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<tr>
<td><strong>Citation</strong></td>
<td>PLoS One, 2013, v. 8 n. 10, article no. e78395</td>
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<td><strong>Issued Date</strong></td>
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Efficient Immortalization of Primary Nasopharyngeal Epithelial Cells for EBV Infection Study

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Abstract

Nasopharyngeal carcinoma (NPC) is common among southern Chinese including the ethnic Cantonese population living in Hong Kong. Epstein-Barr virus (EBV) infection is detected in all undifferentiated type of NPC in this endemic region. Establishment of stable and latent EBV infection in premalignant nasopharyngeal epithelial cells is an early event in NPC development and may contribute to its pathogenesis. Immortalized primary nasopharyngeal epithelial cells represent an important tool for investigation of EBV infection and its tumorigenic potential in this special type of epithelial cells. However, the limited availability and small sizes of nasopharyngeal biopsies have seriously restricted the establishment of primary nasopharyngeal epithelial cells for immortalization. A reliable and effective method to immortalize primary nasopharyngeal epithelial cells will provide unrestricted materials for EBV infection studies. An earlier study has reported that Bmi-1 expression could immortalize primary nasopharyngeal epithelial cells. However, its efficiency and actions in immortalization have not been fully characterized. Our studies showed that Bmi-1 expression alone has limited ability to immortalize primary nasopharyngeal epithelial cells and additional events are often required for its immortalization action. We have identified some of the key events associated with the immortalization of primary nasopharyngeal epithelial cells. Efficient immortalization of nasopharyngeal epithelial cells could be reproducibly and efficiently achieved by the combined actions of Bmi-1 expression, activation of telomerase and silencing of p16 gene. Activation of MAPK signaling and gene expression downstream of Bmi-1 were detected in the immortalized nasopharyngeal epithelial cells and may play a role in immortalization. Furthermore, these newly immortalized nasopharyngeal epithelial cells are susceptible to EBV infection and supported a type II latent EBV infection program characteristic of EBV-infected nasopharyngeal carcinoma. The establishment of an efficient method to immortalize primary nasopharyngeal epithelial cells will facilitate the investigation into the role of EBV infection in pathogenesis of nasopharyngeal carcinoma.

Introduction

Nasopharyngeal carcinoma (NPC) is a common cancer among southern Chinese. It is closely associated with Epstein-Barr virus (EBV) infection [1]. Immortalized nasopharyngeal epithelial (NPE) cells generated from high risk population (Cantonese) will be valuable tools to study EBV infection and its role in the NPC pathogenesis. Access to non-malignant NPE tissues is extremely limited and surgically biopsied nasopharyngeal tissues are small in size; hence presenting tremendous challenges to establish immortalized NPE cells for EBV infection study. Establishment of an efficient and reliable method to immortalize primary NPE cells will greatly facilitate research study in NPC. Viral oncoproteins, notably SV40T and combined action of E6 and E7 from high risk HPV (type 16 and 18), have been commonly used in cell immortalization. In combination with telomerase, high efficiency of immortalization could be achieved. The viral oncogenes could effectively inactivate G1/S cell cycle checkpoint through inactivation of p53 and Rb proteins, releasing cells to progress into cell cycle.
The expression of human telomerase reverse transcriptase (hTert) further compensates the continuous erosion of telomere in dividing cells to prevent onset of cellular senescence; and in combination with either SV40T or HPV16E6/E7 could effectively immortalize many types of human cells. Our laboratory has previously achieved immortalization of NPE cells using either SV40T or HPV16 E6/E7 alone [2]. The process of immortalization was long and the success rate was low. Furthermore, neither SV40 nor HPV has been implicated in the pathogenesis of NPC. The presence of these viral oncogenes may interfere with the actions of EBV encoded products and limit their applications for study of EBV infection in NPC pathogenesis. Immortalization of NPE cells has been achieved by expression of hTert alone but occurred at a very low efficiency [3]. A more efficient and reliable method to immortalize NPE cells remains to be sought.

Bmi-1 represents a good choice for immortalization of primary NPE cells. It is commonly overexpressed in NPC and could be detected in 38.7% of NPC biopsies [4]. Hence, NPE cells immortalized by Bmi-1 will be more representative cell model for EBV infection study. While the immortalization ability of Bmi-1 in primary NPE cells has been demonstrated in an earlier study [4], detailed examination of events associated with the immortalization of NPE cells by Bmi-1 have not been characterized. In this study, we have examined in details the efficiency of Bmi-1 to immortalize primary NPE cells and have characterized some of the crucial events underlying its immortalization action. An efficient method to immortalize primary NPE cells by combined actions of Bmi-1, hTert and silencing of p16 was also established. Furthermore, primary NPE cells immortalized by this protocol were shown to be susceptible to EBV infection which will facilitate their applications in the study of EBV infection in NPC pathogenesis.

The Bmi-1 belongs to the polycomb group family [5] which remodels chromatin protein and deregulates genes commonly involved in carcinogenesis [6,7]. In earlier reports, Bmi-1 was reported to suppress the expression of p16 and p14 at the INK4A locus to overcome cellular senescence [8]. Downregulation of Bmi-1 suppressed proliferation of lymphoma cells. Telomerase activity was detected in cells overexpressing Bmi-1 [9]. The abilities to inhibit p16 and activate telomerase are considered to be the two major properties of Bmi-1 to immortalize human epithelial cells. However, the immortalization actions of Bmi-1 appear to differ among human epithelial cells from different tissue origins. While weak telomerase activity was reported in human mammary epithelial cells upon introduction of Bmi-1, telomerase activity was not commonly detected in dermal keratinocytes and small airway epithelial cells [10]. Suppression of p16 expression was observed when Bmi-1 was overexpressed in human mammary epithelial cells and was attributed for the extension of in vitro lifespan of human mammary epithelial cells [9,10]. However, another study showed that Bmi-1 expression in oral keratinocytes effectively extended in vitro lifespan without significant reduction of p16 expression [11]. Events underlying Bmi-1 immortalization of primary NPE cells are largely undefined. Characterization of these events will facilitate the efficient immortalization of this special type of epithelial cells for NPC study.

Materials and Methods

Cell culture and cell lines

Primary NPE cells (NP105, NP361, NP446 and NP550) were established by explanting small size biopsies of non-malignant nasopharyngeal tissues from patients admitted to the Queen Mary Hospital, University of Hong Kong, Hong Kong. Prior written informed consents for the use of these tissues for research investigation were obtained from patients or guardians on the behalf of the children participants. The collection and use of these specimens have been approved by the Human Research Ethic Committee of the University of Hong Kong. The biopsy of NP105 was taken from tonsil of a 4-year-old female and the tissue of NP550 was taken from the right nasopharyngeal region of a 76-year-old male with NPC and was examined to be tumor free before use for explant culture. Patients information of NP361 and NP446 biopsies have been described [12]. Details of the culture methods have been previously described [2,3]. The NPE2 Bmi-1 cell was an immortalized NPE cell line and the detail has been published [4]. Akata-EBV and a NPC cell line (C666-1) were cultured in RPMI-1640 medium (Sigma, St. Louis, MO) supplemented with 10% fetal bovine serum (GIBCO, Invitrogen, Carlsbad, CA), 100 U/ml penicillin and 100 μg/ml streptomycin. All the cells were maintained in a 37 °C incubator with 5% CO2 in air.

Expression of Bmi-1 and hTert in NPE cells

 Infective retrovirus was prepared according to previous publications [3,4,13]. The cells after transduction with Bmi-1 were selected with 100 ng/ml puromycin for 2 weeks. The expression of Bmi-1 was confirmed by Western blot analysis. The human telomerase reverse transcriptase (hTert) gene was then transduced into the Bmi-1-expressing cells. Details for the transduction of hTert and detection of telomerase activity were previously described [3].

Immunofluorescence staining for cytokeratin

Immunofluorescent staining of acidic cytokeratin (clone AE1; Zymed Laboratories, Inc.) and basic cytokeratin (clone AE3; Zymed Laboratories, Inc.) was performed as previously described [12].

Cytogenetic analysis of the immortalized NPE cells

Spectral karyotyping was performed as previously described [14].

Flow cytometric analysis

Cells were fixed in ice-cooled 70% ethanol. After replacing with phosphate buffered saline, the suspension was incubated with 10 μg propidium iodide and 10 μg RNase A at 37°C for 20 minutes. The flow cytometry data was acquired using FACS Cantoll flow cytometer (BD Biosciences, San Jose, CA, USA) and the results were analyzed by ModFit LT2.0 software (Coulter Electronics). EBV infection rate in the immortalized
NPE cells was determined based on the GFP expression after 2 days infection. Cell suspension was analyzed by LSR Fortessa Analyzer (BD Biosciences).

Western blotting analysis
Details procedures of Western blot analysis have been published previously [15]. The primary antibodies used were: actin (I-19; 1:1000; Santa Cruz Biotechnology, Santa Cruz, CA), anti-vimentin (VIM 3B4; 1:1000; Boehringer Mannheim, Germany), Bmi-1 (F6; 1:200; Millipore), Cytokeratin 8/18 (LP34; 1:1000; Novocastra Laboratories, Newcastle, UK), Cytokeratin 8 (4.1.18; 1:1000; Chemicon, Temecula, CA), anti-vimentin (VIM 3B4; 1:1000; Boehringer Mannheim, Germany), Cytokeratin 13 (Ks 13.1; 1:1000; Chemicon), Cytokeratin 19 (b170; 1:1000; Novocastra Laboratories), EGFR (1005; 1:500; Santa Cruz Biotechnology), p16 INK4A (1:500; Cell Signaling Technology, Danvers, MA), p21 (F-5; 1:500; Santa Cruz Biotechnology), p16 INK4A (1:500; Cell Signaling Technology, Danvers, MA), anti-vimentin (VIM 3B4; 1:1000; Boehringer Mannheim, Germany), Bmi-1 (F6; 1:200; Millipore), Cytokeratin 5/6/18 (LP34; 1:1000; Novocastra Laboratories, Newcastle, UK), Cytokeratin 8 (4.1.18; 1:1000; Chemicon, Temecula, CA), anti-vimentin (VIM 3B4; 1:1000; Boehringer Mannheim, Germany), Cytokeratin 13 (Ks 13.1; 1:1000; Chemicon), Cytokeratin 19 (b170; 1:1000; Novocastra Laboratories), EGFR (1005; 1:500; Santa Cruz Biotechnology), p16 INK4A (1:500; Cell Signaling Technology, Danvers, MA), p21 (F-5; 1:500; Santa Cruz Biotechnology), p53 (DO-7; 1:2000; Dako, Denmark), phospho-p44/42 MAPK (Erk1/2) (Thr202/Tyr204) (1:500; Cell Signaling Technology) and phospho-MEK1/2 (Ser217/221) (1:500; Cell Signaling Technology).

RNA extraction and RT-PCR analysis
Total RNA was extracted from the culturing cells at their exponential growth phase, reversely transcribed to cDNA and subjected to PCR analysis as previously described [13]. The primers sequences for the genes in Bmi-1 driven pathways and for the EBV-encoded transcripts have been published [16,17].

Bio-Plex phosphoprotein assay
Quantitative detection of phosphorylated protein of Akt (Ser473), JNK (Thr183/Tyr185), MEK1 (Ser217/Ser221), NFκB p65 (Ser536), p38 MAPK (Thr180/Tyr182), Stat3 (Tyr705) and Stat6 (Tyr407) in cell lysate was performed using a bead-based Bio-Plex phosphoprotein assay kit (Bio-Rad Laboratories, Hercules, CA). Details procedures have been described [12].

Knocking down p16 in NPE cells
Lentiviral vector for silencing of p16 (pRRL.SIN-18shp16) was kindly provided by Prof. Judith Campisi at the University of California, USA. Lentivirus was prepared by cotransfecting p16 silencing vector with a lentivirus packaging vector mix (Invitrogen) into the 293T packaging cell line. The lentivirus-containing supernatant and hexadimethrine bromide (4 μg/ml) (Sigma) were added to the NPE cells. The silencing of p16 was confirmed by Western blot analysis.

EBV infection of NPE cells
EBV infection of immortalized NPE cells was achieved by either coculturing of NPE cells with lytically-induced EBV-infected Akata cells or incubating the NPE cells with cell-free EBV supernatant. The experimental procedures for the coculture with Akata cells have been described previously [18]. For the cell-free EBV infection, supernatant from the lytically-induced Akata cells was harvested. Epithelial cells were incubated with the supernatant with centrifugation.

Results

Efficiency of Bmi-1 to immortalize primary NPE cells
We have examined the immortalization property of Bmi-1 in primary cultures of NPE cells. The impact of Bmi-1 on the cellular and molecular properties of the immortalized NPE cells and the susceptibility of Bmi-1-immortalized cells to EBV infection were characterized. Primary NPE cell cultures used in this immortalization study were established from non-malignant nasopharyngeal biopsies obtained from NPC patients and tonsillectomies (Table S1 in File S1). The course and events during immortalization of four primary cultures of non-malignant nasopharyngeal cultures were carefully examined. Two of these primary cultures, NP361 and NP550, were established from non-malignant nasopharyngeal epithelial biopsies from NPC patients. The remaining two primary cultures, NP105 and NP446, were established from tonsillectomized tissues from non-NPC patients. All these nasopharyngeal biopsies were confirmed by histopathological examination to be tumor free. Spectral karyotyping analysis in primary nasopharyngeal epithelial cells from these biopsies revealed normal and diploid karyotype (46 chromosomes) with no cytogenetic aberrations (data not shown).

Similar to primary cultures of other human epithelial cells, these non-malignant NPE cells all underwent cellular senescence as evidenced by their enlarged cell morphology and expression of senescence-associated beta galactosidase (data not shown). The number of passages that could be achieved before onset of cellular senescence varied among the four primary NPE cultures (Table S1 in File S1). The two primary NPE cultures established from non-malignant nasopharyngeal biopsies obtained from NPC patients (NP361 and NP550) could be propagated for a longer passage (PD 20 and PD 24 respectively) before onset of cellular senescence while the primary NPE cells derived from non-NPC patients (NP105 and NP446) underwent cellular senescence at a much shorter passage in culture (PD 6 and PD 10 respectively) (Table S1 in File S1). This may indicate the presence of occult genetic alterations in tumor adjacent nasopharyngeal epithelial tissues supporting a longer in vitro lifespan. A larger sample size will be warranted for this observation to be conclusive.

Bmi-1 was ectopically expressed in these four non-malignant NPE cell cultures and examined for its ability to immortalize them (Figure 1A). In all the four primary NPE cultures, expression of Bmi-1 successfully delayed their onset of cellular senescence and markedly extended their in vitro lifespans.

EBV infection of immortalized NPE cells
EBV infection of immortalized NPE cells was achieved by either coculturing of NPE cells with lytically-induced EBV-infected Akata cells or incubating the NPE cells with cell-free EBV supernatant. The experimental procedures for the coculture with Akata cells have been described previously [18]. For the cell-free EBV infection, supernatant from the lytically-induced Akata cells was harvested. Epithelial cells were incubated with the supernatant with centrifugation.

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which has previously been reported by our group to have ability to immortalize human NPE cells in combination with telomerase [12]. Overexpression of cyclin D1 or CDK4, in combination with telomerase and other genetic alterations such as EGFR, has been reported by others to immortalize oral keratinocytes [19]. The CDK4 used in this study is a mutant form (CDK4R24C), harboring a mutation of arginine to cysteine at codon 24, rendering the cells to be insensitive to growth inhibitory action of p16. Compared to these genetic elements, Bmi-1 was shown to be the most efficient in extending the in vitro lifespan of primary NPE cell cultures (Figure 1B and Table S2 in File S1).

Immortalization of primary NPE cells by combined actions of Bmi-1 and hTert

We then sought for the additional crucial elements required for immortalization of primary NPE cells by Bmi-1. Telomerase activation is critical for cell immortalization. The ability of Bmi-1 in combination with hTert (catalytic unit of telomerase) to immortalize primary cultures of NPE cells was then examined. Our previous studies have showed that immortalization of NPE cells by telomerase alone is a rare event and additional genetic alterations are involved [3,12]. In this study, expression of telomerase alone also failed to immortalize these four primary NPE cell cultures despite the successful activation of telomerase by hTert expression (Fig. S1). In combination, Bmi-1 and hTert successfully immortalized two (NP446 and NP550) out of the four primary cultures of NPE cells examined (Figure 2A). The lifespan of the remaining two primary NPE cell

Figure 1. Growth history of primary nasopharyngeal epithelial cells. (A) Immortalization history of primary nasopharyngeal epithelial cells by Bmi-1. NP105, NP361, NP446 and NP550 stopped proliferation at population doublings 6 to 24. The cells were subjected to immortalization by transduction with Bmi-1. The introduction of Bmi-1 could extend the life span of the cells. (B) Expression of genetic elements in primary nasopharyngeal epithelial cells. Genetic elements, Bmi-1, LMP1, cyclin D1 and p16-insensitive form of CDK4, were expressed in primary nasopharyngeal epithelial cells. Among them, Bmi-1 was the most efficient in extension of lifespan of cells. (C) Ectopic expression of Bmi-1, LMP1, cyclin D1 and CDK4 in the primary nasopharyngeal epithelial cells. The cells could maintain the expression of the exogenous genes until the late stage as confirmed by Western blot.

doi: 10.1371/journal.pone.0078395.g001
cultures (NP105 and NP361) was also markedly extended by the combined action of Bmi-1 and hTert (Figure 2A).

Characterization of NPE cells immortalized by Bmi-1 and hTert

The properties of these two newly immortalized NPE cells are described here. The Bmi-1/hTert-immortalized NPE cell lines (NP446 and NP550) are non-tumorigenic when injected subcutaneously to BALB/cAnN-nu (Nude) mice at 10^6 to 10^7 cells per mouse and observed for 3 to 6 months period. Under phase contrast microscope, the Bmi-1/hTert-immortalized cells appeared smaller in size compared to their parental cells (Figure 2B). The epithelial nature of the immortalized NP446 and NP550 cells was confirmed by the presence of keratin revealed by immunocytochemical staining using keratin-specific antibodies (AE1 and AE3) (Figure 2C). Western blotting analysis of keratin profiles of Bmi-1/hTert-immortalized NPE cells revealed the presence of keratins 5, 6, 8, 13, 18 and 19 (Figure 2D) which are the common keratin profile expressed in primary NPE cells [2].

Ability of Bmi-1 to induce telomerase activity in human NPE cells

Previous study has reported that Bmi-1 could induce telomerase activity in human mammary epithelial cells to facilitate cell immortalization [9]. We then examined if Bmi-1 could activate telomerase activity in our NPE cells by the sensitive TRAP assay (Figure 2E). Telomerase activity was not

Figure 2. Properties of Bmi-1/hTert-immortalized cells. (A) Immortalization of nasopharyngeal epithelial cells by expression of Bmi-1 and activation of telomerase. Immortalization could be observed in NP446 and NP550 cell lines by expression of Bmi-1 together with hTert. (B) Microscopic images of normal human nasopharyngeal epithelial cells and immortalized nasopharyngeal epithelial cells (240x). Normal cells of NP446 and NP550, at PD 7 and 15 respectively. They were still proliferative. The normal cells after introduction of Bmi-1 alone still became enlarged and stopped proliferating at later passage. The introduction of Bmi-1 together with hTert to the normal cells extended their life span in culture and the cells were immortalized. (C-D) The epithelial origins of the cells are confirmed by expression of cytokeratin. (C) Immunocytochemistry staining of cytokeratin. (D) Western blot confirmed expression of cytokeratin. (E) Telomerase activity by TRAP assay. Successful induction of telomerase activity in immortalized cells after introduction of hTert. Bmi-1 alone could also induce the telomerase activity in NP550 cell line. However, a relatively low telomerase activity could be observed in the cells after the introduction of Bmi-1 alone or the cells from primary culture.

doi: 10.1371/journal.pone.0078395.g002
detectable in NP446 cells after expression of Bmi-1 (NP446Bmi-1). A low level of telomerase activity was however detected in NP550 cells after expression of Bmi-1 (NP550Bmi-1). However, the low level of telomerase detected in NP550Bmi-1 cells was not sufficient to achieve immortalization as NP550Bmi-1 continued to undergo cellular senescence. Telomerase activity was also not detected in NP105 and NP361 cells expressing Bmi-1 alone (data not shown). In contrast, robust telomerase activity was detected in the Bmi-1/hTert-immortalized NPE cells (NP446Bmi-1/hTert and NP550Bmi-1/hTert cells). We can conclude that Bmi-1 expression is not efficient to activate telomerase in primary NPE cells to achieve immortalization.

Karyotypic analysis of the Bmi-1/hTert-immortalized NPE cells

Detailed examination by spectral karyotyping analysis reveals the presence of some chromosomal abnormalities in the two Bmi-1/hTert-immortalized NPE cells. The abnormalities are mostly numerical in nature. The karyotypes of NP446Bmi-1/hTert cells were largely nearly diploid with the gain of chromosome 5 (Figure 3A). Interestingly, the NP550Bmi-1/hTert cells revealed a near tetraploid karyotype which was also confirmed by flow cytometric analysis of DNA content (Figure 3B). Nearly the whole set of chromosomes was duplicated in the immortalized NP550Bmi-1/hTert cells. In addition, there were a few other structural chromosomal changes observed, particularly involving chromosome 20. These structural changes may occur after tetraploidization as they were not detected in early passages. Detailed examination of DNA content by flow cytometry also revealed normal bimodal distribution of DNA content at G1/S and G2/M phase of primary NP550 cells (Figure 3B). At PD 27, an additional DNA peak could be detected at 8N DNA content revealing the emergence of a G2/M tetraploid population (8.64%) of NP550Bmi-1/hTert cells (Figure 3B). At PD 85, the entire population of NP550Bmi-1/hTert cells revealed a 4N DNA population (76.04%) at G1 phase and 8N DNA population (10.54%) at G2/M phase, with 0% of cells with 2N DNA content at G1 phase, indicating completely takeover of cell population by the tetraploidy of Bmi-1/hTert-immortalized NP550 cells. The underlying events leading to this switching in ploidy in NP550Bmi-1/hTert cells are not completely understood. One speculation is that the tetraploidization of NP550Bmi-1/hTert may confer growth advantage to NP550Bmi-1/hTert cells and facilitated their immortalization.

Alteration of gene expression in Bmi-1/hTert-immortalized cells

We further examined the expression level of p16, p53 and p21 during the course of immortalization of NP446Bmi-1/hTert and NP550Bmi-1/hTert. Previous studies have reported that Bmi-1 could suppress p16 expression in immortalized cells. In both NP446 and NP550 cells, expression of Bmi-1 alone or in combination with hTert did not reveal an immediate suppression of p16 expression (Figure 4A). We were only able to observe decrease of p16 expression at a later stage of immortalization. In NP446, decrease of p16 expression was only observed at PD 51 which was 30 passages after transduction of Bmi-1. The p16 levels fluctuated to some extents in NP550Bmi-1/hTert cells during the course of immortalization. Consistent downregulation of p16 was only observed at a much later passage (after PD 81). It could be concluded that suppression of p16 offers selective growth advantage to Bmi-1-expressing cells which facilitates their immortalization. The selective growth advantage of p16 silencing was also observed in our previous study using telomerase alone to immortalize primary NPE cells [3]. We did not observe obvious changes or trends of p53 expression during the course of immortalization in NP446 and NP550 cells (Figure 4A). In contrast, downregulation of p21 was observed during the course of immortalization of NP446 and NP550 cells.

Upregulation of MAPK pathway in the Bmi-1/hTert-immortalized NPE cells

We then examined if specific signaling pathways are selected during immortalization of NPE cells by Bmi-1 and hTert using the Bioplex phospho-protein detection assay. The signaling pathways examined included Akt, JNK, MAPK, NFκB, p38 MAPK, Stat3 and Stat6. The most prominent pathways activated in the immortalized NP446Bmi-1/hTert and NP550Bmi-1/hTert was MEK/MAPK signaling (Figure 4B). Activation of MAPK signaling in Bmi-1/hTert-immortalized NP446 and NP550 cells was further confirmed by Western blotting (Figure 4C). Notably, activation of MAPK signaling in NP446 and NP550 cells could be confirmed by Western blotting using specific phosphorylated antibodies for activated MEK1/2 and ERK1/2 at early passages, before decrease of p16 expression. We also observed a moderate increase in EGFR expression in both immortalized cells suggesting a selective advantage of MAPK activation for Bmi-1/hTert-immortalized cells. In NP446, activation of MAPK was not observed immediately after Bmi-1 expression indicating that activation of MAPK signaling is not a direct action of Bmi-1 or hTert and may represent a selected phenotype in Bmi-1 and hTert immortalized cells. Our previous study has also shown the overexpression of EGFR and MAPK signaling during immortalization of NPE cells by LMP1 and telomerase which is supportive of a role of MAPK activation in immortalization of primary NPE cells [12].

Upregulation of Bmi-1 driven gene expression in immortalized NPE cells

Bmi-1 has been shown to be involved in the maintenance of stem cells in tissues [20]. To confirm the functional activation of Bmi-1 after expression in the NPE cells, we have examined the expression status of the multiple genes downstream of Bmi-1 activation. The expression of these genes have also been reported in Bmi-1-driven stem cell development pathway. The expression of these Bmi-1-driven genes at different passages of immortalized cells was examined by semi-quantitative RT-PCR analysis. In contrast to the cell cycle regulatory genes and MAPK signaling, we observed immediate activation of many of these Bmi-1 downstream gene targets in immortalized NP446Bmi-1/hTert and NP550Bmi-1/hTert cells (Figure 5).
Expression of Bmi-1 alone could also upregulate many of these genes in NPE cells before immortalization which may contribute to the extension of lifespan by Bmi-1. Interestingly, the expression levels of some genes appeared to be enhanced and stabilized after hTert expression.

Essential role of p16 inactivation in the immortalization of NPE cells by Bmi-1 and hTert

Our results showed that Bmi-1 could effectively extend the lifespan of primary NPE cells and inhibit onset of cellular senescence. In combination with telomerase activation, the ability to extend the lifespan of NPE cells was even more prominent. Nonetheless, expression of Bmi-1 and hTert was only successful in immortalizing two out of four primary cultures of NPE cells. Additional events are required to achieve efficient immortalization of NPE cells. Examination of these NPE cells revealed that p16 silencing may be a crucial event for immortalization of NPE cells by Bmi-1 and hTert. In both NP446 and NP550 cells immortalized by Bmi-1 and hTert, we observed suppression of p16 at late passages. The downregulation of p16 may not be the direct action of Bmi-1 as it was only observed at late passages. The role of p16 in the onset of cellular senescence is well recognized. A consistently high level of p16 was observed in NP105 and NP361 cells which failed to be immortalized despite the effective expression of Bmi-1 and activation of telomerase (Figure 6A). We proceeded to examine if knocking down p16 expression in NP105 and NP361 cells using a lentiviral plasmid expressing p16 led to efficient immortalization.

Figure 3. Karyotypic analysis of the immortalized NPE cells. (A) Spectral karyotyping of the major clones in the Bmi-1 immortalized cells. Majority of clones in NP446Bmi-1/hTert are quite normal and only contained one extra chromosome 5. NP550Bmi-1/hTert are nearly-tetraploid. (B) Flow cytometry analysis of normal NP550 cells and NP550Bmi-1/hTert immortalized cells at different stage of immortalization. Cell cycle distribution of the nasopharyngeal epithelial cells derived from primary culture and different passages of the NP550Bmi-1/hTert immortalized cells revealed by flow cytometry. A switch to tetraploid could be detected after expression of Bmi-1 in NP550. In NP550Bmi-1/hTert, a switch to tetraploid could be detected as early as at PD 27.

doi: 10.1371/journal.pone.0078395.g003
Figure 4. Alteration of gene expression in immortalized NPE cells. (A) Expression of genes involved in G1/S cell cycle entry. p16 and p21 were downregulated in the later passage of immortalized cells. (B-C) Activation of EGFR-MAPK signaling in immortalized cells. (B) Upregulation of phospho-MEK1 but not the other phosphorylated protein was detected in the immortalized cells compared to the control. Means and SDs were calculated from triplicate wells of two independent experiments. ***P < 0.005 and P < 0.001, Student t test. (C) The levels of EGFR, phospho-MEK1/2 and phospho-ERK1/2 expressions in cells were quantitated by ImageJ. Control expression in primary cells was set as 1. A gradual increase in EGFR and its downstream signaling, the phospho-MEK1/2 and phospho-ERK1/2, could be observed during immortalization of NP446Bmi-1/hTert and NP550Bmi-1/hTert.

doi: 10.1371/journal.pone.0078395.g004
shRNA against p16 might result in immortalization of these two NPE lines. Indeed, both NP105Bmi-1/hTert and NP361Bmi-1/hTert cells resumed their proliferation upon effective knock-down of p16 and achieved immortalization (Figures 6B, 6C and 6D). Hence we have effectively immortalized four out of four primary NPE cells using combination of Bmi-1 overexpression, hTert activation and p16 silencing.

EBV infection of the Bmi-1/hTert-immortalized cells

EBV infection has long been postulated to play a critical role in NPC pathogenesis. The lack of representative cell systems for EBV infection has hindered the progress in the understanding of the role of EBV in NPC pathogenesis. A detailed examination of the application of our immortalized NPE cells for EBV infection will be beyond the scope of this current study. It is essential to examine if the NPE cell lines immortalized by Bmi-1, hTert and silencing of p16 are susceptible for EBV infection. We have previously reported the successful EBV infection in a telomerase-immortalized NPE cell line [17]. Infection of immortalized NPE cells by EBV could be achieved by co-culturing with EBV-producing Akata cells. Alternatively, EBV infection could also be achieved using concentrated viral supernatant harvested from EBV-producing Akata cells. The EBV used to infect the NPE cells was GFP-tagged which facilitated the identification of EBV-infected cells by fluorescent imaging. Immortalized NPE cells were infected with EBV by both co-culture and direct cell free infection (Figure 7A and Table 1). Our results showed that the immortalized NPE cells could be effectively infected by either method. We were able to detect expression of multiple transcripts of EBV gene (EBER1, EBNA1, LMP1 and LMP2) in all the immortalized NPE cells established in this study (Figure 7B). The expression of EBV lytic genes (BZLF1 and BRLF1) was low or undetectable in our EBV-infected immortalized NPE cells. The profiles of EBV gene expression detected in infected NPE cells were representative of type II latent EBV infection which is characteristic of EBV

Figure 5. Expression of genes involved in Bmi-1 driven stemness pathway. All the stem cell-ness genes in the Bmi-1 driven pathway were upregulated in the immortalized cells and could be detected at both early and later passages of immortalization.

doi: 10.1371/journal.pone.0078395.g005
infection in NPC. The dynamics of EBV gene expression in freshly infected and stably infected NPE cells are under current investigation.

Discussion

Overexpression of \textit{Bmi-1} is common in many types of human cancer [5,9,21], including nasopharyngeal carcinoma [4]. Overexpression of \textit{Bmi-1} has been reported in over one-third of NPC. \textit{Bmi-1} immortalized NPE cells may provide representative cell models for study of NPC pathogenesis. The immortalization function of \textit{Bmi-1} has been previously reported in multiple cell types. Earlier studies have reported that expression of \textit{Bmi-1} alone could immortalize fibroblasts [22], human mammary epithelial cells [9] and human nasopharyngeal epithelial cells [4]. However, the immortalization action of \textit{Bmi-1} appears to vary among cell lines from different origins [10]. Most importantly, expression of \textit{Bmi-1} was often observed to be insufficient to immortalize primary human epithelial cells [10,23]. In this study, we also observed that expression of \textit{Bmi-1} alone failed to immortalize primary human NPE cells and additional events are required. A recent study indicated that additional events such as inhibition of TGF-\(\beta\) signaling may also be involved in the extension of life span of primary epithelial cells after \textit{Bmi-1} expression [23].

In this study, the ability of \textit{Bmi-1} to extend lifespan was confirmed in primary NPE cells supporting its role in NPC pathogenesis reported in our earlier study [4]. The underlying mechanisms involved in the extension of lifespan of primary

![Figure 6. Immortalization of NP105 and NP361 cells by knocking down p16.](image_url)

A) P16 expression in NP105 and NP361 cells. Primary cells (NP550) and cells that cannot be immortalized by Bmi-1/hTert (NP105 and NP361) expressed a high level of p16. (B-D) Knocking down p16 in NP105 and NP361 cells could lead to immortalization of both cell lines.

doi: 10.1371/journal.pone.0078395.g006
epithelial cells by Bmi-1 are not well characterized. Most studies attributed this property to the suppression action of p16 by Bmi-1 [9,10]. However, in this study, effective suppression of p16 expression was not observed after expression of Bmi-1.

Additional events may be involved in the extension of lifespan in Bmi-1 expression cells.

To achieve immortalization, telomerase activation is essential. In earlier studies, the activation of telomerase was reported to be induced by Bmi-1 expression during immortalization of epithelial cells [8,9,11]. A low level of telomerase was observed in human mammary epithelial cells after expression of Bmi-1 but was not sufficient for immortalization. The ability of Bmi-1 to activate telomerase during immortalization may be cell-type dependent. We were not able to detect significant telomerase activation in all our primary NPE cells examined. Similarly, telomerase activation was also not observed in skin keratinocytes and small airway epithelial cells [10]. In oral keratinocytes immortalized by HPV16 E6 and Bmi-1, telomerase activation was observed only after the cells have emerged from telomere erosion-induced crisis[11]. In our study, telomerase activation by hTert was required for efficient immortalization of primary NPE cells.

Expression of Bmi-1 and hTert was only successful in the immortalization of two primary NPE cells (NP446 and NP550) out of four examined (50%). Characterization of NP446 and NP550 cells immortalized by Bmi-1 and hTert reviewed additional events may be involved during immortalization. One of the biological properties of Bmi-1 is self-renewal and maintenance of human stem cells [5,24]. A comparative genomics approach has revealed genes driven by Bmi-1 which have been reported to be involved in maintenance of the stem cell properties [16]. In this study, we observed effective and immediate activation of Bmi-1-targeted genes after expression

### Table 1. All the Bmi-1/hTert-immortalized NPE cells could be effectively infected by EBV.

<table>
<thead>
<tr>
<th></th>
<th>EBV infection rate (% GFP positive)</th>
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<tbody>
<tr>
<td></td>
<td>Co-culture infection</td>
</tr>
<tr>
<td>NP446Bmi-1/hTert</td>
<td>19.6</td>
</tr>
<tr>
<td>NP550Bmi-1/hTert</td>
<td>54.7</td>
</tr>
<tr>
<td>NP105Bmi-1/hTert/sh-p16</td>
<td>23.5</td>
</tr>
<tr>
<td>NP361Bmi-1/hTert/sh-p16</td>
<td>36.8</td>
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</table>

Both co-culture and direct cell free EBV infection could be achieved in the immortalized cells.

doi: 10.1371/journal.pone.0078395.t001
of Bmi-1 in primary NPE cells, prior to suppression of p16 expression. Their involvement in immortalization of NPE cells warrants further investigations. Furthermore, downregulation of the p21 protein and activation of MAPK signaling were observed in both Bmi-1/hTert-immortalized NP446 and NP550 cells suggesting their involvement in immortalization of NPE cells. Interestingly, we observed a change of ploidy of NP550 cells during immortalization by Bmi-1 and hTert. The ploidy of primary NP550 cells, which was largely diploid, drifted to become near-tetraploid after immortalization by Bmi-1 and hTert. The exact mechanism involved in the change of ploidy is under investigation and may involve activation of aurora kinase A (unpublished observation). Expression of Bmi-1 appears to provide a selective growth advantage of these near-tetraploid NP550 cells. Aberrant mitosis leading to gross chromosomal rearrangement and ploidy changes may induce cell cycle arrest and apoptosis. Bmi-1 expression has been shown to promote survival of cells to chemotherapeutic agents [25] and plays a role in DNA damage repair. Expression of Bmi-1 may promote survival of cells harboring aberrant chromosomal changes, which otherwise may trigger cellular apoptosis induced by mitotic abnormalities. Further investigations are warranted to determine if Bmi-1 expression may promote survival of cells harboring genetic alterations.

The examination of p16 expression during the immortalization of primary NPE cells by Bmi-1 reveals a complex pattern. The ability of Polycomb group proteins to suppress p16 expression for extension of in vitro lifespan of cells and eventually immortalization of fibroblasts has been previously reported [7,8,26]. Bmi-1 has been reported to suppress p16 expression through transcriptional repression [4,8]. The suppression of p16 expression by Bmi-1 may be cell type dependent. Suppression of p16 expression was not prominent in NPE cells after expression of Bmi-1 and was only observed at later passages after expression of Bmi-1 and hTert. In NP446 cells, significant suppression of p16 was only observed after PD 50 and may be a selective event to support Bmi-1 and hTert immortalization of NPE cells. The p16 expression levels in NP550 cells also fluctuated after Bmi-1 and hTert expression. Consistent suppression of p16 expression was only observed at late passages (after PD 81). In our previously reported study, p16 deletion is a critical event selected by immortalized NPE cells after expression of telomerase [3]. Suppression of p16 clearly offers a growth advantage for NPE cells and is actively selected during immortalization. In NP105 and NP361 cells, expression of p16 remained high after ectopic expression of Bmi-1 and hTert, and they failed to become immortalized. Silencing of p16 expression was required to immortalize these two cell lines clearly indicating the importance of p16 inactivation in immortalization of NPE cells.

Bmi-1 expression, telomerase activation and p16 inactivation are all common events present in NPC [27]. The ability of Bmi-1 expression, telomerase activation and p16 silencing to efficiently immortalize primary NPE cultures has enabled us to establish immortalized NPE cells from high risk NPC population for NPC studies. Furthermore, NPE cells immortalized by combined action of Bmi-1, hTert and silencing of p16 are susceptible to EBV infection and enable establishment of type II latent EBV infection characteristic of EBV infection in NPC. The role of EBV in NPC pathogenesis is still enigmatic at this stage. Events relating to infection of EBV and persistent of EBV infection in premalignant nasopharyngeal epithelium remain poorly defined. Efficient immortalization of primary NPE cells susceptible to EBV infection will facilitate investigations of EBV infection in NPC pathogenesis.

Supporting Information

Figure S1. Activation of telomerase in primary nasopharyngeal epithelial cells. Telomerase was activated in primary nasopharyngeal epithelial cells by transduction with hTert. However, immortalization could not be achieved. (TIFF)

File S1. Contains Tables S1 and S2. Table S1: Information of primary nasopharyngeal epithelial cultures. Table S2: Highest population doublings that can be achieved by the primary nasopharyngeal epithelial cells expressing different genetic elements (Bmi-1, LMP1, cyclin D1 and CDK4). Cells expressing Bmi-1 could achieve the highest population doublings compared to those expressing other genetic elements. (DOC)

Acknowledgements

The authors acknowledged the Faculty Core Facility of the Li Ka Shing Faculty of Medicine, University of Hong Kong for providing the equipment for acquisition of flow cytometry data. pLXIN-hTert expression vector was a kind gift from Dr. J. C. Barrett at the National Institute of Environmental Health Sciences, USA.

Author Contributions

Conceived and designed the experiments: YLY CM MZ SWT. Performed the experiments: YLY PSP WD CMT PMH. Analyzed the data: YLY WD SWT. Contributed reagents/materials/analysis tools: MZ YJ APWY. Wrote the manuscript: YLY SWT.
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