenetic changes by hyperacetylation of histones, which seems to link ING function rather towards DNA repair pathway. In line with this, it has been shown that ING1 can interact with Gadd45a [45,46], and both ING1b and ING2a with PCNA [47,48] to mediate nucleotide excision repair.

Furthermore, ING1 and ING2 have been reported to directly bind to the promoter of CDK inhibitors, p16 and p21, respectively [49,50]. In line with this, ING1 up-regulates p16 transcription via p300 HAT activity and induces cellular senescence (Figure 1), while the underlying mechanism that ING2 positively regulates p21 transcription remains unclear.

**Figure 1:** Model of cellular senescence induction by the ING proteins as epigenetic regulators.

ING1 and p300/CBP interact with p16 promoter and positively regulate its transcription. As cyclin-dependent kinase (CDK) inhibitor, p16 inhibits the activity of cyclin-CDK complexes, thus, preventing the phosphorylation of retinoblastoma protein (pRb) keeping it active. Active pRb remains bound with E2F, a pro-proliferative transcription activator, and suppresses its activity. This leads therefore to the inhibition of cell proliferation and triggers the cellular senescence pathway. ING2 interacts with the promoter of p21, another CDK inhibitor, and activate p21 transcription via an unknown mechanism. On the other hand, ING2 can bind to cyclin D1 promoter via its PHD to the histone mark of H3K4me3. The recruitment of mSin3a/HDAC1 complex suppresses the transcription of the cyclin D1 gene. Both up-regulation of p21 and down-regulation of cyclin D1 reduce the cyclin-CDK complexes activity, leading to cellular senescence.

These suggest that ING tumor suppressors can either activate or inhibit the target genes transcription, and that ING proteins change their signaling dependent on the environment that induce a specific interaction with different factors. This may subsequently induce ING distinct pathways, e.g. cellular senescence or DNA repair. Indeed, both functions are important for tumor suppression.

Evidence suggests that ING1 is involved in regulating the replicative lifespan, as the knockdown of ING1b expression results in increased number of replications [35]. In addition the expression of ING1b was found to be 8 to 10 times higher in senescent cells compared to young proliferating human fibroblasts [35]. Although the induction of cellular senescence may be a multifactorial process, data suggest that the overexpression of only one of the tumor suppressors ING1 or ING2 leads to the induction of cellular senescence [22,23]. This indicates an overlapping functional role of ING1 and ING2. The overexpression of ING1b in non-tumorigenic primary human fibroblasts resulted in growth arrest with the induction of cellular senescence [22]. Similarly, the expression of ING2a in early passage of primary human fibroblasts also showed cellular senescence inducing capability [23]. The functional consequences of ING1 or ING2 to induce cellular senescence have also been evaluated in vivo. In a mouse model of mammary tumorigenesis, inactivation of ING1 indicated an increased tumor burden, whereas inactivation of ING2 showed the opposite result [49].

**Figure 1:** Model of cellular senescence induction by the ING proteins as epigenetic regulators.
senescence in transformed cancer cells, such as by re-expression, are not yet clear.

The molecular pathway to induce cellular senescence is also under investigations. A functional link of both ING isoforms has also been reported to increase p53 protein stability by posttranslational modification that enhances the transcriptional activity of p53 and thereby triggering the cellular senescence phenotype [34,51,52]. However, it seems that both ING1 and ING2 can trigger the cellular senescence via more than one pathway by revealing also a p53-independent pathway of ING-mediated cellular senescence [36,53] as well as via the p16-pRb pathway [49] (Figure 1).

Notably, the PHD region plays an essential role to induce cellular senescence. Human fibroblasts transfected with PHD mutants of ING1 with a deficient histone binding ability were not capable to undergo cellular senescence [54], thus supports the important role of PHD and histone binding of ING proteins for the induction of cellular senescence. This finding strongly links the epigenetic regulation of ING tumor suppressors with cellular senescence induction also in non-tumor cells.

The ING - PHD: a highly conserved ING-domain between plants, insects and mammals

The ING PHD domain is relevant for both epigenetic control and induction of cellular senescence. Interestingly, the human ING1b and ING2a proteins are not only sharing high homologies in their amino acid sequences, but is also found to have high amino acid homology to ING proteins of other species, especially in the PHD region (Figure 2).

(B) Human ING2a (NP_001555.1) is aligned to mouse ING2 (NP_075992.2), zebrafish INg2 (NP_001002448.1), fruit fly ING (NP_650656.1), C. elegans ING homolog (NP_496909.1), A. thaliana ING2 (NP_974026.1), and O. sativa PHD finger protein ING (NP_001048939.1).

The NCBI BLAST program (http://blast.ncbi.nlm.nih.gov) was used to identify the most homologous proteins to human ING1b or human ING2a in other species. In addition, the whole protein alignment between human INGs and other species were performed with EMBSSO needle program (http://www.ebi.ac.uk/Tools/) to calculate the percentage identity and homology of amino acid sequences. The amino acid positions of each protein are indicated. The plant homeodomain (PHD) region of each species is separately aligned and compared as the percentage identity to the human PHD. Among other isoforms in D. melanogaster (fruit fly), C. elegans and O. sativa, the identified ING homologs exhibit the highest similarity to both human ING1 and ING2.

In other species including mouse (Mus musculus), zebrafish (Danio rerio), fruit fly (D. melanogaster), nematode (C. elegans), and plant (Arabidopsis thaliana and Oryza sativa) ING homologues were identified. Among these species, the mouse ING1 (NCBI ref: NP_036049.2) and ING2 (NP_075992.2) proteins share the highest identity (90% and 96%) and homology (95% and 99%) of amino acid sequences to the human ING1b (NP_937862.1) and ING2a (NP_001555.1) proteins, respectively (Figure 2). Notably, both PHD amino acid sequences of ING1 and ING2 from mouse ING proteins are 100% identical to PHD of human ING proteins indicating an important function. This is further supported by the fact that more than 50% of the amino acid sequences in the PHD region of plants (A. thaliana and O. sativa) INGs are identical with the human PHD of INGs. Of note, the plants and fruit fly PHD domains share higher identity to human PHD as compared to the PHD region of C. elegans (Figure 2). The protein alignments suggest that ING proteins and their PHD seem to be under strong evolutional selection and therefore, existing and being conserved in many species.

Phenotype of genetic ING models in vivo

The role of ING factors in vivo was analyzed using mice knock-out models. Despite the high homology of ING factors between plants and mammals, which implies an important functional role, a relatively weak phenotype in the KO mice was surprising. KO mice of either ING1 or ING2 were viable but promote tumor development [36,37,55]. ING1 KO mice revealed a high incidence of B cell lymphoma development [36,55]. ING2 KO mice on the other hand, were observed with the development of soft-tissue sarcomas [37]. Of note, the male ING2 KO mice exhibited the particular phenotype of being infertile and having small testes. These mice showed deficient spermatogenesis, altered meiotic recombination, and failed to complete meiosis II [37]. Interestingly, ING2 also seems to play a role in preimplantation development [56]. The ING2 expression was observed to be rapidly increased during the 2-cell to 4-cell cleavage-stage. In line with this, KD of ING2 in mouse zygote slows down the embryonic development [56].

The relatively mild phenotype of KO mice of these highly conserved factors suggests that ING1 and ING2 share similar functions and may compensate for the loss of function of the ING2 or ING1 null mutant, respectively. Also other ING family members might take over some essential functions for null mutants of one ING gene. Thus, we propose that the presence of multiple ING genes might serve as a redundant
and security viability system to reduce disadvantageous mediated by mutations of one of the ING genes.

Similarly, in C. elegans the depletion of the ING homolog protein suggests that this protein inhibits ionizing radiation-induced germ-cell apoptosis [57]. Moreover, nematodes expressing a mutant ING protein exhibit a weak uncoordinated phenotype.

Although ING proteins are rarely studied in plants, the functions and effects of PHD fingers of other factors are well established. Many proteins in plant contain putative PHD fingers and were found to be involved in various developmental processes including flowering, development of anthers, and inflorescences [58-60].

ING factors epigenetically regulate the gene expression of both mRNA and miRNA genes [42,50,61,62]. However, not much is known and questions remain open to better understand the epigenetics of ING factors epigenetically regulate the gene expression of both normal cell cycle by ING isoforms, (IV) sensing of cellular stress and its signaling that affects ING-factors to induce cellular senescence, apoptosis or DNA repair. Further it is unclear (V) which of the ING isoform functions is lost or decreased during early tumorigenesis.

Also the functional overlapping role of tumor suppressive function by each ING factor is unclear. However, there must be important biological reasons for the ING proteins to be naturally selected with a very high preservation of their amino acid sequence. Thus, many questions remain open to better understand the epigenetics of ING pathway as tumor suppressors and with relevance to human aging.

Acknowledgement

We are grateful to Dr. Mohsen Esmaeli and Tim Schmaeche for critically reading the manuscript.

References


