Neurturin Gene Therapy Protects Parasympathetic Function to Prevent Irradiation-Induced Murine Salivary Gland Hypofunction

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Head and neck cancer patients treated with irradiation often present irreversible salivary gland hypofunction for which no conventional treatment exists. We recently showed that recombinant neurturin, a neurotrophic factor, improves epithelial regeneration of mouse salivary glands in ex vivo culture after irradiation by reducing apoptosis of parasympathetic neurons. Parasympathetic innervation is essential to maintain progenitor cells during gland development and for regeneration of adult glands. Here, we investigated whether a neurturin-expressing adenovirus could be used for gene therapy in vivo to protect parasympathetic neurons and prevent gland hypofunction after irradiation. First, ex vivo fetal salivary gland culture was used to compare the neurturin adenovirus with recombinant neurturin, showing they both improve growth after irradiation by reducing neuronal apoptosis and increasing innervation. Then, the neurturin adenovirus was delivered to mouse salivary glands in vivo, 24 hr before irradiation, and compared with a control adenovirus. The control-treated glands have ~50% reduction in salivary flow 60 days post-irradiation, whereas neurturin-treated glands have similar flow to nonirradiated glands. Further, markers of parasympathetic function, including vesicular acetylcholine transporter, decreased with irradiation, but not with neurturin treatment. Our findings suggest that in vivo neurturin gene therapy prior to irradiation protects parasympathetic function and prevents irradiation-induced hypofunction.

INTRODUCTION

Every year approximately 630,000 new head and neck cancer patients are diagnosed globally.1 Therapeutic irradiation (IR) is one of the primary treatments, and ~64% of survivors treated with IR suffer from permanent xerostomia, the subjective symptom of dry mouth.2 There is also an acute reduction in salivary gland flow rates of ~50% within a week of IR, which can diminish to ~20% by 7 weeks.3 This salivary hypofunction is often accompanied by xerostomia, although xerostomia may not always correlate with salivary flow rates.4 The primary pathophysiological mechanism that causes salivary hypofunction is not clear, although the loss of acinar cells and disruption of muscarinic receptor-stimulated water secretion have been proposed.4 Currently, there are no therapies to prevent or permanently repair this IR damage.2,4

We previously showed that during murine submandibular gland (SMG) development the parasympathetic ganglion (PSG) is critical to maintain a population of epithelial keratin 5-positive (K5+) progenitors, which are required for gland development.2 Acetylcholine signaling from the PSG, via muscarinic M1 receptors and EGFR in the epithelium, increased K5+ progenitor proliferation. We then showed that IR of the developing SMG in ex vivo culture reduced innervation by causing apoptosis of ~30% of distal endbud epithelial cells that produced neurturin (NRTN), a neurotrophic factor. NRTN binds its receptors GFRz2/RET and is important for parasympathetic nervous system development and function.6,7 NRTN-deficient mice have reduced PSG and defects in SMG innervation.8 In our previous experiments, the loss of the NRTN-expressing endbud cells disrupted epithelial-neuronal interactions, resulting in a loss of innervation, and subsequently, 3 days after IR, apoptosis of PSG neurons occurred. However, when IR SMGs were treated with recombinant NRTN, which binds GFRz2 on the nerves, reduced neuronal apoptosis was observed. Consequently, NRTN restored parasympathetic function, which improved innervation and epithelial regeneration after IR.9 It has also been shown, using reversible duct ligation as a model to study gland regeneration after damage, that epithelial regeneration after removal of the ligature only occurred if the parasympathetic innervation was intact. The remaining epithelium regenerated in a similar

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manner to gland development, i.e., by branching morphogenesis.\textsuperscript{10} Taken together, these data suggest that NRTN could be a candidate for treatment of IR-induced salivary hypofunction by targeting the parasympathetic innervation to stimulate epithelial regeneration and saliva secretion.

Salivary glands are directly accessible through ducts in the oral cavity, and gene transfer to the gland epithelia is accomplished by delivery of a vector by retro-ductal cannulation. Several studies demonstrated the success of non-integrating adenovirus serotype 5 vectors in transducing adult mouse SMGs following retrograde ductal cannulation.\textsuperscript{11–14} Adenoviral vectors show sustained transgene expression in the SMGs for up to 6 months after exposure to fractionated IR.\textsuperscript{14} As a result, a phase 1 clinical trial was completed, using adenoviral-mediated transfer of aquaporin-1 cDNA to the salivary epithelium in patients previously treated by IR for head and neck cancer. The study confirmed that gene therapy is feasible in humans to potentially increase fluid secretion and relieve xerostomia experienced by patients after IR.\textsuperscript{15,16}

A gene therapy vector expressing NRTN was used in human clinical trials in the central nervous system as a potential treatment for Parkinson’s disease.\textsuperscript{17} Here, we delivered a serotype 5 adenovirus expressing NRTN (AdNRTN) prior to fractionated IR of SMGs. The AdNRTN-treated SMGs produced similar amounts of saliva as control non-IR mice 60 days after IR. We propose that NRTN protects the PSG from IR damage by promoting PSG neuronal survival, subsequently allowing regeneration of the remaining epithelium. Thus, NRTN gene therapy may be a potential strategy to prevent permanent IR-induced hypofunction in humans being treated for head and neck cancer.

RESULTS

AdNRTN and Recombinant NRTN Protein Have Similar Effects in Fetal SMG after IR

Our initial experiments were to determine whether AdNRTN treatment was similar to recombinant NRTN treatment in \textit{ex vivo} fetal SMG culture. We hypothesized the virus may infect both mesenchyme and epithelium, which would then secrete NRTN to improve innervation after IR. First, we tested a range of viral doses by treating \textit{ex vivo} embryonic day 13.5 (E13.5) SMGs for 24 hr with concentrations of AdNRTN ranging from $10^2$ to $10^5$ viral particles/cell (vp/c) and then a 5 Gy dose of IR. We calculated the average cell number in an E13.5 SMG (20,000 cells/gland) to determine the vp/c. After a further 72 hr of \textit{ex vivo} culture, there was a significant increase in endbud number with an AdNRTN concentration of $10^4$ vp/c compared with IR SMGs without AdNRTN treatment (Figure 1A). The endbud number is a measure of branching morphogenesis and SMG growth, and is expressed as a fold change over the number of endbuds at the
beginning of the experiment. Further increases in the dose of AdNRTN virus did not improve SMG growth post-IR.

For the following experiments, we used E14.5 SMGs in ex vivo culture (Figure 1B) to directly compare the AdNRTN (10⁴ vp/c) with treatment with recombinant NRTN protein. This assay was similar to our previous studies, where we showed that a 5 Gy IR dose reduced endbud number due to apoptosis of the proliferating epithelium, but allowed survival of the PSG and axonal bundles. Both AdNRTN and recombinant NRTN protein treatment 24 hr before IR improved gland growth 72 hr post-IR (a total of 96 hr ex vivo culture), and the morphology of the epithelial endbuds appeared more similar to non-IR glands (Figure 1B). The irradiated SMGs treated with either AdNRTN or recombinant NRTN also had a 2-fold increase in endbud number compared with those treated with a control GFP-expressing virus (AdC) (Figure 1C). Moreover, viral infection alone did not stimulate endbud growth because AdC treatment was similar to IR alone (data not shown).

We measured gene expression with qPCR 72 hr post-IR and compared non-IR SMGs and IR SMGs treated with either AdNRTN or recombinant NRTN with the AdC treatment (Figure 1D). We measured expression of Nrtn its coreceptor, Ret, and the neuronal marker Tubb3. Overall, Nrtn expression was not reduced by 5 Gy IR, and the surviving cells expressed similar levels of Nrtn as recombinant NRTN-treated SMGs. Although we confirmed that AdNRTN treatment increased Nrtn expression (~1.5-fold) after IR, Ret expression was similar in all groups. The expression of Tubb3 is relatively increased after 5 Gy IR (Figure 1D), because IR has direct effects on the epithelium, increasing apoptosis. Thus, the surviving neurons, which are more resistant to IR, result in increased Tubb3 expression by qPCR.

We next used whole-mount immunostaining to assess PSG axonal outgrowth and NRTN expression 72 hr post-IR after AdNRTN treatment. After 5 Gy IR, the PSG and axons are clearly present along the ducts, although the epithelium is reduced in size (Figure 2A). At higher magnification (Figure 2B) it is clear that the defasciculation of axons around the distal endbuds are reduced with IR. Both AdNRTN and recombinant NRTN increased epithelial budding and increased axonal projections surrounding them (Figure 2A). The PSG axon bundles (red, TUBB3⁺) appeared thicker surrounding the buds in SMGs treated with either AdNRTN or NRTN. NRTN protein (green, NRTN) was also present in the epithelial buds of AdNRTN- and NRTN-treated SMGs (higher magnification, Figure 2B). There appeared to be groups of cells with high NRTN expression after AdNRTN treatment (white arrows, Figure 2B), suggesting that the virus may be preferentially infecting a subpopulation of cells. These findings suggest that AdNRTN may protect the PSG after IR during ex vivo SMG culture to improve innervation and distal endbud regeneration.

**AdNRTN Reduces Apoptosis and Promotes PSG Survival in Fetal SMGs after IR**

We previously showed that recombinant NRTN treatment reduces neuronal apoptosis and restores parasympathetic function after irradiation, and we needed to confirm that AdNRTN treatment would have a similar effect. Therefore, we measured immunostaining of cleaved caspase-3, an apoptotic marker, in the PSG after IR and treatments. Both AdNRTN and NRTN reduced the number of caspase-3-positive neuronal cell bodies (TUBB3⁺) in the PSG of IR SMGs in ex vivo culture (Figure 3A). Quantitation of fluorescence showed there was a significant ~2.5-fold reduction in apoptotic cells in the PSG when IR SMGs were treated with either AdNRTN or NRTN (Figure 3B). Thus, AdNRTN and NRTN have similar effects in reducing neuronal apoptosis after IR.

**In Vivo AdNRTN Treatment followed by Fractionated IR Rescues Saliva Secretion 60 Days Post-IR**

We then tested whether AdNRTN treatment before IR has a similar effect in adult murine SMGs in vivo. Adult mice were treated with a retro-ductal infusion of AdNRTN (10⁴, 10⁵, or 10¹⁰ vp/gland [vp/g]) or an empty vector adenoviral control (AdC) by ductal cannulation into both SMGs. The following day, ~24 hr later, the SMGs were treated for 5 consecutive days with fractionated IR (5 × 6 Gy). After 60 days, the whole stimulated salivary volume was measured for 15 min by using capillary tubing to drain saliva from the floor of the mouth after intraperitoneal pilocarpine administration (Figure 4). It has been previously shown that by...
60 days post-IR there is a reproducible chronic loss of saliva.\textsuperscript{18–20} As expected, IR significantly reduced salvia secretion more than 50% compared with non-IR controls (Figure 4), and treatment with the control vector (AdC+IR) did not improve saliva secretion. However, treatment with all three doses of AdNRTN before IR improved salivary secretion so that it was similar to non-IR control levels (Figure 4). Saliva secretion did not change in non-IR mice treated with AdNRTN 10^10 vp/g (AdNRTN10^10 vp/g non-IR). These data suggest that treatment with AdNRTN before IR may be a useful strategy to protect secretory function.

**In Vivo AdNRTN Treatment Does Not Alter SMG Histology but Increases Markers of Parasympathetic Vesicular Transport**

After in vivo saliva collection, the SMGs were harvested and analyzed by qPCR, histology, and immunostaining. We first confirmed that we could detect increased expression of \textit{Nrtn} 60 days post-IR, which correlated with increased doses of AdNRTN treatment (Figure 5A). We also confirmed we could detect adenovirus DNA in the SMGs 60 days after treatment (Figure S1). We used qPCR to show there was also a significant increase in expression of \textit{Ret}, the kinase receptor that is a coreceptor with GFRz2 for NRTN. \textit{Ret} expression decreased \textasciitilde0.7-fold after IR and increased to \textasciitilde1.2-fold of non-IR levels with AdNRTN treatment (Figure 5B). Furthermore, expression levels of parasympathetic and sympathetic genes, \textit{Vacht} and \textit{Th}, respectively, were restored with AdNRTN treatment 60 days after IR damage (Figure 5B). There were no significant differences in the expression of \textit{Gfra2} and \textit{Tubb3} or other parasympathetic and sympathetic markers, \textit{Chat} and \textit{Adra2b}, respectively, between the IR and IR with AdNRTN groups (Figure 5B).

We then analyzed adult SMG tissue histology and markers of innervation after AdNRTN treatment. The cellular morphology of H&E-stained SMG tissues appeared comparable among all in vivo treatments (Figure 6A). These observations are consistent with previous reports, where major atrophy and tissue fibrosis are absent 2 months after IR damage in this strain of mice and appear a few months later.\textsuperscript{21} In addition, after staining SMGs for TUBB3, a pan-neuronal marker, increased innervation was observed with IR (Figures 6B and 6C). However, the protein expression of the parasympathetic marker, vesicular acetylcholine transporter (VACHT), but not the sympathetic marker tyrosine hydroxylase (TH), significantly increased with \textit{in vivo} AdNRTN treatment compared with non-IR (Figures 6B and 6C). VACHT and TH are essential for acetylcholine neurotransmitter release and for catecholamine synthesis, respectively.\textsuperscript{22,23} Therefore, changes in VACHT suggest that AdNRTN is acting directly upon the parasympathetic nerves.

**NRTN Does Not Increase Proliferation in Head and Neck Cancer Cell Lines**

A possible concern using gene therapy with a neurotrophic factor is the potential off-target proliferative effects on any remaining head and neck squamous carcinoma cells. To begin to address this issue, we treated 3 different head and neck cancer cell lines (WSU-HN12, WSU-HN6, WSU-HN8\textsuperscript{24}) with recombinant NRTN and measured cell proliferation (Figure S2). These cell lines were previously analyzed by RNA-sequencing (RNA-seq) analysis\textsuperscript{24} and did not express GFRz2, the receptor for NRTN. Here, we found that the direct treatment of the tumor cells with recombinant NRTN did not induce off-target proliferation.

**DISCUSSION**

Here, we provide proof-of-principle evidence that gene therapy \textit{in vivo} with an AdNRTN protects the parasympathetic innervation of the murine SMG 60 days after IR, resulting in stimulated saliva production similar to non-IR controls. There was no discernable difference in the histology of the SMGs at 60 days post-IR with any of the treatments, although there was also an overall increase in neuronal staining and an increase in staining of markers of both sympathetic and parasympathetic nerves in the AdNRTN-treated SMGs. We speculate that the mechanism by which AdNRTN protects saliva production may be similar to that reported in our previous \textit{ex vivo} experiments. We previously proposed that recombinant NRTN was a survival factor and reduced apoptosis of the PSG neurons 3 days after IR-induced apoptosis of the \textit{Nrtn}-expressing epithelial cells.\textsuperscript{25} Whether PSG apoptosis occurs 3 days after IR \textit{in vivo} remains to be determined. The increase in neuronal staining with markers of both sympathetic and parasympathetic nerves suggests that the balance of innervation by these two autonomic pathways may be important for regeneration.
A gene therapy trial to treat IR-induced xerostomia was aimed at repairing fluid secretion in the surviving ductal epithelial cells long after the IR damage. The phase 1 clinical trial reported some success using an Aquaporin-1-expressing adenovirus. Aquaporin-1 is a water channel not normally expressed in salivary ductal epithelium and was hypothesized to increase salivary fluid flow from the surviving undamaged ductal cells post-IR. Our proof-of-principle in vivo study using AdNRTN before IR is based on the hypothesis that providing neurotropic support for the parasympathetic neurons will allow improved salivary function and regeneration of surviving epithelial progenitors after IR. We speculate that the improved salivary function may be a result of reduced neuronal apoptosis and an increase in PSG survival via neuronal NRTN/GFRα2/RET signaling. This is supported by the increased expression of VACHT in neurons after AdNRTN treatment. Studies have shown a strong relationship between the levels of VACHT expression and acetylcholine release into the synaptic cleft in the peripheral nervous system, suggesting that NRTN is increasing parasympathetic activity. Moreover, the increase in TH by AdNRTN treatment also denotes a potential increase in the synthesis of catecholamines, markers of sympathetic neurotransmission. Sympathetic signaling was reported to have inhibitory effects on epithelial regeneration by blocking muscarinic-induced EGFR activation. Nevertheless, these changes in VACHT and TH suggest that AdNRTN delivery directly affects GFRα2-expressing parasympathetic nerves and indirectly affects sympathetic nerves, although the mechanisms of this crosstalk remains to be investigated.

The murine SMG is used as a model to study how to prevent, repair, or regenerate function after IR damage. However, a caveat with the murine SMG model at 60 days post-IR is that it does not present the atrophy, fibrosis, and inflammatory infiltrates, which are a histopathological hallmark of human gland post-IR. The pathophysiological mechanisms causing hypofunction in the absence of histological damage are not clear. A likely mechanism relates to disruption of muscarinic signaling that drives salivary fluid secretion. We speculate that NRTN stimulates parasympathetic innervation and restores secretion of neurotransmitters, increasing cholinergic signaling via muscarinic receptors. While our studies are promising, future studies will be required using animal models such as the minipig, which respond to IR in a similar manner to human glands. Whether stimulation of parasympathetic innervation and subsequent epithelial regeneration can prevent the fibrotic and inflammatory response, or whether AdNRTN could be used to treat IR-induced hypofunction long after the initial IR damage remain to be determined.
MATERIALS AND METHODS

Ex Vivo SMG Culture Treatment and Irradiation

Fetal SMG explants from ICR embryos were cultured as previously described. 13,14 E13.5 or E14.5 SMGs were placed in culture dishes and incubated with fresh medium containing BSA, treated with either a single dose of recombinant mouse NRTN or adenovirus expressing NRTN cDNA. The optimal dose of recombinant NRTN was previously determined to be 1 ng/mL. 15 For adenovirus treatment, 10^5–10^6 vp/c or an empty control vector were added to the media. Explants were irradiated 24 hr later in a Gammacell 1000 irradiator with a single dose of irradiation (5 Gy). SMGs were cultured for a further 24–48 hr. Explants were either lysed for RNA or fixed for immunostaining. The endbud number was counted at the beginning of the experiment (time 0) and at the end of the assay (72 hr) and expressed as a ratio (endpoint/T0) and normalized to the control irradiated treatment (% IR control). All experiments showing endbud numbers were obtained using 4 SMGs/group, and each experiment repeated at least three times.

Adenovirus Vector Construction

The AdLTR_EF1α-hNRTN vector used for these experiments, here abbreviated to AdNRTN for simplicity, is a hybrid E1-deleted serotype 5 recombinant adenovector with a human NRTN cDNA (hNRTN; Origene Technologies, Rockville, MD, USA), which can persist long-term transgene expression in the salivary gland. 13 In brief, to construct plasmid pACLTR_EF1α-hNRTN, the plasmid pACLTR_EF1α was digested with BamHI filled in with T4 DNA polymerase and then digested with EcoRI to ligate ~0.6 kb of hNRTN; pCMV6-5L5-hNRTN (Origene) was digested with Xba I and filled in with T4 DNA polymerase and digested with EcoRI to obtain the hNRTN cDNA. The recombinant vector AdLTR_EF1α-hNRTN was produced by homologous recombination of pACLTR2EF1α-hNRTN with pJM17 in C7 cells. The titers (particles/mL) of purified vectors were determined by qPCR using primers from the E2 region. 13

Gene Expression Analysis

Real-time PCR was performed as previously described. 38 Primers were designed using Beacon Designer software. Melt-curves and primer efficiency were determined as previously described. Gene expression was normalized to non-IR SMGs and the ribosomal housekeeping gene Rps29 (data not shown). qPCRs were done in triplicate, and N = 11–12 SMGs analyzed. Vacht expression was not detected in 5 of 11 SMGs. Error bars represent the SEM. ANOVA with post hoc Dunnett’s test compared with non-IR, *p < 0.05; **p < 0.01; ***p < 0.001.

For the irradiated mice, only head and neck area was irradiated by placing each animal in a specially built Lucite jig so the animal could be immobilized without the use of anesthetics. 40 Additionally, the jig was fitted with a Lucite cone surrounding the head and preventing head movement during radiation. Fractionated irradiation at 5 × 6 Gy (6 Gy/day for 5 days) was delivered 24 hr after vector delivery by a Therapax DXT300 X-ray irradiator (Pantak). After irradiation, animals were removed from the jig, housed (four animals per cage) in a climate- and light-controlled environment, and allowed free access to food and water.

Blood, saliva, and tissue were collected on day 60. For saliva collections, anesthetized mice were stimulated using 1 μL/g body weight of a pilocarpine solution subcutaneously (0.5 mg/mL). Whole saliva was collected with a 75-mm hemocrit tube (Drummond) into 1.5 mL pre-weight Eppendorf tubes for 15 min and frozen immediately. Blood samples were collected from the retro-orbital sinus after the saliva collection. To separate the serum phase from blood, we centrifuged samples at 12,000 rpm for 3 min, and the serum supernatant was collected and stored at −80°C. Mice were sacrificed in a carbon monoxide chamber and the SMGs were removed.

Figure 5. Treatment of SMGs In Vivo with Increasing Doses of AdNRTN Increases Nrtn Transcripts within the Gland and Improves Some Neuronal Markers after IR

(A) qPCR analysis of Nrtn transcripts in SMGs 60 days after IR and treatment with increasing doses of AdNRTN. (B) Additional qPCR of genes related to NRTN signaling in IR SMGs treated with AdNRTN 10^9 compared with IR treatment alone. The expression of the NRTN receptors Gria2 and Ret kinase, neuronal markers Tubb3, and the parasympathetic markers Chat and Vacht and the sympathetic markers Th and Adra2b. Gene expression was normalized to non-IR SMGs and the ribosomal housekeeping gene Rps29 (data not shown). qPCRs were done in triplicate, and N = 11–12 SMGs analyzed. Vacht expression was not detected in 5 of 11 SMGs. Error bars represent the SEM. ANOVA with post hoc Dunnett’s test compared with non-IR, *p < 0.05; **p < 0.01; ***p < 0.001.
Histopathology
Intact SMGs were surgically removed from each mouse following euthanasia. SMGs were either lysed for RNA or protein, cryopreserved, or fixed in paraformaldehyde (PFA) 4%. After PFA fixation, 4-μm paraffin sections were stained with H&E and visualized in an Axiovert 25 light microscope (Zeiss, NY, USA) at 40× magnification. H&E images were taken with a Nikon 5000 camera (Nikon, Japan). Cryopreserved in vivo SMG tissues were used for immunofluorescence analysis.

Immunofluorescence Analysis
Whole-mount fetal SMG immunofluorescence analysis has been described previously. Ex vivo SMGs and PSGs were fixed with 4% PFA for 20 min at room temperature and/or ice-cold acetone/methanol for 5 min. In vivo SMGs were cryosectioned at 200 μm and fixed similar to the ex vivo SMGs. Tissues were blocked overnight with 10% donkey serum (Jackson Laboratories, ME, USA), 1% BSA, and mouse on mouse (MOM) immunoglobulin G (IgG) blocking reagent (Vector Laboratories, CA) in 0.01% PBS-Tween 20. SMG tissues were incubated with primary antibodies overnight at 4°C: goat anti-GFRα2 (1:100; R&D Systems), goat anti-NRTN (AB477, 1:100; R&D Systems), mouse anti-TUBB (clone TU1 at 1:200; R&D Systems), goat anti-vesicular acetylcholine transporter (VACHT; 1:100; Millipore), chicken anti-TH (1:400; Neuromics), rabbit anti-caspase-3 (1:200; Cell Signaling), rat anti-perlecan (1:400; Chemicon), and rabbit anti-E-cadherin (1:200; Cell Signaling). Nuclei were stained with Hoechst C10337 (Invitrogen). Antibodies were detected using Cy2-, Cy3-, or Cy5-conjugated secondary Fab fragment antibodies (Jackson Laboratories, ME, USA). Fluorescence was analyzed using a Zeiss 710 LSM microscope (Zeiss, NY, USA). Quantitation of the caspase-3-positive cells in the PSG of fetal SMGs was performed according to our previous publication. Quantitation of Tubb3, VACHT, TH, GFRα2, and Hoechst fluorescence in the in vivo adult SMG sections was calculated from z stacks consisting of eight confocal images separated 3 μm apart and scanned through the section at four different random quadrants (N = 4). The Tubb3, GFRα2, VACHT, and TH fluorescent staining were normalized to the nuclear staining.

Statistics
We used GraphPad Prism 7.0 software (GraphPad Software, CA, USA). Data were log transformed, and one-way ANOVA was used comparing groups with the experimental control. Graphs show the mean ± SEM.

Cell Culture
The following cell lines were a gift from S. Gutkind (University of California San Diego [UCSD]): WSU-HN12, WSU-HN6, WSU-HN8. They were cultured in DMEM (11965-092; Thermo Fisher Scientific) with 10% FBS, in 5% CO₂, at 37°C.

WST-1 Proliferation Assay
Cells were seeded at 300 cells/well on 96-well dish, allowed to adhere overnight, then serum starved for 24 hr. Cells were then stimulated with Nrtin (0.01 and 1 ng/mL) (R&D Systems, Minneapolis, MN, USA), diluted in 0.1% BSA in PBS. Cell proliferation was assayed in both serum-free and 1% FBS with a WST-1 colorimetric cell proliferation assay according to the manufacturer’s instructions (05015944001; Sigma, St. Louis, MO, USA).
SUPPLEMENTAL INFORMATION

Supplemental Information includes two figures and can be found with this article online at https://doi.org/10.1016/j.omtm.2018.02.008.

AUTHOR CONTRIBUTIONS

Conceptualization, M.P.H., J.N.A.F., and I.M.A.L.; Investigation, J.N.A.F., C.Z., I.M.A.L., C.M.G., A.P.C., J.M.S., and V.N.P.; Writing, J.N.A.F., I.M.A.L., V.N.P., and M.P.H.; Supervision, V.N.P. and M.P.H.

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