Establishment of a Zebrafish Xenograft Model for \textit{in Vivo} Investigation of Nasopharyngeal Carcinoma

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Abstract
Nasopharyngeal carcinoma (NPC) is a unique malignant tumor of the head and neck. Despite higher survival rates by the combination of radiotherapy and chemotherapy, the recurrence or metastasis of NPC still occurs at about 10%. Therefore, there is urgent demand to develop more effective \textit{in vivo} models for preclinical trials to investigate the mechanisms of NPC development and progression and to explore better treatment approaches. In this study, we transplanted human NPC CNE1 cells into zebrafish embryos to establish a xenograft model of NPC, where the proliferation and invasion behaviors of NPC cells were investigated \textit{in vivo}. Combining \textit{in vitro} and \textit{in vivo} analyses, we found that activating transcription factor 7 (ATF7) was involved in the occurrence and development of NPC regulated by peptidyl-prolyl cis-trans isomerase NIMA-interacting 1 (Pin1). The zebrafish NPC xenograft model established here thereby provides an \textit{in vivo} tool for exploring the occurrence and development of NPC, which may help to identify new tumor markers and develop new therapeutic strategies for the treatment of NPC.

Keywords
zebrafish, nasopharyngeal carcinoma, xenograft, Pin1, ATF7

Introduction
Nasopharyngeal carcinoma (NPC) is a unique head and neck malignant tumor, which originates from nasopharyngeal epithelium. It is one of the malignant tumors with high incidence rate in South China and Southeast Asian countries$^{1,2}$. Studies have shown that the occurrence of NPC is related to Epstein-Barr virus (EBV) infection, genetic susceptibility, and environment$^3$. Due to the special anatomical location and high radiosensitivity of NPC, current main treatment method of NPC is to combine radiotherapy and chemotherapy$^{1,4}$. Although it has a higher survival rate, the recurrence or metastasis rate of NPC is about 10% in patients received radiotherapy combined with chemotherapy$^{5,6}$. In addition, chemotherapy resistance is usually accompanied by a poor prognosis, which is also a major obstacle for the treatment of patients with recurrent NPC. Therefore, to improve the survival rate of patients is still a major challenge in the treatment of NPC. There are urgent needs to develop more effective \textit{in vivo} NPC models for preclinical trials, so as to explore the mechanisms involved in NPC occurrence and progression.

Model organisms are important for studying and understanding human diseases. Traditional vertebrate models include mice, rats, and so on. Among them, mouse is the most widely used model to establish xenotransplantation models for evaluating the preclinical efficacy of anticarcinogen$^7$. Despite the increase of various immunodeficient and humanized mouse models, there are still inherent limitations with these models. For example, mice are not suitable for large-scale screening of small molecules due to their high cost, long life cycle, and large breeding space needed. In addition, it is difficult to intravitaly image internal cancers in mice. To complement these shortages in mice models, zebrafish xenograft models were developed. Zebrafish has very conserved genome compared with human and owns highly homologous...
oncogenes and tumor suppressor genes, making it an ideal model for identifying carcinoma-related genes and screening therapeutic compounds. Furthermore, zebrafish has already been shown as a suitable animal model for studying the pathological mechanisms of many diseases due to its availability, operability, transparent body, and low cost. Most importantly, its adaptive immune system does not mature until 4 weeks after fertilization. Xenograft models establishment in zebrafish embryos can avoid immune rejection efficiently. Moreover, by combining the fluorescent labeling of tumor cells, the transparent fish body allows to access and trace the transplanted cells in vivo in single-cell resolutions, and to observe the migration of tumor cells and the formation and metastasis of tumors by imaging time-lapse movies.

In this study, we aimed to establish a zebrafish NPC xenograft model to provide an in vivo system to investigate the migration and invasion of NPC cells and also the formation of NPC in vivo. Meanwhile, the role of the peptidyl-prolyl cis-trans isomerase NIMA-interacting 1 (Pin1) on the NPC formation and on the regulation of the activating transcription factor 7 (ATF7) was investigated using this in vivo model. The results indicated that the zebrafish xenograft model could be an efficient model system for studying the tumorigenesis and the progress of NPC in vivo.

Materials and Methods

Cell Culture and Transfection

The NPC CNE1 cells were cultured in RPMI 1640 medium (Gibco, Thermo Fisher Scientific) supplemented with 10% fetal bovine serum (FBS; Thermo Fisher Scientific, Waltham, MA, USA) at 37°C in a humidified chamber containing 5% CO2. For establishment of CNE1-LV3 and CNE1-shPin1 cells, CNE1 cells were sequentially infected with LV3 and shPin1 lentiviral particles (the lentivirus packaging vector pGLV3/H1/GFP was purchased from GenePharma Co., Ltd, Suzhou, China) using polybrene transfection enhancer (Sigma-Aldrich, Darmstadt, Germany) and screened with 1 μg/ml puromycin. Pin1 shRNA sense: 5′-GCTCAGGCCG AGTGTACTA-3′ and antisense: 5′-TAGTACACTCGGCCCT GAGC-3′. The plasmids used in this experiment were constructed previously in our group. A total of 5 × 104 CNE1 cells were seeded to a 6-well plate 24 h before transfection. Pin1 overexpression vector or negative control vector was mixed together with lipofectamine 2000 for transfection when the cells were at a 60% to 70% confluence. Cells were harvested for protein extraction at 36-48 h after transfection.

Zebrafish Husbandry

Wild-type zebrafish AB and transgenic zebrafish Tg(kdrl: mCherry) were kept according to standard laboratory procedures. Adult zebrafish were raised in a circulatory system at 28.5°C for 14 h of light and 10 h of darkness. For zebrafish mating, sexually mature zebrafish were placed in pairs in the mating tank overnight, and the following day, they begin spawning when the light is on. Embryos were collected, rinsed, and placed in Petri dishes containing embryo water. Embryos were incubated in an incubator at 28.5°C. To prevent pigment formation in larvae, 0.003% (w/v) phenylthiocarbamide/N-phenylthiourea (PTU; Sigma-Aldrich) was added to the embryo water at 24 h post-fertilization (hpf). Handling of zebrafish was complied with the legislation of “Guangdong Laboratory Animal Management Regulations.”

NPC Xenografts and Tumor Cell Growth Quantification

Tumor cells at 85% fusion in the T25 culture flask were dissociated using 0.25% trypsin (Gibco, Thermo Fisher Scientific) and were counted before centrifugation (3,000 rpm, 3 min). Then, the supernatant was removed, and the cells were resuspended with complete medium and collected in a 1.5-ml EP tube with 3 × 105 cells/ml. Tumor cells were transplanted into the perivitelline space around the yolk sac or common cardinal vein (CCV) of 2 days post-fertilization (dpf) zebrafish embryos (300–400 cells/fish) by a microinjection needle (1.0 mm × 0.6 mm) and CellTram Vario oil-pressure microinjector (Eppendorf, Hamburg, Germany); 20 zebrafish embryos were used for transplantation each group and host zebrafish were placed in an incubator at 28.5°C for 12 h, then transferred to a 34°C incubator and imaged at 3, 5, 7, and 9 dpf, respectively. Zebrafish were imaged under Leica fluorescence stereomicroscope (M205FA; Leica, Nussloch, Germany). Tumor volume was determined by quantifying the two-dimensional (2D) image area with ImageJ and the obtained areas were multiplied by the average GFP fluorescence intensity. This method has now become the standard method in the field.

Time-Lapse Imaging of Extravasated Tumor Cells From Vessels

At 24 h post-transplantation (hpt), the larvae were imaged for NPC CNE1 cells migration by the spinning-disk confocal super resolution microscope (IXplore SpinSR; Olympus, Tokyo, Japan). Larvae were anesthetized in embryo water containing 0.02% tricaine (Sigma-Aldrich), fixed with 1% low-melting-point agarose (Thermo Fisher Scientific), and then imaged under 488 and 561 nm lasers for 11 h (every 20 min). At 48 hpt, the xenografted larvae were imaged continuously. Z projection of the confocal serial images was generated by Olympus CellSens software.

Western Blot

Cell proteins were extracted when cells grow to 100% confluence. Protein lysate (100 μl) [1 ml RIPA (radioimmunoprecipitation assay buffer) and 10 μl PMSF (phenylimethanesulfonyl fluoride); Solarbio, Beijing, China] was added to the 60-mm
culture dish (Corning, Darmstadt, Germany). Equal amounts of cellular proteins were separated with 12% SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) and transferred onto PVDF (polyvinylidene difluoride) membranes, which were then blocked with 5% skim milk for 2 h. Then, the PVDF membranes were incubated with primary antibodies overnight at 4°C, washed three times with TBST, followed by horseradish peroxidase (HRP)-conjugated secondary antibodies for 1 h at room temperature, and then washed three times with TBST before imaging. The primary antibodies include rabbit anti-Pin1 (ab192036; Abcam, Cambridge, UK), rabbit anti-ATF7 (ab87844, Abcam), and rabbit anti-Cyclin D1 (ab40754, Abcam). Mouse anti-β-actin (GB12001; Servicebio, Wuhan, China) and mouse anti-GAPDH (GB12002; Servicebio) were served as loading controls. Goat anti-rabbit (HS101-01; TransGen Biotech, Beijing, China) and goat anti-mouse (HS201-01; TransGen Biotech) were used as secondary antibodies. The signal bands were detected with an ECL detection kit (Thermo Fisher Scientific) and a chemiluminescence imaging system (GelView 6000M; BioLight, Xiamen, China), and the results were analyzed by ImageJ software.

**Cellular Immunofluorescence**

Cells were seeded in 24-well plates (2 × 10⁵ per well) containing round glass slides (14 mm in diameter), and when cells grow to 80% confluence, the glass slides with cells on were fixed with acetone on ice for 8 min, cells were subsequently blocked with 1% bovine serum albumin (BSA) and 0.5% Tween at room temperature for 30 min, and incubated with primary antibodies overnight at 4°C. The next day, cells on the glass slides were washed three times with the block solution and incubated with fluorescent-conjugated secondary antibody for 2 h at room temperature, after which cells were washed three times with the block solution. The primary antibodies consist of mouse anti-Pin1 (sc-46660; Santa Cruz, Dallas, TX, USA), rabbit anti-Pin1 (ab192036; Abcam), rabbit anti-ATF7 (ab87844; Abcam), and rabbit anti-Cyclin D1 (ab40754; Abcam). DAPI (D9564; Sigma-Aldrich) staining was used to show the cell nucleus. Images were acquired under confocal laser scanning microscope (FV3000; Olympus) and analyzed with ImageJ software.

**Immunofluorescence of Zebrafish Larvae**

The method was based on a previous study. Briefly, zebrafish larvae at 7 dpf (9 dpf of developmental stage) were fixed with 4% paraformaldehyde (PFA) at 4°C overnight. The following day, the skin of the zebrafish larvae was removed and the skinned larvae were dehydrated with methanol, then the larvae were placed in methanol at −20°C for 2 h. The dehydrated larvae were incubated with 3% H₂O₂ for 1 h at room temperature in dark. Larvae were rehydrated sequentially in 75%, 50%, and 25% methanol. Using 1% PT [1% Triton X-100 in phosphate-buffered saline (PBS)] to treat larvae four times for 15 min each. Then larvae were blocked with PBTN (4% BSA and 0.02% NaN₃ in 1% PT) at room temperature for 2 h and incubated with primary antibodies at 4°C overnight. The second day, the larvae were washed eight times with 1% PT for 30 min each and followed by incubation with fluorescent-conjugated secondary antibodies at 4°C overnight. The larvae were washed eight times with 1% PT for 30 min each. The primary antibodies include mouse anti-AE1/AE3 (ab86734; Abcam), mouse anti-Pin1 (sc-46660; Santa Cruz), rabbit anti-Pin1 (ab192036; Abcam), rabbit anti-ATF7 (ab87844; Abcam), rabbit anti-Cyclin D1 (ab40754; Abcam), and goat anti-GFP (ab5450; Abcam). DAPI (D9564; Sigma-Aldrich) staining was used to show the cell nucleus. Images were acquired under confocal laser scanning microscope (FV3000; Olympus) and spinning-disk confocal super resolution microscope (IXplore SpinSR; Olympus), and analyzed with ImageJ software.

**Statistical Analysis**

Statistical analyses were performed by Student’s *t* test, one-way analysis of variance (ANOVA) and two-way ANOVA. Statistical significance was defined as *P* < 0.05. All statistical analyses and quantitative image plotting were performed using GraphPad Prism 7 software. All data were generated and statistically analyzed from at least three independent experiments. We state that all the experiments and data are with high replicability.

**Results**

**Establishment of Zebrafish NPC Xenograft Model**

To establish zebrafish xenograft model for human NPC, we transplanted GFP-labeled human NPC CNE1 cells (Fig. 1A) into the perivitelline space around the yolk sac of 2 dpf wild-type zebrafish embryos (Fig. 1B). The xenografted zebrafish larvae were incubated at 34°C and imaged at stages of 1 dpf, 3 dpf, 5 dpf, and 7 dpf, respectively, to record and quantify the transplanted tumor cells in the fish body (Fig. 1B). Tumor cell–transplanted embryos at 1 dpf were used as controls for the comparison of tumor cell proliferation within hosts at 3 dpf, 5 dpf, and 7 dpf, respectively. As a result, the transplanted human CNE1 cells significantly increased at the stages of 5 dpf and 7 dpf compared with the number of tumor cells in fishes at 1 dpf (Fig. 1C). These suggest that human NPC CNE1 cells transplanted into zebrafish embryos could survive and proliferate well in vivo.
Migration, Invasion, and Proliferation of Transplanted NPC CNE1 Cells in Zebrafish

Tumor metastasis is one of the main causes of death in patients, which may even occur in the early stages of tumor development when the primary tumor is relatively small\(^1\). In conventional mammalian xenograft models, early-stage tumor micrometastases cannot be detected by real-time imaging analysis. In contrast, the zebrafish here provides a unique in vivo model that allows to visualize early micrometastases of tumor cells, and especially the invasion of cells after migrating out of the vessels. In the above assay by transplanting CNE1 cells into the perivitelline space around the embryonic yolk sac, we have observed proliferation of tumor cells in fish body. Next, to investigate the metastatic process of tumor cells in zebrafish, we transplanted tumor cells into the CCV of 2 dpf zebrafish embryos to construct an experimental metastasis model (Fig. 2A). After transplantation of the tumor cells, embryos were incubated at 34°C and were imaged at 1, 3, 5, and 7 dpt to trace the localization of tumor cells in zebrafish circulatory system (Fig. 2A). The results showed that CNE1 cells could reach most parts of the zebrafish body, including head, trunk, and tail regions, with blood circulation (Fig. 2A). It is noteworthy that in the caudal vein plexus of the zebrafish, the migrated GFP-labeled CNE1 cells survived and proliferated with the development of the embryos (Fig. 2B). By immunofluorescence staining of tumor markers of AE1/AE3, it was found that the transplanted and migrated NPC tumor cells were tumorigenic after the extravasation from the caudal vein plexus in the tail region (Fig. 2C).
Figure 2. Transplantation of human NPC CNE1 cells into the CCV of zebrafish to observe the behavior of tumor cells. (A) Top: diagram of the establishment of zebrafish xenograft model. Bottom: experimental protocol. (B) Left: tumor cells transplanted into the circulatory system migrate to the tail region of zebrafish with blood flow and proliferated there. Black arrows indicate GFP-labeled tumor cells in the tail region. Scale bar: 3 mm. Right: the relative fluorescence intensity of tumor cells in zebrafish trunk and tail regions was quantified. The amount of tumor cells at 1 dpt was used as the baseline for comparison with that at other time points. Data were quantitatively analyzed by one-way ANOVA analysis (n = 4 fishes analyzed). (C) Zebrafish whole-mount immunofluorescence detection of the epithelial-derived tumor marker of AE1/AE3. Tumor cells that reached the tail region with blood flow were able to become tumorigenic in the zebrafish. Dashed line frames beneath are the high magnification of the region indicated. The nuclei of CNE1 cells are shown by white dots, and the nuclei of zebrafish are shown by white asterisks. The size of the nuclei of CNE1 cells is obviously larger than those of zebrafish. Scale bar: 50 μm. (D) Tumor cells were transplanted into the Tg(kdrl:mCherry) zebrafish circulatory system, and tumor cells migrated out from vessel lumen within 24–48 hpt in zebrafish tail. High magnifications of the regions (white dashed line) are shown on the right. White arrows point to the vessels (dorsal aorta) in the zebrafish. Scale bar: 50 μm. Data are shown as mean ± SD. NPC: Nasopharyngeal carcinoma; CCV: common cardinal vein; GFP: green fluorescent protein; PTU: phenylthiourea; dpt: days post-transplantation; dpf: days post-fertilization; hpt: hours post-transplantation; NS: not significant. **P < 0.01; ***P < 0.001.
During the metastatic process, an important event is the invasion of cancer cells into the blood vessels to enter circulation, and later exiting the vasculature at a distant site. Therefore, we next sought to capture the migration behaviors of tumor cells out of the blood vascular wall of the caudal vein plexus by real-time images. To this end, Tg(kdrl:mCherry) transgenic zebrafish line which labels the vascular endothelial cells by mCherry fluorescent proteins was utilized. The results revealed that with 24 h after transplantation (from 24 to 48 hpt), CNE1 cells extravagated out from the dorsal aortic wall and invaded into intervascular space of dorsal aorta and posterior cardinal vein (Fig. 2D). Meanwhile, it was observed that the extravasation of tumor cells was accompanied by the morphological change of the vascular walls, and the arterial vessel was re-lumenized at 48 hpt when CNE1 cells completely migrated out (white arrows in Fig. 2D). The above results demonstrate that zebrafish xenograft model established can provide an ideal in vivo system for live tracing the processes of metastasis and invasion of tumor cells.

**Knockdown of Pin1 Inhibits the Proliferation of NPC Cells**

Pin1 specifically binds to and isomerizes the phosphorylated serine/threonine-proline (pSer/thro-pro) motif, leading to the alteration of protein structure, function, and stability. The structural and functional alterations of these phosphorylated proteins are closely related to the development of cancer. Previous studies have shown that Pin1 is overexpressed in NPC primary tumors and tumor cell lines, and that Pin1 activates the MAPK/JNK signaling pathway and upregulates the expression level of Cyclin D1, hence inducing the growth of NPC cells. Here, to further verify the role of Pin1 in NPC in our established zebrafish xenograft models, we constructed Pin1-knockdown CNE1 cell lines of NPC by transfecting pGLV3-shPin1 plasmid which expresses green fluorescent protein (GFP). Meanwhile, the empty pGLV3 vector-transfected CNE1 cell was used as a control. The transfected CNE1 cells were sorted by flow cytometry and imaged (Fig. 3A). Next, the expression of Pin1 was detected by western blot and immunofluorescence, respectively. The results indicated that the expression of Pin1 in the Pin1-knockdown cells was significantly reduced compared with that in the control group (Fig. 3B and Fig. S1A). In addition, the western blot and immunofluorescence staining of Cyclin D1 confirmed its reduced repression in Pin1-knockdown CNE1 cells (Fig. 3C, D).

Next, we transplanted GFP-labeled control or Pin1-knockdown CNE1 cells into the perivitelline space around the yolk sac of 2 dpf zebrafish embryos to investigate the survival and proliferation of these cells in vivo (Fig. 4A). As a result, it showed that both CNE1-LV3 and CNE1-shPin1 cells transplanted into the embryos could survive and proliferate within 7 dpt. However, the proliferation and growth of Pin1-knockdown CNE1 cells were significantly inhibited when comparing with that of the control group (Fig. 4A), which is consistent with the previous reports. To further explore the metastatic and invasive activity of CNE1 cells after knockdown of Pin1, we transplanted CNE1-LV3 and CNE1-shPin1 cells into the CCV of 2 dpf embryos, followed by real-time imaging of the xenograft embryos at 1, 3, 5, and 7 dpt, respectively (Fig. S1B). Our results revealed that due to the reduced Pin1 expression, the proliferation and invasion of CNE1-shPin1 cells in the caudal vein plexus region of zebrafish larvae were significantly inhibited (Fig. S1B). In contrast to the transplanted CNE1-LV3 control cells, majority of the transplanted CNE1-shPin1 cells died out within 7 days after transplantation (white arrows in Fig. S1B). In addition, we examined the expression levels of Pin1 and Cyclin D1 in tumor cells in vivo after transplanted into zebrafish embryos for 7 days by immunofluorescence staining. The results indicated that the expression of both Cyclin D1 and Pin1 was reduced in Pin1-knockdown CNE1 cells colonized in vivo when compared with that in the CNE1-LV3 cells, which was consistent with the findings of the expression of Cyclin D1 and Pin1 in vitro after loss of Pin1 (Fig. 4B).

**The Expression Level of ATF7 Is Correlated With Pin1 in NPC**

ATF7 is a member of the ATF/cAMP response element binding (CREB) protein family. Although it has already been identified for more than 20 years, its role in tumor development is still poorly understood. In contrast, the role of another ATF/CRE family member of ATF1 in NPC has already been widely explored. Previously, we have found that both ATF7 and ATF1 strongly interacted with Pin1 by a mammalian two-hybrid assay. Therefore, we assume that Pin1 might be a potential regulatory factor of ATF7. To verify this, we analyzed the expression level of ATF7 by western blot and immunofluorescence staining after knockdown of Pin1, respectively. The results indicated that the expression of ATF7 was suppressed after inhibition of the expression of Pin1 (Fig. 5A and Fig. S2). To further confirm the correlation between the ATF7 and Pin1, control or Pin1 containing plasmids were first transfected into CNE1 cells, and the expression of Pin1 and ATF7 was then analyzed by western blot. It was found that the expression of ATF7 increased significantly in Pin1-overexpressed CNE1 cells than that in control plasmid-transfected cells (Fig. 5B).

Subsequently, we examined the expression of ATF7 in CNE1 cells transplanted in vivo. The results revealed that in the inhibition of Pin1, the expression of ATF7 in CNE1-shPin1 cells was reduced in compared with that in CNE1-LV3 cells after transplantation in zebrafish for 7 days (Fig. 5C). Taken together, the results from in vivo and in vitro analyses suggested that Pin1 could efficiently affect the expression of ATF7, which might be also involved in Pin1-regulated carcinogenesis and development of NPC.
Figure 3. The expression of Cyclin D1 in tumor cells was reduced after knockdown of Pin1. (A) After transfection of LV3 control or shPin1 plasmids which express GFP as selecting marker, the CNE1-LV3 and CNE1-shPin1 cells were sorted first by FACS and imaged. Scale bar: 50 μm. (B) Western blot analysis was performed to detect the expression of Pin1 in tumor cells. The relative expression levels of Pin1 were quantified and the data were quantitatively analyzed by Student’s t test. Western blot (C) and immunofluorescence staining (D) of Cyclin D1 expression in CNE1 cells after knockdown of Pin1. Scale bar: 20 μm. Data were quantitatively analyzed using Student’s t test. Data are shown as mean ± SD. GFP: green fluorescent protein; FACS: fluorescence activating cell sorter. *P < 0.05; ***P < 0.001; ****P < 0.0001.
Figure 4. Inhibition of Pin1 expression suppressed tumor cell proliferation in zebrafish. (A) Combined fluorescence and bright field images of zebrafish xenograft model transplanted with CNE1-LV3 or CNE1-shPin1 into the perivitelline space around the yolk sac at 1, 3, 5, and 7 dpt, respectively. The relative fluorescence intensity of the tumor cells in zebrafish was quantified. The amount of tumor cells at 1 dpt was used as the baseline with which the amount of tumor cells at other time points was compared. The differences in cell proliferation between the two groups were quantitatively analyzed by two-way ANOVA (CNE1-LV3 group, n = 6 transplanted fishes analyzed; CNE1-shPin1 group, n = 11 transplanted fishes analyzed). Scale bar: 1 mm. (B) Whole-mount embryo immunofluorescence staining of Pin1 and Cyclin D1 expression in zebrafish with tumor cells transplanted at 7 dpt. Selected areas of images (white dashed line) are shown with a higher magnification below. Scale bar: 50 μm; 10 μm in the amplified image. The mean fluorescence intensity of Pin1 and Cyclin D1 was quantitatively measured, and the data were quantitatively analyzed by Student's t test (CNE1-LV3 group, n = 6 transplanted fishes analyzed; CNE1-shPin1 group, n = 4 transplanted fishes analyzed). Data are shown as mean ± SD. dpt: days post-transplantation; EGFP: enhanced green fluorescent protein. **P < 0.01; ***P < 0.001; ****P < 0.0001.
Figure 5. The expression of ATF7 was correlated with the expression of Pin1. (A-B) Western blot analysis was used to detect the expression of ATF7 in Pin1-knockdown or Pin1-overexpressed CNE1 cells. The relative expression levels of Pin1 and ATF7 were quantified (n = 4 independent repeats). (C) Whole-mount embryo immunofluorescence staining of ATF7 expression in tumor cells transplanted in zebrafish. Images of selected areas (white dashed line) are shown at higher magnification below. Scale bar: 50 μm; 10 μm in the amplified image. The mean fluorescence intensity of Pin1 and ATF7 was analyzed quantitatively. The data were quantitatively analyzed by Student’s t test (CNE1-LV3 group, n = 5 transplanted fishes analyzed; CNE1-shPin1 group, n = 3 transplanted fishes analyzed). Data are shown as mean ± SD. GAPDH: glyceraldehyde-3-phosphate dehydrogenase; EGFP: enhanced green fluorescent protein. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001.
In summary, our data provide strong evidences that \textit{zebrafish} could provide a simple, rapid, and visual xenograft model that allows for human cancer research compared with traditional rodent models.

\textbf{Discussion}

Although the combination of chemotherapy and radiotherapy has contributed to the increased survival rate of the NPC patients (5-year survival rate around 85% to 90%)\textsuperscript{3,4}, distant metastasis, recurrence, and chemoresistance are still challenges in the treatment for NPC. Therefore, studies on pathogenesis of NPC, identification of NPC-related biomarkers, optimization of treatment strategies for different patients, and development of novel treatment methods are urgently needed. In the traditional \textit{in vitro} tumor models, analysis of the interaction between tumor and host is hardly possible for the study of behavior and complexity of tumor, which are essential events in tumor growth and progression. Therefore, translational studies using \textit{in vitro} model for clinical application are still challenging\textsuperscript{27}, and to develop more efficient \textit{in vivo} models will help to explore accurate therapeutic strategies for cancer patients. In this study, we established a xenograft NPC model using classic model animal of \textit{zebrafish} and confirmed that human NPC cells can survive and proliferate in \textit{zebrafish} embryos. In addition, we combined the perivitel-line space transplantation with the CCV transplantation and observed the extravasation behaviors of NPC cells out of the blood vessels for the first time. We proved that human tumor cells can proliferate, migrate, and invade in \textit{zebrafish} body. Based on these observations, we propose that the \textit{zebrafish} xenograft model enables to visualize and investigate NPC cell metastasis \textit{in vivo}.

\textit{Pin1} is a key signal-regulating protein in tumors, which is highly expressed in invasive tumors, and is closely related to the development and prognosis of tumors\textsuperscript{25–27}. Existing studies have confirmed that \textit{Pin1} is involved in the occurrence and development of NPC\textsuperscript{14,24}. As a downstream gene of multiple signaling pathways, \textit{Cyclin D1} mainly plays a role in the process of G1 phase entering to S phase of cell cycle and has important influence on cell growth related to tumor deterioration and metastasis\textsuperscript{27}. In breast cancer, \textit{Pin1} overexpression is associated with increased expression of \textit{Cyclin D1}\textsuperscript{28}. In hepatocellular carcinoma, \textit{Pin1} overexpression has also been shown to upregulate \textit{Cyclin D1}\textsuperscript{29}. In NPC, overexpression of \textit{Pin1} activates MAPK/JNK and NF-κB pathway, thereby regulating the expression of \textit{Cyclin D1}\textsuperscript{32}. In this study, to evaluate the feasibility of \textit{zebrafish} xenograft model for tumor studies, we verified the regulatory effect of \textit{Pin1} on \textit{Cyclin D1} expression in transplanted CNE1 cell in \textit{zebrafish}. The finding was in consistent with those obtained \textit{in vitro} and in mice that loss of \textit{Pin1} inhibits the expression of \textit{Cyclin D1} thus inhibiting the proliferation of NPC cells\textsuperscript{32}.

\textit{ATF7} is a member of CREB protein family, and many members of the ATF/CREB family are associated with cancer progression. For example, \textit{ATF1} promotes the invasiveness of thyroid papillary tumors\textsuperscript{30}, \textit{ATF3} promotes the invasiveness of prostate tumor cells\textsuperscript{31}, and overexpression of \textit{ATF4} leads to drug resistance in human cancer cell lines\textsuperscript{32}. It has been demonstrated that \textit{ATF7} expression is upregulated in hepatocellular carcinoma tissues and that inhibition of \textit{ATF7} expression inhibits hepatocellular carcinoma cell proliferation and promotes apoptosis\textsuperscript{33}. However, the role of \textit{ATF7} in NPC has not yet been reported. In this study, we found that the expression of \textit{ATF7} in NPC cells was correlated with that of \textit{Pin1}. Therefore, we hypothesized that \textit{Pin1} might be a new regulator of \textit{ATF7}, and \textit{Pin1} might function in promoting the development of NPC by regulating the expression of \textit{ATF7}. However, the mechanism of the regulation of \textit{Pin1} on \textit{ATF7} has not been clarified, and the mechanism of \textit{ATF7} in NPC development needs to be further explored.

Although many advantages make \textit{zebrafish} a suitable \textit{in vivo} model for cancer research, there are still some inherent disadvantages, such as the different incubation temperatures for \textit{zebrafish} and mammalian cells, and diverse tumor microenvironment. Regarding the culturing temperature, the optimal growth temperature for human tumor cells is 37°C, whereas it is 28°C for \textit{zebrafish}. Therefore, the incubation temperature of 34°C was tested and finally chosen as the compromise temperature in our study. This temperature may lead to changes of \textit{zebrafish} metabolism\textsuperscript{14}. Nevertheless, the \textit{zebrafish} xenograft model in our study provides an intact microenvironment close to that in humans for tumor growth. In addition, the unique imaging advantages of transparent fish bodies, combining with multiple transgenic labeling of various organs, allow us to real-timely record the behaviors of tumor cells for a long period (7 days after tumor cell transplantation). In summary, we established a \textit{zebrafish} xenograft NPC model and briefly verified the involvement of \textit{ATF7} regulated by \textit{Pin1} in the occurrence of NPC.

\textbf{Conclusion}

The \textit{zebrafish} xenograft model provides an ideal \textit{in vivo} system for rapid identification of tumor formation and progressing and efficiently contributes to explore the mechanisms of NPC carcinogenesis and development.

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\textbf{Author Contributions}

E.H. performed the experiments, analyzed the data, and drafted the manuscript. H.H. and L.W. help with the cell experiment and transplantation. B.L. generated the stabilized cells. Z.H. gave suggestions and guidance. J.Z. initiated this study and monitored the whole progress of this project.
Ethical Approval
The whole experimental procedures were examined by the Clinical Ethics Committee of the Affiliated Hospital of Guangdong Medical University, China.

Statement of Human and Animal Rights
This study does not contain any studies with human subjects or with rodent vertebrates. Handling of zebrafish was performed in accordance with Guangdong State Regulations on Laboratory Animal Management (2010).

Statement of Informed Consent
Not applicable.

Declaration of Conflicting Interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Supplemental material
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