Liberation from fibrogenesis or tumorigenesis via cellular senescence by a novel small molecule

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Abstracts

Uncontrolled proliferative diseases such as fibroses or cancers are fatal human disorders. Previously, we found that a chromone-scaffold derivative called ONG41008 had a strong anti-fibrotic effect on in vitro fibrogenesis as well as in a murine lung fibrosis model. It later occurred to our attention that while ONG41008 remarkably attenuated proliferation of diseased human lung myofibroblasts (DHLF), resulting in replicative senescence (RS) typified by cell flatness, normal human lung fibroblasts were not affected. Video demonstration revealed that RS was evident within 48hr after ONG41008 treatment. No ONG41008 affected activated caspase 3 and mitochondrial membrane potential in DHLF. An interactome study suggested that metabolic shift, chromatin remodeling, or cell cycle control may be required for the RS. This observation prompted us to be engaged in cellular senescence of tumor cells. Clearly, senescent cells were conspicuously but temporarily observed in A549, adenocarcinomic human alveolar epithelial cells, giving us confidence that dysregulated cell proliferation could be a common underlying principle conserved in both DHLF and A549. An early phase of stimulation of A549 by ONG41008 led to RS followed by multinucleation (MNC), which has been known to be oncogene-induced senescence (OIS). MNC was immediately followed by apoptosis. Concomitant with massive upregulation of p16 and translocation to the nuclei, complete cell death of the remaining A549 occurred. Induction and nuclear translocation of p21was also noted in both A549 and DHLF stimulated with ONG41008. No induction of TP53 was seen but phosphorylation of TP53 was substantially increased in A549. Both immunocytochemistry and western blots corroborated these common senescent imaging features. With comparative analyses, it is clear that ONG41008 exhibited much lesser toxicity on normal human lung fibroblast than SAHA (suberoylanilide hydroxamic acid) and Nintedanib.

Taken together, all these studies strongly suggest that ONG41008 is a potent inducer of RS or OIS, presumably resulting in cessation of cell cycle are at G1 or G2 stage and/or systemic cell death. To our best knowledge, liberation of uncontrolled proliferative cells from fibrogenesis or tumorigenesis by a small molecule *in vitro* is an unprecedented case. ONG41008 could be a potential and safe drug for a broad range of fibrotic diseases or tumorigenic diseases.

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Key words

Anti-proliferation, Cellular senescence, Fibrosis, Cancer, and Small molecule

Introduction

Uncontrolled cell cycle progression has been well appreciated in terms of close association with rise of fibrosis or cancer [1]. Cells need to tightly regulate cell cycle entry and exit [2], otherwise leading to disastrous consequences involving tumorigenesis [3]. Although targeting cell cycle progression by small molecules has been considered an intriguing cancer modality undesirable medical risks such as systemic inhibition of stem cell differentiation or aggravation of homeostatic immunity may happen [4]. In this regard, natural drug scaffolds such as chromone-scaffold (CS) largely discovered in flavones or isoflavones may be able to provide a great deal of medical benefits to these proliferative diseases because these natural products have been largely proven to be safe and to give rise to unexpected desirable effects on suppression of uncontrolled proliferation shown by pathogenic myofibroblasts or aggressive tumor cells [5]. Metastasis plays a conclusive role in cessation of onset of cancer and has been a cancer target [6]. And it has been increasingly clear that metabolic regulation is closely coupled to cell cycle in cancer cells. Exclusive dependence of tumor cell growth on aerobic glycolysis, alternatively called lactate fermentation, has been well established [7]. Similarly, myofibroblasts utilize aerobic glycolysis and tend to generate hypoxia, playing a central role in the initiation and perpetuation of fibrosis [8].

Flavones are members of the polyphenol family, a group of over 10,000 compounds have been exclusively found in the plant kingdom [9]. In general, these phytochemicals protect plants from radiation damage [10]. Due to their anti-oxidant or anti-inflammatory potentials, flavones have long been used to treat inflammatory diseases such as arthritis and asthma [11]. Chromone, 1,4-benzopyrone-4-one, is a central chemical scaffold constituting flavones and isoflavones [12]. We recently reported that eupatilin, a CS-containing compound from an *Artemisia* species and its analog called ONG41008 substantially inhibited fibrogenesis *in vitro*

and fibrosis *in vivo* via actin depolymerization followed by dismantling latent TGFβ complex (LTC). This resulted in inhibition of epithelial-mesenchymal-transition (EMT) [13]. Here, we show that ONG41008 is a potent inducer of replicative senescence (RS) in pathogenic myofibroblasts and oncogene-induced senescence (OIS) in A549, a lung adenocarcinoma cell, respectively, thereby being escaped from fibrogenesis and tumorigenesis. These therapeutic capabilities of ONG41008 in terms of anti-proliferation may open a door as a powerful new therapeutic modality as regenerative medicine for treating fibrosis as well as cancer.

Results

ONG41008-treated DHLF exhibited cell growth arrest, leading to replicative senescence

(RS).

Disease human lung fibroblasts from IPF patients (DHLF) are αSMA+ pathogenic myofibroblasts also expressing an array of muscular collages. We sought to identify differential therapeutic modes by two first-in class idiopathy pulmonary fibrosis (IPF) drugs, nintedanib and pirfenidone as compared to that of ONG41008. Due to a drug-repurposing from cancer to IPF, nintedanib has been well appreciated for induction of cellular toxicity whereas pirfenidone is known to be hepatotoxic and seems to indirectly modulates immune cells function, thereby producing lesser active TGFβ [14] [15]. We lately reported that ONG41008 was a TGFβ-biogenesis inhibitor, immediately being capable of establishing cessation of TGFR signaling, resulting in reprogramming EMT to MET [13]. As shown in Figure 1A, nintedanib was able to induce a robust cell death in 24hr and a longer exposure in 48hr virtually killed all DHLF. IC50 was 10.17µM at 72hr. Pirfenidone remained unresponsive in terms of cell survival. Interestingly, ONG41008 marginally affected cell survival and its maximal survival rate remained over 60% regardless of increase in ONG41008 concentrations, resulting in no IC50 value. This strongly indicate that a systemic growth arrest may largely occur at the ONG41008 treatment of DHLF. Our previous 14 Day non-GLP Tox. study suggests that ONG41008 was essentially non-toxic to mice and rats and no perturbed pathologic features were observed (data not shown). As a control experiment, normal human lung fibroblasts (NHLF) were similarly treated as in the case of DHLF. Surprisingly, while nintedanib gave rise to a drastically low cell survival rate and pirfenidone remained unresponsive as in the case of DHLF. However, ONG41008 didn't essentially affect NHLF, not allowing us to calculate IC50.

But, the IC50 of nintedanib was around 30μM (Figure 1B). The susceptibility of DHLF to nintedanib in terms of apoptosis was more enhanced than that of NHLF.

In order to rule out involvement of apoptosis in ONG41008-mediated growth arrest in DHLF, three apoptotic means were exploited; 1) activated Caspase3 assay, 2) mitochondrial membrane potential, and 3) LDH cytotoxicity assay. FCCP (trifluoromethoxy carbonylcyanide phenylhydrazone) was used as a control for mitochondrial membrane potential. As shown in Figure 2A, 2B, and 2C, only nintedanib was able to cause a remarkable apoptosis, giving rise to their respective IC50 or EC50, suggesting that the systemic growth arrest induced by ONG41008 was not a consequence of apoptosis and would rather be related to cellular senescence. A hallmark of RS is cell flatness [16]. We previously found that GATA6 plays an important role in perpetuating EMT in DHLF [13]. We assumed that expression of GATA6 would discontinue in the absence of TGFβ but in the presence of ONG41008. A RS model was set up with time lapses microscopy along with increasing ONG41008 concentrations. As shown in Figure 3A, GATA6 expression was consistently seen in the entire RS setting, which remains to be further elucidated. RS was generated at 5µM ONG41008 and became more evident at 10μM. The maximal extent of the RS occurred at 20μM. DHLF in the absence of ONG41008 remained unchanging. This study singled out a conclusion that ONG41008 was a potent inducer of cellular senescence, namely RS, in DHLF. Moreover, two sets of video demonstration clearly showed that RS was evident in the presence of 10µM ONG41008 (Supplementary Video Imaging I at high cell density and II at low cell density). An RNA-seq analysis was done, giving rise to a nuclear interactome as shown in Figure 3B. EGR1 (Early Growth Response 1) was a priming protein, which interacted with an array of proteome involving CDC45 (Cell Division Cycle 45), Transforming Acidic Coiled-Coil Containing Protein 3 (TACC3), or Cyclin Dependent Kinase 1 (CDK1), which play pivotal roles in onset

of nuclear events in cell differentiation, DNA replication, cell cycle control or cellular senescence [17] [18] [19] [20]. We also discovered that unique transcriptomes were also responsible for driving replicative senescence. RNA-seq analysis was again conducted to compare difference in gene expression between DHLF and 10µM ONG41008-treated DHLF. Upregulated genes were sorted out by p-values. As shown in Supplementary Figure 1 and 2, three interactomes seemed to be working cooperatively for being led to RS; the first one was metabolic interactome typified by pyruvate dehydrogenase kinase (PDK)1, the second one was actin biogenesis, and the last one was related to histone modification, DNA replication, and generation of a muscle-neuron signature whereby the DHLF expression signature could be transiently going through to the RS. The reason being generation of a muscle-neuron signature remains to further explored. Interestingly, vast majority of top-ranked genes affected by ONG41008 were soluble factors or receptors (Supplementary Table I). We imagined that prior to euchromatin formation driving RS expression of this array of cytokines or interaction with cognate-receptors might happen for priming cellular senescence.

ONG41008-mediated RS may be required for participation of TP53, p21 and p16 in DHLF

It has been appreciated that several tumor suppressor proteins are responsible for cessation of cell cycle progress [21]. TP53 undergoes phosphorylation and is mainly coupled to p21 or p16, leading to cell cycle inhibition. As shown in Figure 4A, TP 53 was not induced but rapidly translocated to the nucleus 1hr after 1 μ M ONG41008 treatment. The same held true that p21 in DHLF treated with 1~10 μ M ONG41008 was upregulated and translocated to the nuclei (Figure 4B). With respect to p16, it was also upregulated and translocated to the perinuclear zone and the nuclei (Figure 4C). It has been well documented that the cooperative

works between TP53, p21 and p16 plays pivotal roles in cell cycle control (Ref. 22). We conducted a cell cycle analysis of DHLF stimulated with ONG41008. As shown Figure 4D, ONG41008 rapidly ceased cell cycles of DHLF at G1 or G2.

Cellular senescence-mediated apoptosis of A549 by ONG41008

Pathogenic myofibroblasts and tumor cells are in common in many ways; first, aerobic glycolysis is the major catabolism of glucose. Second, uncontrolled cell division occurs. Third, immune escape is a major cause for perpetuation of their existence. The observation that DHLF were rapidly converted to RS by ONG41008 prompted us to see if cancer cells would similarly behave. Cellular senescence was established in A549 with the treatment of 10μM ONG41008 for 24hr, 48hr or 72hr, and immunohistochemistry was conducted enabling us to see morphological change, namely cell flatness [22]. As seen in Figure 5A, flat cell morphology was evident in A549 treated with ONG41008 for 24hr and multinucleated A549 began to appear (Figure 5A and 5B). It has been appreciated that multinucleation of cancer cells is caused by called oncogene-induced senescence (OIS) [23]. Akt1 activation has been detected in A549 [24]. In fact, our ICC data shows that the phosphorylated-Akt1existed in A549 and its levels were augmented in proportional to increasing concentrations of ONG41008 (Supplementary Figure 3). The greatest number of multinucleated A549 was observed at 48hr (Figure 5B), after which the number was significantly reduced (Figure 5C). These studies suggest that ONG41008 was a potent inducer of OIS [25].

Taken together, ONG41008 was able to induce OIS in 549 such that ONG4100-exposed A549 underwent a systemic apoptosis.

Phosphorylation of TP53 and nuclear translocation of phosphorylated TP53, p21, and p16 rapidly occurred prior to the OIS of A549 stimulated with ONG41008

As in the case of induction of RS shown in DHLF we further explored changes of TP53, p21, and p16, which are able to induce OIS in A549. As shown in Figure 6A and 6B, while TP53 protein expression remained unchanging but its nuclear location could be seen at 1µM ONG4008, p21 was upregulated and similarly nuclear localized. p16 was upregulated in proportional to varying concentrations of ONG41008 and was nuclear localized as well as distributed at the perinuclear zones (Figure 6C). We discovered that cell morphology became more homogenous at 72hr after ONG41008 treatment than that of 24hr-long ONG41008 treatment, which may reflect elimination of multinucleated A549. And a maximal translocation of p16 to the nuclei seemed to be completed at 72hr. These p16-satutated A549 would be supposed to undergo apoptosis (Figure 6D and Figure 7B). Since phosphorylation of TP53 is known to be essential for its capability of regulating cell cycle arrest [26], we conducted western blot analysis by using a phospho-specific TP53 antibody. Increase in the concentrations of ONG41008 was proportional to degree of phosphorylation of TP53 (Figure 7A) and total amounts of TP53 remained same. We also noticed that both p21 and p16 were induced in a concentration-dependent manner, which corroborated the results of ICC (Figure 6A, 6B, and 6C). Respective molecular positioning was denoted by yellow circles. We continued to stimulate A549 in the presence of both TGF\$\beta\$ to make A549 more proliferative in conjunction with ONG41008. It was observed that a significant fraction of A549 died off from 6 days and complete cell death occurred at 15 days whereas control A549 remained vibrant (Figure 7B). While blue circles represented control A549 growth, blue and red circles denoted control A549 and ONG41008-treated A549, respectively. All cells exposed to 10µM ONG41008 seemed to receive apoptotic signals during 72hr.

In addition to A549, a human aggressive breast cancer cell lines called MCF7 were brought into focus; MCF7 is a human primary breast cancer cell line. These cells and A549 were stimulated with 20µM ONG41008 for 48hr and ICC was conducted. A549 MCF7 showed a robust cellular senescence (Figure 8A). The number of each MNC were counted (Figure 8B. This data may suggest that the degree of MNC formation was varying due to cellular context. We also conducted a cell cycle analysis of A549 stimulated with ONG41008.

Taken together, ONG41008 was able to induce cellular senescence in cancer cells, most likely being OIS, leading to apoptosis.

Comparison of anti-cancer capabilities between ONG41008 and SAHA

MOA (mode of action) of ONG41008 to eliminate cancer cells seems to be induction of OIS. There are numerous anti-cancer drugs targeted to cell cycle control, DNA replication, or chromatin remodeling, etc. We explored how cellular survival rate and mitochondrial membrane potential were affected by these drugs. We decided to compare between ONG41008 and SAHA (suberoylanilide hydroxamic acid), a reversible pan-histone deacetylase (HDAC) inhibitor, because effect of ONG41008 on cell cycle regulation and that of SAHA on chromatin remodeling seemed to be inter-related to cell cycle division control as well. It should be noted that an important nod in the A549-ONG41008 was HIST1H4K (H4 Clustered Histone 12) involved in DNA replication and DNA repair (Supplementary Figure 4). As shown in Figure 9A, while substantial attenuation of cell survival by SAHA rapidly occurred at 24hr and seemed to reach a maximum at 72hr, an inhibitory effect by ONG41008 began at 48hr and continued to be increased to 4 days till all cells were eliminated. The cell killing kinetics made sense in that initiation of OIS by ONG41008 begin at 48hr and continued to be increased till 72hr based upon our ICC data as seen in Figure 5. The same held true for mitochondrial membrane

potential (Figure 9A). Either IC50 showed around 25-fold differences. These data suggest that

ONG41008-mediated apoptosis was a consequence of induction of OIS whereas SAHA

exhibited a rapid cell death due to its HDAC inhibitory effect on a broad range of chromatin

remodeling processes, likely a reason being many undesirable effects associated with SAHA.

The extent of apoptotic capacity and time duration by ONG41008 was distinguished from those

of SAHA or other anti-cancer drugs. In order to ascertain whether ONG41008, SAHA and

Nintedanib, an originally designed anti-cancer drug, differentially affected NHLF a set of

CCK-8 assays were conducted. As shown in 9B, it is clear that while both SAHA and

Nintedanib exhibited tremendous toxic effects on NHLF ONG41008 did not give rise to cell

toxicity.

An A549-ONG41008 interactome

Fas may be a central nod, which was the death-domain containing cell surface receptor in OIS-

mediated apoptosis in A549 (Figure 10). As an additional evidence supporting that systematic

apoptosis could be initiated from additional nodes of A549-ONG41008 interactome. By and

large, the interactome study showed that FAS (a cell surface death receptor), NGFR (Neuronal

Growth Factor Receptor), a death-domain containing cell surface receptor, and PGF (Placental

Growth Factor) were linked to TRIAP1 (TP53 Regulated Inhibitor of Apoptosis 1) as an

apoptosis nod of the interactome (Supplementary Figure 4). Furthermore, a TP53-regulated

cell cycle hub was linked to the apoptosis hub through which CDKNIA also called p21 was

situated between these two interacting hubs. The TP3-regulated cell cycle hub comprised

CEBPA, TP53INP1, BTG2, SSN1, SSN3, and HIST1H4K. Gene ontology was summarized in

the pertinent boxes in Supplementary Figure 4. Interestingly, we noted that GDF15, a human

TGFβ family member, may play an interesting role, probably in connecting two distinct

systems and stabilizing them. GDF has been shown to be able to inter-connect two distinct biological systems [27].

Discussion

Fibrosis and cancer are still intractable and fatal diseases because fundamentals of these diseases are directly related to cell cycle regulation. Cell cycle control is meticulously regulated such that proliferation, differentiation or apoptosis should be harmonized. Dysfunction of this tight regulation may lead to vulnerability initiating serious diseases. Uncontrolled proliferation is the core of the vulnerability. The current anti-fibrotic and anticancer drugs are largely targeted to uncontrolled proliferation [28]. A number of side effects are associated with these drugs, leaving behind us regrets. Immune checkpoint inhibitors (ICIs) are becoming a very successful modality for dealing with cancer with much lesser side effects [29]. Immunity is able to distinguish self from non-self, conforming to *in vivo* cancer immunity. Here, we show that RS and OIS generated by ONG41008 were able to make DHLF and A549 liberated from uncontrolled proliferative status, respectively. Although we haven't tested many normal cells ONG41008, at least, was capable of clearly distinguishing pathogenic myofibroblasts from normal lung fibroblasts, giving a great surprise to us. When ONG41008 was orally administered into mice or rats no pathological abnormalities or toxicities were seen at 1g of ONG41008 (preclinical study number: U-18156), suggesting that ONG41008 is a safe drug for normal cell types.

Pathogenic myofibroblasts and cancer cells are in common such that metabolic features gaining ATP stem from aerobic glycolysis, resulting in dysregulated redox potential represented by NAD/NADH ratio. ATP levels or intracellular pH may become similar. Chromone scaffold (CS) is the core structure found in flavones or isoflavones [30], which has

been known to capture UV in such a way that plants are protected from UV damage. Although

over 10,000 kinds of CS-containing derivatives (CSD) have been discovered the number of

CSD exhibiting anti-fibrotic capacity seems to be limited, suggesting that not only CS but also

a variety of side chains linked CS may contribute to anti-fibrosis. It has been reported that CS

plays a role in anti-cancer activity; SB203580 [31]. Since ONG41008 is able to distinguish

pathogenic myofibroblasts (DHLF) and cancer cells (A549) we would like propose that the CS

of ONG41008 may be able to recognize some elements comprising intracellular micro-

environments involving NAD, NADH, pH or ATP as an energy sensor.

All thing considered, we believe that ONG41008 is a potent inducer of cellular

senescence (RS and OIS) and pathogenic myofibroblasts or cancer cells can be liberated from

their diseased uncontrolled proliferation loops.

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Competing financial interests

B-S Youn retain the shares of Osteoneurogen, M-K Meang retains a stock option, and SB Kim

are employed by OsteoNeuroGen. The current contents of the ONG41008 data has been

subjected to a Korean provisionary patent.

Author contributions

B-S Youn conceived the idea and wrote the manuscript. MK Maeng was largely involved in

executing major experiments involving methods and materials. SB Kim played a major role

in conducting drug responsiveness of DHLF and NHLF.

Footnotes:

The following abbreviated terms were mainly used in this manuscript;

CS: Chromone Scaffold

CSD: Chromone Scaffold Derivatives

DHLF: Diseased Human Lung Fibroblasts from IPF patients

NHLF: Normal Human Lung Fibroblasts

RS: Replicative Senescence

OIS: Oncogene-Induced Senescence

MNC: Multinucleation

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Figure legends

Figure 1. Comparison of survival rate of DHLF and NHLF treated with ONG41008,

Nintedanib, or Pirfenidone A-B) DHLF (Diseased Human Lung Fibroblasts) and NHLF

(Normal Human Lung Fibroblasts) were stimulated in the absence of TGF β with various

concentrations of the drugs as indicated up to 30µM for 24hr, 48hr and 72hr. Cell survival rate

was measured by CCK-8. IC50 of Nintedanib was calculated within Sigmoidal, Fourth Party

Logistics by GraphPad Prism 7.00.

Figure 2. No apoptotic effects on the induction of the ONG41008-mediated growth

arrest of DHLF DHLF were collected and re-seeded into each culture dishes pertinent to

corresponding apoptotic assays; A) Activated caspase3 assay was performed using ONG41008,

Nintedanib, or Pirfenidone. B) Mitochondrial membrane potentials were measured. FCCP was

used as a control being a mitochondrial oxidative phosphorylation uncoupler. C) Lactate

dehydrogenase (LDH) was assayed. IC₅₀ or EC₅₀ was measured at 72hr with Sigmoidal, Fourth

Party Logistics by GraphPad Prism 7.00.

Figure 3. Induction of RS by ONG41008 and an interactome analysis A) DHLF were

treated with different concentrations of ONG41008 along with control (RPMI medium).

Immunocytochemistry (ICC) was conducted with anti-human GATA6, phalloidin, and DAPI.

Morphological change was monitored under phase contrast microscopy. Typical replicative

senescent cells were denoted by "replicative senescence" white-colored. B) An RNA-seq

analysis was conducted using total RNAs prepared from non-treated DHLF or 10µM

ONG41008-treated DHLF. An interactome representing upregulated genes was established via

the String program based upon p-values (>0.005).

Figure 4. Translocation of TP53 to the nuclei and both induction and nuclear

translocation of p21 and p16 in DHLF upon ONG41008 treatment DHLF were treated

with 1µM to 10µM ONG41008 along with control (RPMI medium). ICC were conducted with

anti-human TP53, anti-p21, anti-p16, or phalloidin in conjunction with DAPI; A) Translocation

of p53 to the nuclei was demonstrated. B-C) Induction and translocation of p21 and p16 to

the nuclei were shown, respectively.

Figure 5. Capability of ONG41008 with respect to induction of a robust cellular

senescence leading to multinucleation A549 were stimulated with 10µM ONG41008 for

24hr, 48hr, or 72hr and subjected to ICC by using anti-ZEB1, DAPI, and Phalloidin; A) ICC

for ONG41008-treated A549 at 24hr was conducted in which flat cells representing MNC

denoted with white circles along with control presented. B-C) ICCs for ONG41008-treated

A549 at 48hr and 72hr were conducted, respectively. MNC were also denoted by white circles.

Figure 6. Translocation of TP53 to the nuclei, and both induction and nuclear

translocation of p21, and both induction and perinuclear and nuclear translocation of p16

in A549 upon ONG41008 treatment A-C) A549 were stimulated with varying

concentrations of ONG41008, 1µM to 20µM, and ICC was conducted with anti-TP53, anti-

p21, or anti-p16. D) A549 were treated with 10µM ONG41008 for 72hr during which control

and ONG41008 was everyday replaced with new media and new batch of 10µM ONG41008,

respectively. p16 expression was detected by ICC.

Figure 7. Western blot analysis of TP53, p21, and p16 and microscopic observation of cell death of A549 A) Cell lysates were prepared from A549 treated with different concentrations, 1μM to 50μM for 24hr and subjected to western blot using pertinent antibodies as indicated. Yellow circles represent respective molecular mass. B) A549 were continually treated with 20μM ONG41008 for 15 days in the presence of TGFβ (2.5ng/ml) in order to keep cells vibrant along with control A549. Cell death was monitored under phase contrast microscopy.

Figure 8. Induction of cellular senescence of human aggressive cancer cells upon

ONG41008 treatment A) A549 and MCF7 were stimulated with 20μM ONG41008 for

24hr. ICC was conducted with the use of Phalloidin and DAPI and merged imaging was

compared. B) MNC were calculated and compared between A549 and MCF7.

Figure 9. Comparison of anti-cancer capabilities associated with ONG41008 with SAHA and Nintedanib in A549 and their toxic effects on NHLF. A) Survival rates of A549 treated with ONG41008 or SAHA were measured via CCK-8. IC50s were measured.

B) NHLF were stimulated with ONG41008, SAHA or Nintedanib for various time points as indicated. CCK-8 assays were conducted. IC50s were measured.

ONG41008 While ONG41008 was responsible for inducing replicative senescence in DHLF it was able to induce OIS followed by multinucleation, which caused apoptosis of A549. Regardless of both cell types participation of TP53, p21, and p16 is required for both types of

senescence occurred in DHLF and A549. ONG41008 does not affect induction of TP53. Translocation of TP53, p21, and p16 is necessary for cell cycle arrest.

Materials and Methods

Cell culture and reagents

DHLFs were purchased from Lonza (Basel, Switzerland) and cultured in fibroblast growth medium (FBM, Lonza, Walkersville, MD, USA). Recombinant human TGFβ and PDGF were obtained from Peprotech (Rocky Hill, CT, USA) and used at a final concentration of 5 ng/ml. Chemically synthesized ONG41008 was obtained from Syngene International Ltd. (Bangalore, India), dissolved at a stock concentration of 50 mM in DMSO, and stored in aliquots at -20°C. DMSO with according concentration was used as control. RAW264.7 cell line was purchased from Korean Cell Line Bank (Seoul, Korea) and cultured in RPMI supplemented with 10% FBS and 1% P/S (Welgene, Seoul, Korea). LPS was purchased from Sigma and used at final concentration of 100 ng/ml.

Immunocytochemistry

Cells were fixed using 4% paraformaldehyde, permeabilized with 0.4% TritonX100, blocked with 1% BSA and incubated with Rhodamine Phalloidin(Thermo Fisher, Massachusetts, USA), anti-GATA6(Abcam, Cambridge, UK), Anti-p53(Cell signaling technology, Beverly, MA), p21(Abcam, Cambridge, UK), p16-INK4A(Proteintech, IL, USA), ZEB1(Cell signaling technology, Beverly, MA) for 4 hour at RT. After washing, cells were incubated with Alexa Fluor 488(Abcam, Cambridge, UK) conjugated secondary antibody. Images were analyzed using EVOS M7000(Invitrogen, CA, USA)

Western Blot

A549 cells were seeded at 1×10^6 cells/well in 100mm cell culture dishes and incubated

overnight, followed by treatment with various concentrations of ONG41008. After 24 hr, The

cell lysates were clarified by centrifugation at 14,000 x g for 10 minutes and the supernatant

was collected. The protein concentrations were quantified by Bradford assay (Thermo Fisher,

Massachusetts, USA). Thereafter, 25ug of cellular protein was loaded on 10% SDS-PAGE gel

and transferred to Nitrocellulose membranes. After blocking with 5% BSA, the membranes

were incubated with Anti-p53(Cell signaling technology, Beverly, MA), phosphor-p53(Cell

signaling technology, Beverly, MA), p21(Abcam, Cambridge, UK), p16-INK4A (Proteintech,

IL, USA), GAPDH (Abcam, Cambridge, UK) overnight at 4'C. after washing thoroughly,

membranes were incubated with HRP-conjugated secondary antibody. Protein band was

visualized using ECL reagent (Abfrontier, Korea) and Uvitec HD9(UVITEC, UK)

Live Imaging

DHLFs were seeded at 12 well cell culture plates, and after 24hr, TGF beta (5ng/ml),

Nintedanib (10uM), Pirfenidone (10uM) and ONG41008 (10uM) were treated. Cells were

incubated in EVOS M7000 CO² incubation chamber (Invitrogen, CA, USA), cell morphology

images were captured every 30 minutes. The snapshots were collected and attached as dynamic

video.

CCK-8 assays

Cells were seeded onto 96well plates at a density of 3×10^4 cell/well for 24hr. Then the cells

were treated with 0.01, 0.1, 1, 10, 20, 30uM ONG41008 or Pirfenidone or Nintedanib. After

treatment, cell survival rate was measured by cell counting kit-8 (Dojindo, CK04) as

manufacturer's protocol.

Caspase-3 assay

The caspase-3 activity was measured using Caspase-3 assay kit (abcam, ab37401). Cells treated

with various concentrations of ONG41008 or Pirfenidone or Nintedanib were harvested and

lysed on ice. Thereafter, protein was measured by BCA (Thermo, 23227) and adjusted to 50ug

protein per 50ul cell lysis buffer. Then the caspase-3 activity was examined by following

manufacturer's protocol.

Mitochondrial Membrane Potential assay

Cells were seeded in 96 wells plate for 24hr, then exposed to different concentration of

ONG41008 or Prifenidone or Nintedanib. The cells were then co-incubated with TMRE

(abcam, ab113852), for 30min 37°C in the dark. Cells treated 20uM FCCP were used as

positive control. After that, mitochondrial membrane potential was examined by following

manufacturer's protocol.

LDH assay

LDH release were detected by using the LDH assay kit (abcam, ab56393). The cell culture

plates were centrifuged at 480g for 10min, and supernatants (10ul/well) were extracted into

another 96well plate. Then, 100ul LDG reaction mix was added to each well and incubation

for 30min at room temperature. The absorbance values were measured at 450nm on the

microplate reader.

RNA-seq processing, differential gene expression analysis, and interactome analysis

Processed reads were mapped to the *Mus musculus* reference genome (Ensembl 77) using Tophat and Cufflink with default parameters. Differential analysis was performed using Cuffdiff using default parameters. Further, FPKM values from Cuffdiff were normalized and quantitated using the R Package Tag Count Comparison (TCC) to determine statistical significance (e.g., P values) and differential expression (e.g., fold changes). Gene expression values were plotted in various ways (i.e., Scatter, MA, and Volcano plots), using fold-change values, using an R script developed in-house. The protein interaction transfer procedure was performed using the STRING database with the differentially expressed genes. A 60 Gb sequence was generated, and 10,020 transcripts were read and compared. The highest-confidence interaction score (0.9) was applied from the *Mus musculus* species, and information about interacts were obtained based on text mining, experiments, and databases (http://www.string-db.org/). Due to company information sake the above detailed RNA-Seq or interactome data interpretation would be limited but essential data sufficiently supporting our assertion were provided.

Reverse transcriptase PCR and real-time PCR

Cells cultured in either 12 or 24-well plates were washed twice with cold PBS and harvested using TaKaRa MiniBEST Universal RNA extraction kit (Takara, Japan). RNA was purified using the same kit according to manufacturer's protocol. RNA was reverse-transcribed using the cDNA Synthesis Kit (PCRBio Systems, London, UK). Synthesized cDNA was amplified with StepOne Plus (Applied Biosystems, Life Technologies) and 2× qPCRBio Probe Mix Hi-ROX (PCRBio). Comparisons between mRNA levels were performed using the ΔΔCt method, with GAPDH as the internal control.

Supplemental information

Suppl. Video demonstration I: A 72hr-long video photographing at high DHLF cell density

was conducted.

Suppl. Video demonstration II: A 72hr-long video photographing at low DHLF cell density

was conducted.

Suppl. Table I: Top replicative senescence-initiating genes: soluble factors / receptors. An

RNA-seq analysis was conducted with the use of total RNAs prepared from DHLF or

ONG41008-treated DHLF for 24hr. Top twenty genes showing p>0.005 were selected.

Suppl. Figure 1 and Figure 2: An RNA-Seq was performed as described above. The genes

were sorted out in such a way that three sets of genes showing their p-values were

corresponding to between 0.005 and 0.05, or between 0.05 and 0.5 were selected and selected

genes were further analyzed via the String Program, giving rise to three interactomes.

Suppl. Figure 3. Phospho-AKT expression pattern of A549 treated with ONG41008. A549

were stimulated with increasing concentrations of ONG41008 (1mM to 20mM). ICC was

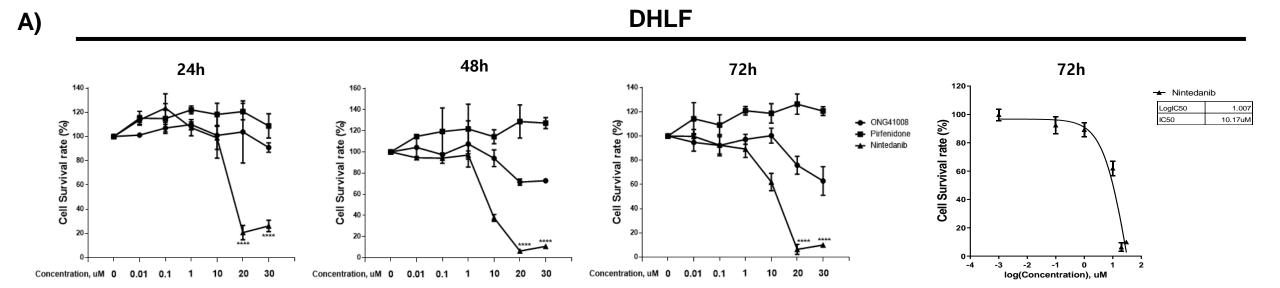
performed using anti-phospho-specific Akt1 and Phalloidin

Suppl. Figure 4. A549-ONG41008 network An A549-ONG41008 interactome was

established. An RNA-seq analysis was conducted with the use of total RNAs prepared from

A549 or ONG41008-treated 549 for 24hr. Top twelve upregulated genes showing p>0.005

were selected and their interactome was established via the String Program.



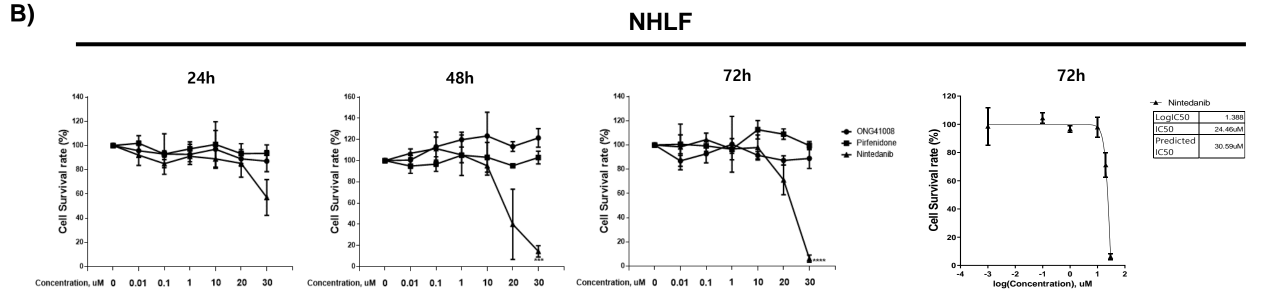
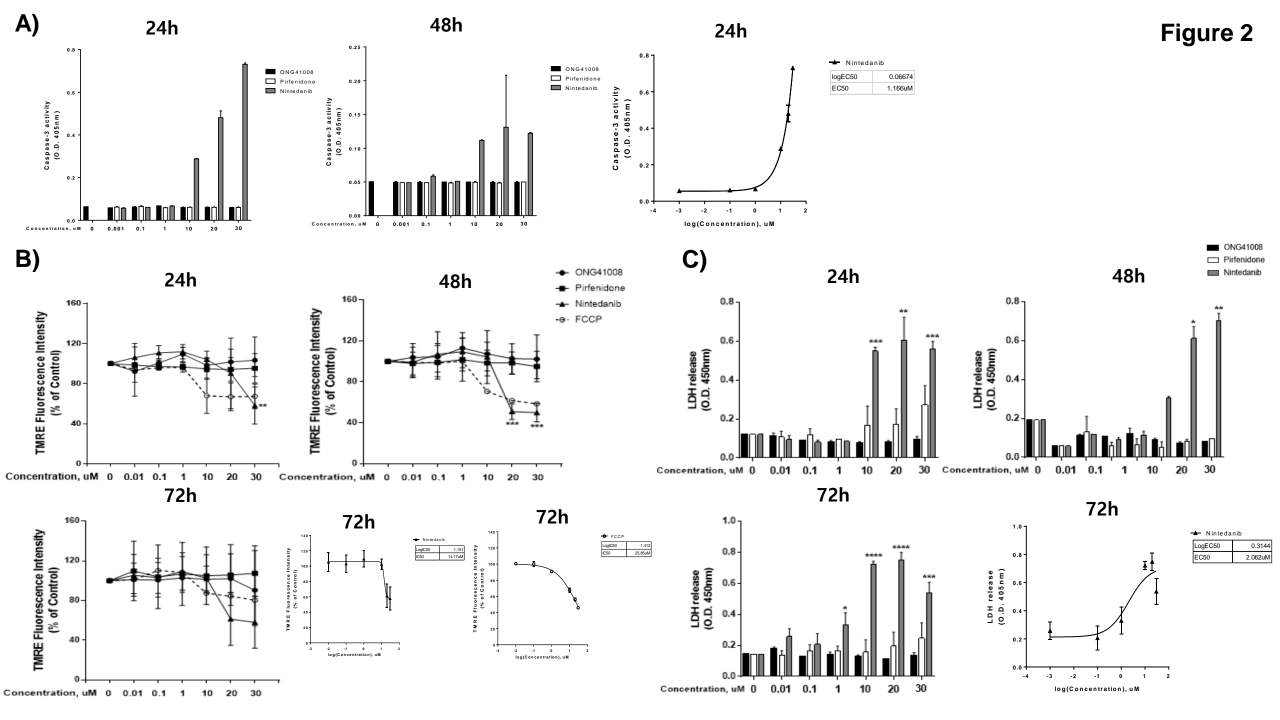


Figure 1



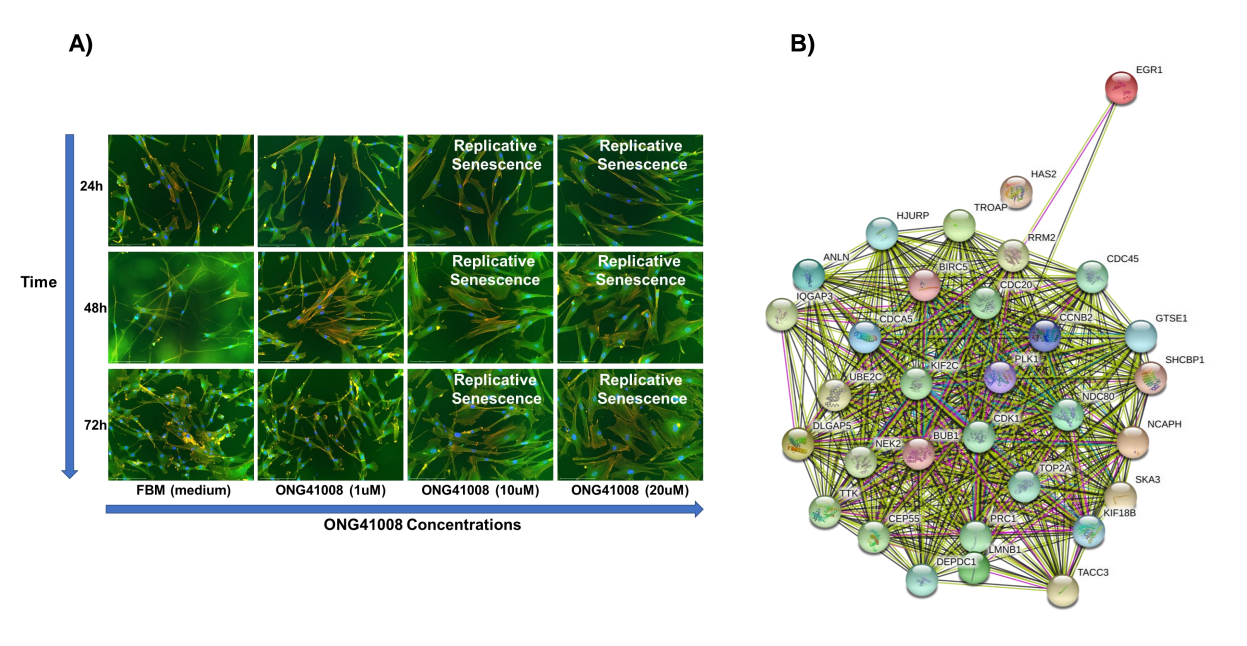
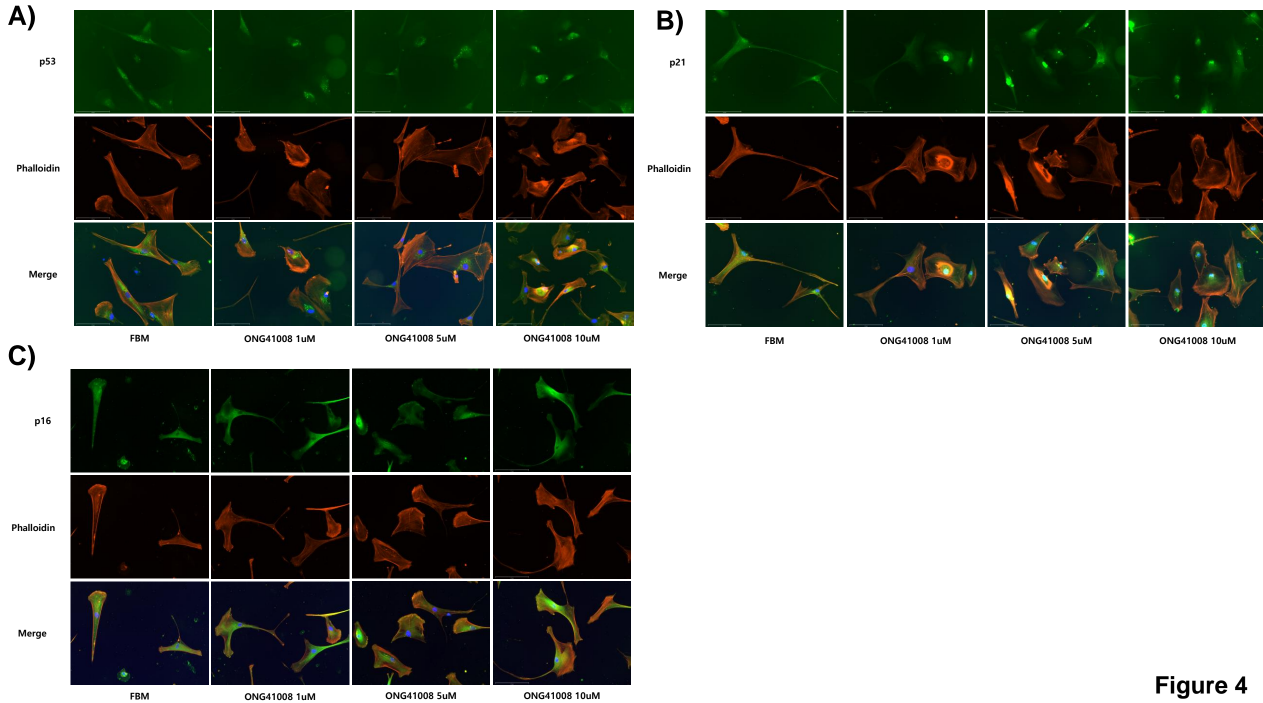


Figure 3



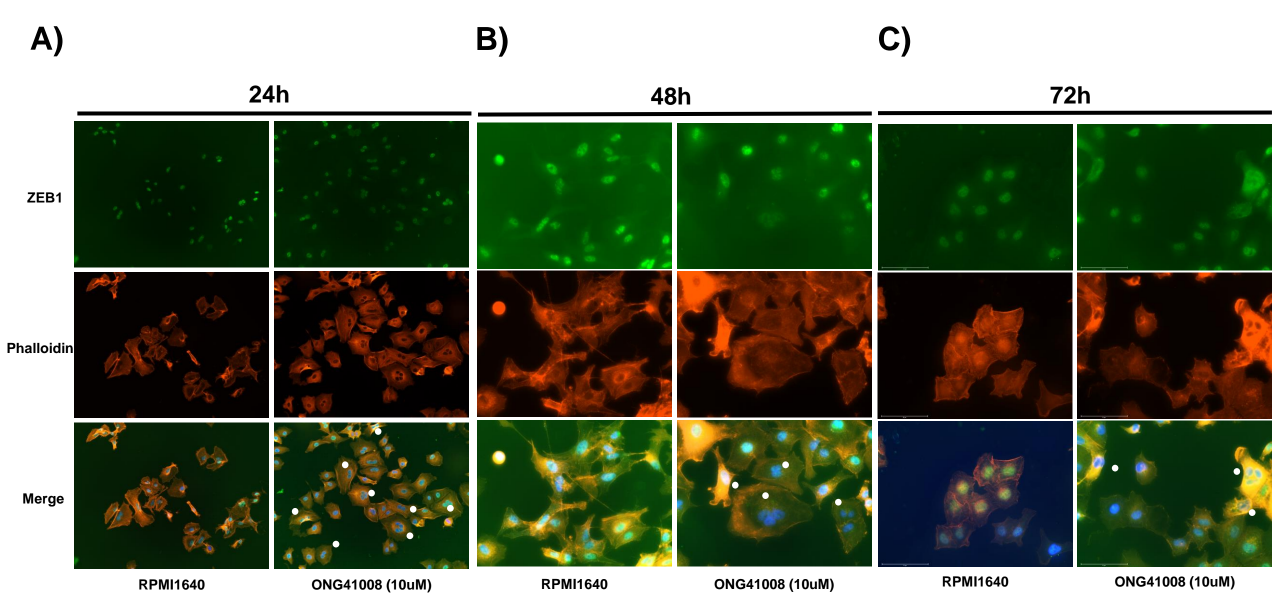
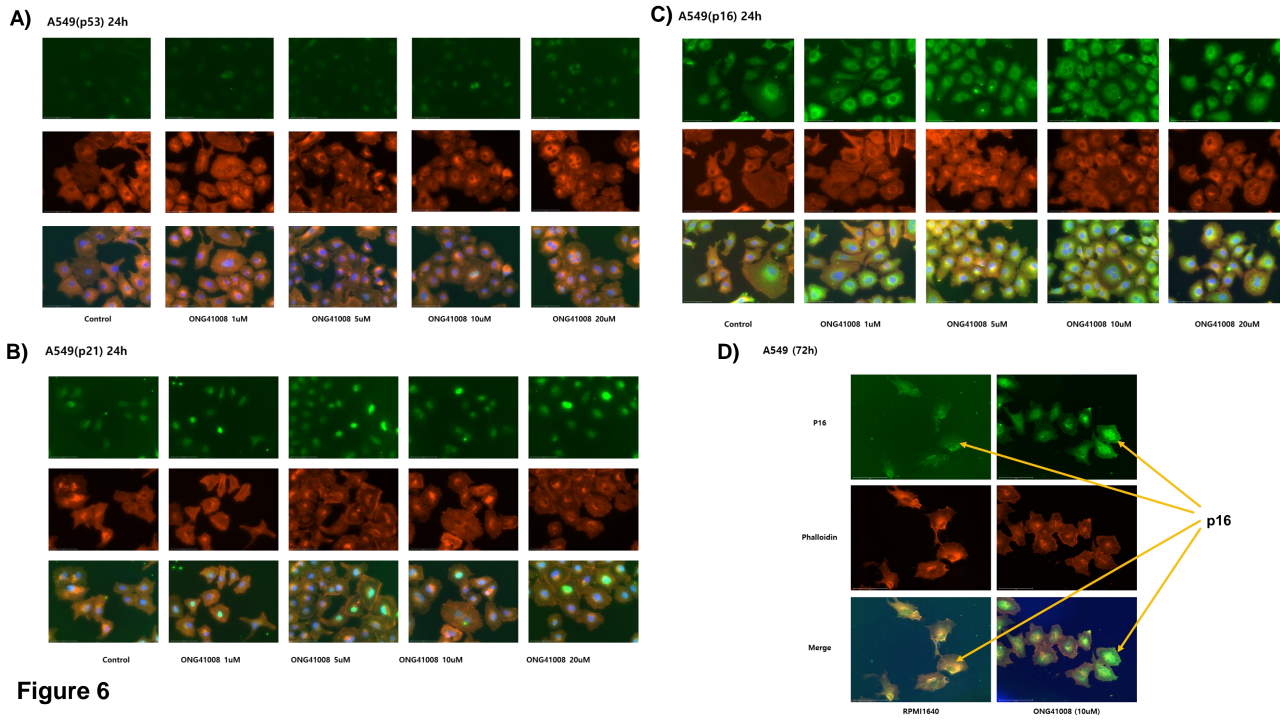
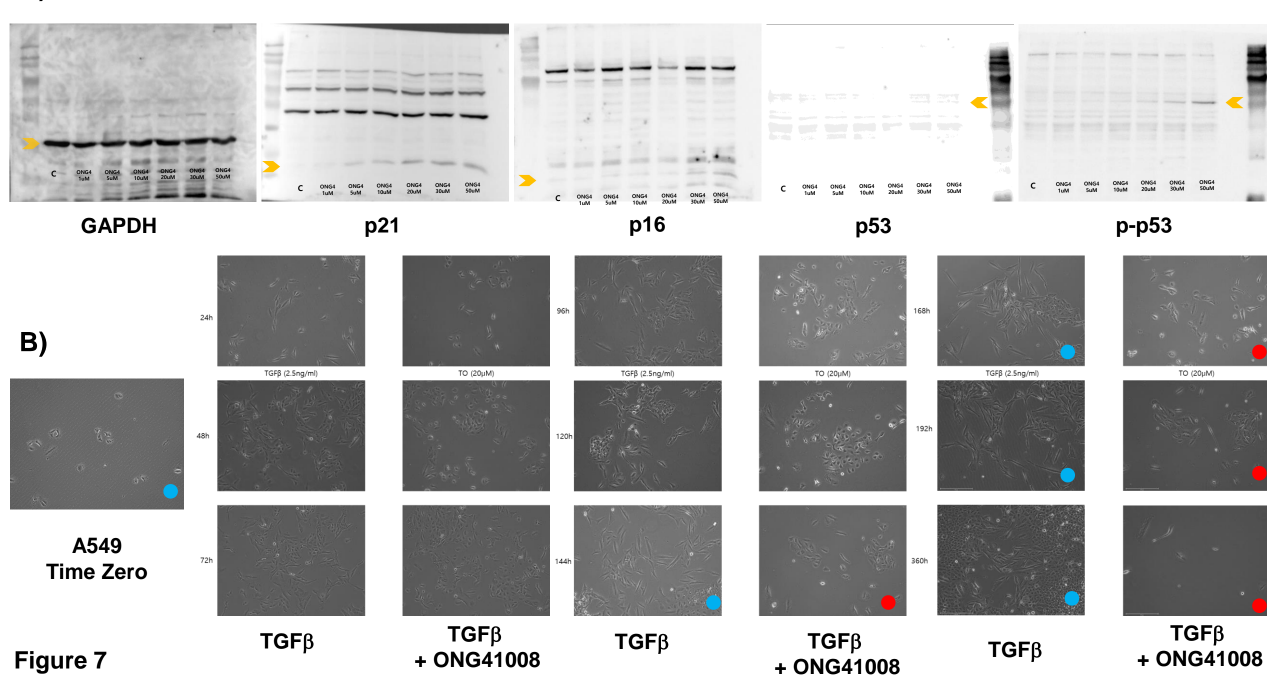


Figure 5



A)



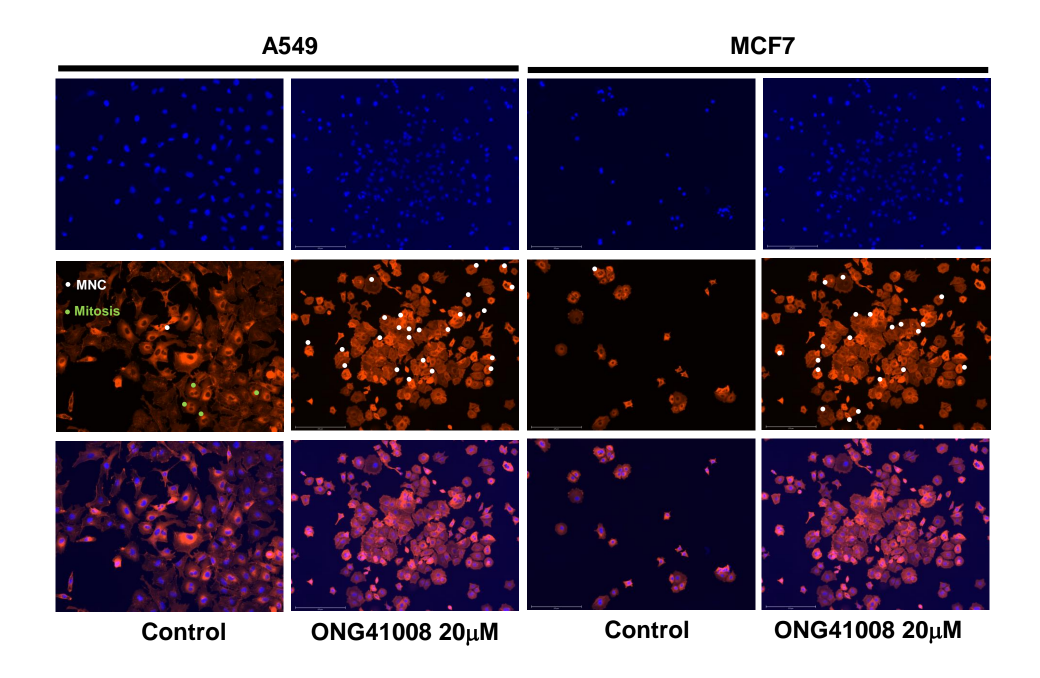
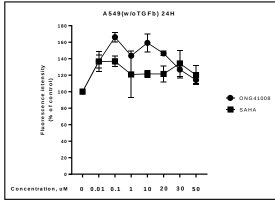
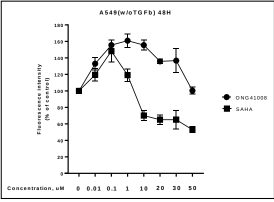


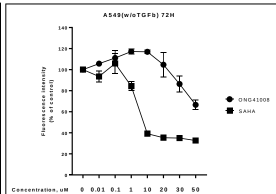
Figure 8

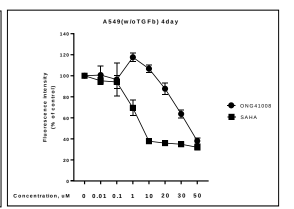


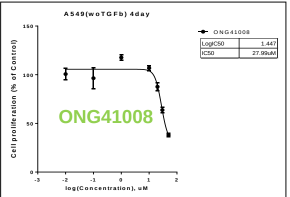
A549



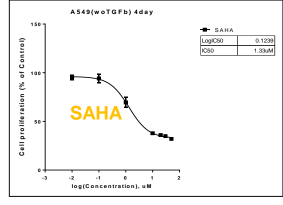




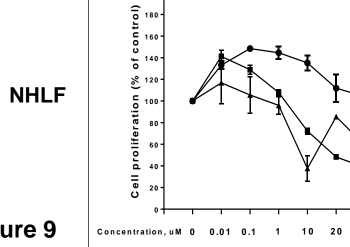




NHLF(woTGFb) 48h

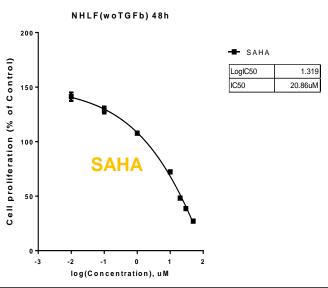


B)



200





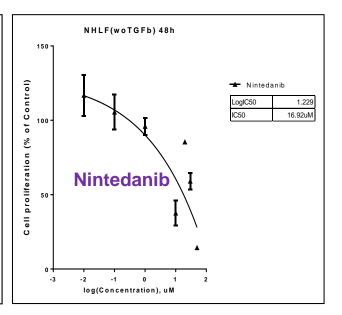


Figure 9

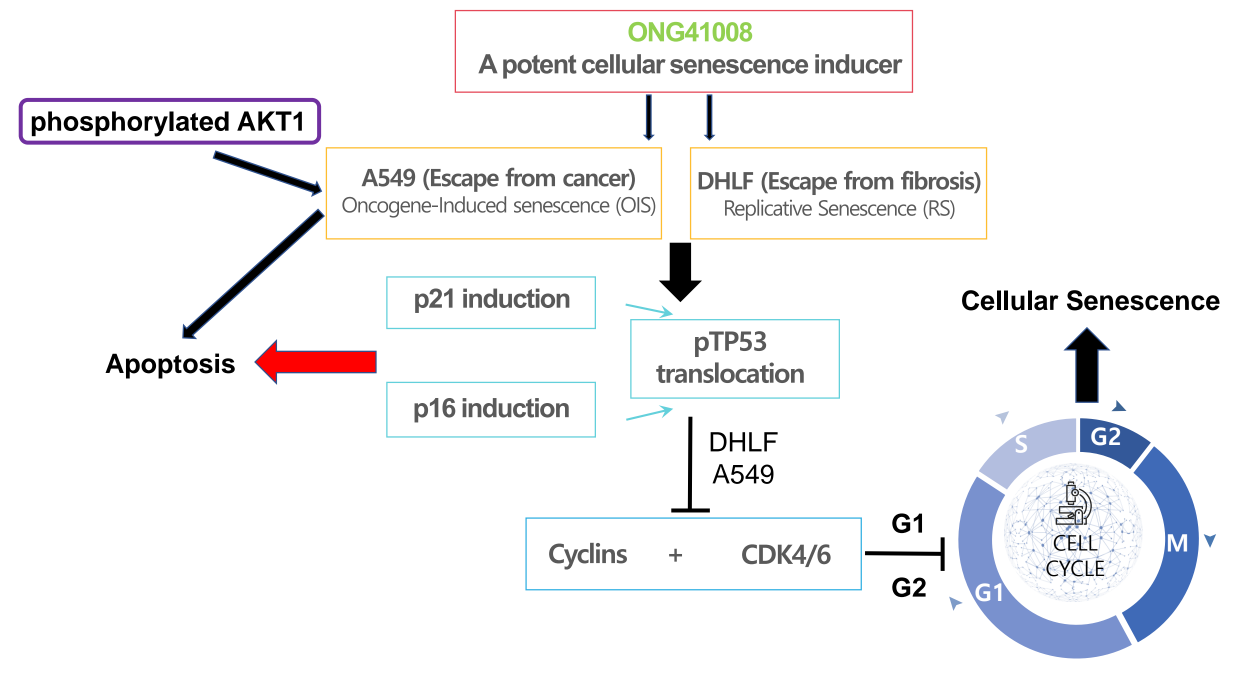


Figure 10