Dysregulation of miR-31 and miR-21 induced by zinc deficiency promotes esophageal cancer

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Zinc deficiency (ZD) increases the risk of esophageal squamous cell carcinoma (ESCC). In a rat model, chronic ZD induces an inflammatory gene signature that fuels ESCC development. miRNAs regulate gene expression and are aberrantly expressed in cancers. Here we investigated whether chronic ZD (23 weeks) also induces a protumorigenic microRNA signature. Using the nanoString technology, we evaluated microRNA profiles in ZD esophagus and six additional tissues (skin, lung, pancreas, liver, prostate and peripheral blood mononuclear cells [PBMC]). ZD caused overexpression of inflammation genes and altered microRNA expression across all tissues analyzed, predictive of disease development. Importantly, the inflammatory ZD esophagus had a distinct microRNA signature resembling human ESCC or tongue SCC miRNAomes with miR-31 and miR-21 as the top-up-regulated species. Circulating miR-31 was also the top-up-regulated species in PBMCs. In ZD esophagus and tongue, oncogenic miR-31 and miR-21 overexpression was accompanied by down-regulation of their respective tumor-suppressor targets PPP2R2A and PDCD4. Importantly, esophageal miR-31 and miR-21 levels were directly associated with the appearance of ESCC in ZD rats, as compared with their cancer-free Zn-sufficient or Zn-replenished counterparts. In situ hybridization analysis in rat and human tongue SCCs localized miR-31 to tumor cells and miR-21 to stromal cells. In regressing tongue SCCs from Zn-supplemented rats, miR-31 and miR-21 expression was concomitantly reduced, establishing their responsiveness to Zn therapy. A search for putative microRNA targets revealed a bias toward genes in inflammatory pathways. Our finding that ZD causes miR-31 and miR-21 dysregulation associated with inflammation provides insight into mechanisms whereby ZD promotes ESCC.

Introduction

Oral-esophageal squamous cell carcinomas (OSCC, ESCC) are major causes of cancer deaths worldwide. Because of absence of early symptoms, ESCC is frequently diagnosed at an advanced stage and has a 5-year survival of 10%. Patients with OSCC (major site, tongue) have a high mortality rate, because of the appearance of second cancers in the esophagus through field carcinization effects (1). Despite the decline in worldwide cancer-mortality rates since the mid 1980s, ESCC and OSCC remain deadly diseases. Thus, clarification of their pathogenesis and development of new prevention strategies are urgently needed.

The major risk factors for developing oral-esophageal carcinomas include alcohol consumption, tobacco use and nutritional deficiencies. Epidemiological and clinical studies have implicated dietary zinc (Zn) deficiency (ZD) in the pathogenesis of ESCC and OSCC (2,3). Zn is an essential trace element required for the activity of >300 enzymes, for proper immune function and for the conformation of >2000 transcription factors that control cell proliferation, apoptosis and signaling pathways (4). ZD predisposes to diseases by adversely affecting these processes.

Our ZD rat oral-esophageal cancer model reproduces the ZD link to human ESCC and OSCC (5) and provides a unique opportunity to decipher the mechanism by which ZD promotes oral-esophageal carcinogenesis. Previously, we showed that weaning rats on a ZD diet for 6 weeks developed a precancerous condition in the upper aerodigestive tract (tongue, esophagus and forestomach [expanded lower esophagus]), with increased cellular proliferation (5) and extensive gene-expression changes, including overexpression of proinflammation genes S100 calcium binding protein a8 and a9 (S100a8 and S100a9) (6). Subsequently, prolonged ZD (21 weeks) was shown to amplify this inflammation in the esophagus by causing overexpression of numerous inflammation genes in addition to S100a8/S100a9, thereby providing an inflammatory microenvironment conducive to ESCC development on repeated exposure to the environmental carcinogen N-nitrosomethylbenzylamine (NMBA). Zn replenishment (ZR) reversed the inflammatory signature and prevented cancer formation (7). In addition, Zn supplementation in nutritionally complete rats suppressed tongue SCC development by attenuating the inflammation (8). miRNAs (miRNAs) are a family of short noncoding RNAs that have emerged as powerful posttranscriptional regulators of gene expression (9). The ability of individual miRNAs to regulate a large number of mRNA species allows them to coordinate complex programs of gene expression that alter a range of biological processes, including cellular proliferation, apoptosis, immune response and signaling events (10). miRNAs expression levels are altered in many types of human cancer, including ESCC and OSCC (11). Furthermore, miRNA expression patterns are associated with chronic inflammatory diseases and other inflammatory conditions (12).

We asked whether overexpression of cancer-related inflammation genes induced by ZD in the rat esophagus (7) is a consequence of a protumorigenic microRNA signature. For this, we examined miRNA profiles in esophagus and six additional tissues (skin, lung, pancreas, liver, prostate and peripheral blood mononuclear cells [PBMC]) after chronic ZD (23 weeks) using the novel nanoString nCounter technology (Seattle, WA) (13,14). The deficient esophagus exhibited a protumorigenic miRNA signature with miR-31 and miR-21 as the top-up-regulated species. The role of these two oncomiRs in esophageal and oral neoplasia was further explored. ZD also induced distinct miRNA expression patterns across all tissues profiled.

Materials and methods

Animals

Male weanling male Sprague-Dawley rats were obtained from Taconic Laboratory (Germantown, NY). Custom-formulated egg-white-based diets were from Harlan Teklad (Madison, WI). ZD- and Zn-sufficient (ZS) diets were identical except for the Zn content, which was 3–4 ppm and 65 ppm, respectively.

Abbreviations: BCIP, 5-bromo-4-chloro-3-indolylphosphate; DA VID, 5-bromo-4-chloro-3-indolyl-β-D-glucuronic acid; DAB, 3,3′-diaminobenzidine dihydrochloride; DAPI, 4,6-diamidino-2-phenylindole; DAPI, 4,6-diamidino-2-phenylindole dihydrochloride; DAVID, Database for Annotation, Visualization and Integrated Discovery; ESCC, esophageal squamous cell carcinoma; FFPE, formalin fixed and paraffin embedded; GO, gene ontology; HIC, immunohistochemistry; ISH, in situ hybridization; miRNAs, microRNAs; NBT, 5-nitro-blue tetrazolium; PBMC, peripheral blood mononuclear cells; Zn, zinc; ZD, Zn deficient; ZS, Zn sufficient; ZR, Zn replenishment.

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References

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Experimental design

The animal study was conducted according to approved Institutional animal protocols. Weaning rats (n = 19) were fed ad libitum a ZD diet for 23 weeks. Control rats (n = 15) were pair fed a ZS diet to match the decreased food consumption of ZD animals (7). At study conclusion, blood was collected from the retro-orbital venous plexus after anesthesia with isoflurane. PBMCs were prepared from whole blood using Histopaque-1077 (Sigma-Aldrich, St. Louis, MO). Esophagus, tongue, skin, lung, liver, prostate and pancreas were isolated. Esophagus and tongue were cut into two portions. One portion was formalin fixed and paraffin embedded (FFPE) for histological, immunohistochemical (IHC), and in situ hybridization (ISH) studies. Esophageal and tongue mucosa were prepared as previously described (15). The prostate was microdissected into dorsal, lateral, ventral and anterior lobes. Tissues were snap frozen in liquid nitrogen and stored at −80°C.

RNA preparation

Total RNA was prepared using TRIzol reagent (Invitrogen, Carlsbad, CA) (7).

miRNA expression profiling

The multiplexed nanoString nCounter miRNA system (nanoString Technologies) (13,14) was used for miRNA expression profiling in esophagus and six additional tissues (skin, lung, pancreas, liver, prostate and PBMC; n = 6 rats/tissue/dietary group). This assay was performed at the Ohio State University Comprehensive Cancer Center Nucleic Acid Facility.

Total RNA (100ng) was used as input material. Small RNA samples were prepared by ligating a specific DNA tag onto the 3’ end of each mature miRNA according to manufacturer’s instruction (nanoString Technologies). These tags normalized the melting temperatures (Tms) of the miRNAs and provided identification for each miRNA species in the sample. Excess tags were then removed and the resulting material was hybridized with a panel of miRNA tag-specific nCounter reporter probes. Hybridization reactions were incubated at 64°C for 18h. Hybridized probes were purified and immobilized on a streptavidin-coated cartridge using the nCounter Prep Station (nanoString Technologies). nCounter Digital Analyzer was used to count individual fluorescent barcodes and quantify target RNA molecules present in each sample. For each assay, a high-density scan (600 fields of view) was performed.

nanoString data analysis

Abundances of miRNAs were quantified using the nanoString nCounter gene-expression system (13). Each sample was normalized to the geometric mean of the top 50 most highly expressed miRNAs. Student’s t-test was used to calculate statistical significances of pair-wise comparisons. Calculations were performed using the R statistical computing environment (http://www.r-project.org/).

Gene ontology (GO) analysis

GO analysis was performed using miRNAs with P < 0.05 and fold change of >1.3. For each miRNA, the common target genes found in two or more datasets were identified. The Database for Annotation, Visualization and Integrated Discovery (DAVID) was used to perform GO analysis was then performed for each tissue using DAVID (Database for Annotation, Visualization and Integrated Discovery) (16).

TaqMan miRNA assay

TaqMan miRNA assays were performed to quantify mature miRNAs using pre-designed probes, rat U87 as the normalizer (Applied Biosystems, Foster City, CA), and the competitive Ct method. Real-time PCR was performed on the ABI Prism 7900HT Sequence detection system (Applied Biosystems).

ISH

miRCURY locked nucleic acid (LNA) microRNA detection probes, namely, miro-miR-21, miro-miR-31, lsa-miR-31, negative controls (miro-miR-31 and miro-miR-21) with mismatches at two positions, were purchased from Exiqon (Vedbaek, Denmark). The oligonucleotides are double DIG-labeled at the 5’- and 3’-ends. ISH was performed on 6 µm FFPE sections as described by Nielsen et al. (17). Following deparaffinization, rehydration in graded alcohol and proteinase K treatment, tissue sections were hybridized with miro-miR-31 probe (20µM), or miro-miR-21 probe (50µM) in hybridization buffer (Exiqon) at 57°C for 14h in a hybridizer (Dako, Glostrup, Denmark). Following stringent washes in SSC buffers, the sections were blocked against unspecific binding of the detecting antibody, using DIG wash and blocking reagent. miRNA was localized by incubation with 4-nitro-blue tetrazolium (NBT) and 5-bromo-4-chloro-3‘-indolylphosphate (BCIP; Roche, Mannheim, Germany). Nuclear fast red (Vector Lab., Burlingame, CA) was used as a counterstain.

Quantitative real time PCR

qRT-PCR was performed using predesigned probes (Applied Biosystems), PSMB6 as the normalizer and the comparative Ct method (7).

IHC

IHC was performed on esophageal/lingual sections to assess nuclear programmed cell death 4 (PDCD4, LS-B1388, Lifesiences, Seattle, WA) after citrate-based antigen retrieval as previously described (7). Protein was localized by incubation with 3-amino-9-ethylcarbazole substrate-chromogen (AEC; Dako, Carpinteria, CA).

Immunoblot analysis

Total protein samples were separated by SDS-polyacrylamide gel electrophoresis and transferred onto polyvinylidene difluoride membranes (7). After incubation with primary antibodies: PDCD4 (LS-B1388, Lifesiences), TGM1 (ab55915; Abcam, Cambridge, MA), PPP2R2A (85689, Cell Signaling, Danvers, MA). RHOB/BR (sc-102084; Santa Cruz Biotech, CA), STA2U (sc-87439; Santa Cruz Biotech) and loading control GAPDH (CB1001, Calbiochem, San Diego, CA), the membrane was incubated with appropriate peroxidase-conjugated secondary antibodies. Immunodetection was by Pierce enhanced chemiluminescence substrate (Thermo Scientific, Waltham, MA) and intensity of protein bands quantified by GS-800 calibrated densitometer (BIO-RAD, Hercules, CA).

Zn measurement

Tissue Zn content was determined using Atomic Absorption Spectrometer Analyst 400 (PerkinElmer, Waltham, MA).

Rat and human tongue SCC samples

 Archived rat tongue SCC tissues (FFPE) were derived from a previous study (8). Human tongue SCC tissues (FFPE) were from Thomas Jefferson University Hospital.

Rat ESCC samples

 Archived rat ESCC tissues (FFPE) and RNAs were derived from a previous study (7).

Statistical analysis

Differences in Zn, gene expression and protein levels were assessed with Student’s t-test. Statistical tests were two-sided and were considered significant at P < 0.05.

Results

Prolonged ZD reduces tissue Zn levels and induces overexpression of inflammation genes

Chronic dietary ZD (23 weeks) led to a significant decline in Zn levels across all tissue types (Figure 1A). Zn levels in serum and testis (two widely used indicators of ZD) were 52% and 40% lower in ZD than control ZS rats. Prostate accumulates the most Zn under normal condition (18) showed the greatest decline in tissue Zn levels (~53%), followed by esophagus (~47%), skin (~43%), pancreas (~42%), tongue (~33%) and lung (~26%). Liver showed the least reduction (~19%).

Chronic ZD induced up-regulation of numerous inflammatory genes in the esophagus (7). Whether it also causes inflammation in the other tissue types was evaluated by analyzing the expression of eight cancer-associated inflammation genes (S100a8, S100a9, Cxcl5, Il1b, Cxcl2, Cxcl4, Ptggs and Tlr4) (7), using qRT-PCR. The ZD tongue is highly susceptible to chemically induced carcinogenesis (5) and was the only tissue that showed overexpression of all eight inflammatory genes (Figure 1B). S100a8 and S100a9 were overexpressed in all tissues studied (Figure 1B). In PBMC, the difference between ZD and ZS samples was suggestive but not statistically significant (S100a8, P = 0.07; S100a9, P = 0.08). Immunoblot and enzyme-linked immunosorbent assay (ELISA) analyses showed that the ZD effect observed at the transcript levels for S100a8, S100a9, Cxcl5 and Il1b were also reflected at the protein level in tongue, lung and pancreas (Figure 1C and 1D).

These data establish that prolonged ZD reduces Zn levels across tissue types and up-regulates expression of key inflammation genes.

Normal ZN tissues show high abundance of tissue-specific miRNAs

The nanoString platform directly measures miRNA expression levels without reverse transcription or PCR amplification, thereby eliminating enzymatic bias (13,14). At the time we initiated this study, the nanoString rat miRNA kit was unavailable. Initially, we used the human miRNA assay to profile PBMC and then the mouse miRNA assay to profile esophage, skin, lung, pancreas, liver and...
prostate. Because of a high degree of sequence conservation of miRNAs across human, mouse and rat, 178 and 236 miRNAs on the human and mouse panels, respectively, have sequences identical to the rat miRNAs. Thus, we profiled 178 rat miRNAs in PBMC and 236 rat miRNAs in the other six tissues (n = 6 rats/tissue/dietary group).

The nanoString profiling platform provided an estimate of miRNA abundance in normal ZS and ZD organs. ZS organs showed high abundance of many tissue-specific miRNAs (see Supplementary Table I online). As shown in Figure 2B, the liver-specific miR-122 (19) was abundantly expressed in the liver; skin-specific miR-203 (20) in the skin; epithelial tissue-specific miR-205 in skin and esophagus; pancreatic islet-specific miR-375 (21) and pancreas-specific miR-216 and miR-217 (22) in the pancreas; and the hematopoietic markers miR-142-3p (23), miR-150, miR-16, miR191, miR-26a, miR-19b (24) in PBMC. Conversely, the ubiquitous oncopgenic miR-21 was expressed with moderate abundance across all seven tissues, and oncopgenic miR-31 in low abundance in these same tissues. These miRNA abundance data from normal rat tissues establish that the nanoString technology is both sensitive and specific.

**ZD induces a prooncogenic miRNA signature in inflammatory esophagus**

Using a cutoff point of P-value <0.05 and ≥1.3 fold difference, we identified 30 dysregulated miRNAs in the inflammatory-deficient esophagus (23 up- and 7 down-regulated) (Figure 2C, see Supplementary Table II online). Importantly, the ZD esophagus had a distinct miRNA signature that resembles the miRNAomes of human ESCC and tongue SCC (25–33). This signature was defined by up-regulation of five oncopgenic miRNAs (miR-31, 3.0 fold; miR-21, 2.7 fold; miR-223, 2.5 fold; miR-142-3p, 2.4 fold; miR-221, 1.6 fold) and down-regulation of two tumor suppressors (miR-204, −2.5 fold; miR-375, −2.0 fold). Because these same miRNAs are aberrantly expressed in human ESCC and tongue SCC (25–33), prolonged ZD induces a miRNA expression profile in the rat esophagus that in human tissue is predictive of ESCC development.

We then determined whether the top-up-regulated miR-31 and miR-21 in ZD esophagus are also overexpressed in the highly tumorigenic (5) and inflammatory ZD tongue (Figure 1, B-D). Using Taqman miRNA assay (Figure 3A), both oncomiRs were significantly overexpressed in ZD tongue as compared to ZS counterpart (miR-31, 2.6 fold, P = 0.015; miR-21, 3.9 fold, P = 0.014; n = 6 rats/dietary
Zn deficit induces oncogenic miRNAs in esophagus

Table I. The miRNA target biological processes enrichment in Zn-deficient tissues by gene ontological (GO)

| Tissue | GO terms                              | No. of genes | En* Score | Adjusted P-value |
|--------|---------------------------------------|--------------|-----------|-----------------|
| Esophagus | Zinc ion binding family               | 119          | 3.4       | 0.010           |
|         | Positive regulation of Cell           | 67           | 2.9       | 0.011           |
|         | Differentiation                       |              |           |                 |
|         | Serine/threonine-protein kinase 10    | 59           | 1.8       | 0.021           |
|         | family                                |              |           |                 |
|         | MAPKK and JNK family                  | 42           | 1.4       | 0.041           |
|         | Negative regulation of actin filament | 31           | 1.3       | 0.042           |
|         | family                                |              |           |                 |
|         | Epithelial cell proliferation         | 22           | 0.9       | 0.049           |
| Skin    | ATP calmodulin pathway                | 56           | 4.1       | 0.018           |
|         | Positive regulation of Cell           | 67           | 2.9       | 0.011           |
|         | Differentiation                       |              |           |                 |
|         | Serine/threonine-protein kinase 10    | 59           | 1.8       | 0.021           |
|         | family                                |              |           |                 |
|         | MAPKK and JNK family                  | 42           | 1.4       | 0.041           |
|         | Negative regulation of actin filament | 31           | 1.3       | 0.042           |
|         | family                                |              |           |                 |
|         | Epithelial cell proliferation         | 22           | 0.9       | 0.049           |
| Lung    | Transcription regulation by RNA-pol II| 187          | 3.1       | 0.015           |
|         | Negative regulation of biosynthesis pathways | 103    | 2.9       | 0.012           |
|         | Zinc ion binding family               | 68           | 1.4       | 0.037           |
|         | Lung development                      | 23           | 1.0       | 0.046           |
| Pancreas | Positive regulation of anti-apoptosis genes | 79      | 2.1       | 0.024           |
|         | Insulin signal pathway                | 68           | 1.2       | 0.035           |
|         | Zinc ion binding proteins             | 20           | 0.9       | 0.046           |
| Liver   | Zinc ion binding family               | 101          | 4.1       | 0.023           |
|         | Glicosyl-lysine family                | 93           | 2.1       | 0.028           |
|         | Post-transcription regulation         | 49           | 1.7       | 0.033           |
| Prostate | Magnesium metabolism family           | 77           | 6.4       | 0.021           |
|         | ATP biosynthetic process              | 63           | 5.5       | 0.022           |
|         | Endocytosis                           | 39           | 4.9       | 0.033           |
|         | Transcription regulatory activity      | 21           | 3.4       | 0.036           |
|         | TGB-beta signaling pathway            | 22           | 2.0       | 0.040           |
|         | Zinc finger domain family             | 19           | 1.4       | 0.046           |
| PBMC    | Zinc finger family                    | 115          | 3.0       | 0.035           |
|         | Ubiquitin-conjugating enzyme          | 82           | 2.8       | 0.039           |
|         | Proteolysis – cellular protein catalytic process | 79  | 1.7       | 0.040           |
|         | Protein-lysin-N-methyltransferase activity | 62  | 1.4       | 0.045           |

* Enrichment Score.

Fig. 2. miRNA expression profiles of ZD esophagus and six additional tissues. (A) Heat maps with supervised clustering of miRNAs expression in seven tissues. Red, green and black colors represent high, low and mean expression levels, respectively (n = 6 rats/dietary group/tissue, except PBMC: n = 7 (ZD) and 5 (ZS); P < 0.05). (B) Normal ZS tissues showing tissue-specific miRNA expression in liver, skin, pancreas and PBMC. (C) Barplots showing fold change of miRNA expression in 7 ZD versus ZS tissues (cutoff of P < 0.05, fold change ≥ 1.3).

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**Fig. 3.** Validation of nanoString miRNA profiling data. (A) Validation of 14 representative miRNAs in various ZD tissues and expression of miR-31 and miR-21 in ZD tongue by TaqMan miRNA assays. (B) ISH localization of miR-31 and miR-21 in esophagus and tongue (23 weeks ZD). miR-31 signal (arrow heads, blue, NBT/BCIP; counterstain, nuclear fast red) was strong/abundant in hyperplastic ZD esophageal epithelium and tongue epithelia versus moderate/infrequent in control ZS tissues. miR-21 signal (blue, arrows) was moderate in basal and stromal cells in ZD tissues but infrequent in ZS tissues. No ISH signal was seen with mm-miR-31 or mm-miR-21 (two mismatches). Scale bar, 25 µm. miRNA was localized by 4-nitro-blue tetrazolium (NBT) and 5-brom-4-chloro-3-indolylphosphate (BCIP). (C) IHC showing reduction of PDCD4 protein expression (nuclear) in ZD esophagus and tongue versus ZS tissues. (D) Immunoblots show down-regulation of D′ or mm-miR-21 (two mismatches). Scale bar, 25 µm. miRNA was localized by 4-nitro-blue tetrazolium (NBT) and 5-brom-4-chloro-3-indolylphosphate (BCIP).

**Liver.** Among the 22 dysregulated miRNAs identified in ZD liver, only oncogenic miR-21 (1.5 fold) has been reported in human hepatocellular cancer (39).

**Prostate.** Zn is important for prostate health. The pathogenesis of prostate cancer involves the transformation of normal prostate epithelial cells that accumulate Zn in malignant prostate cells that do not accumulate Zn (18). ZD prostate that showed the greatest decline in Zn levels (Figure 1A) had 41 down-regulated miRNAs. Several tumor suppressors, including miR-184 (1.5 fold), miR-488 (1.5 fold) and miR-330 (1.3 fold), are hallmarks of the human prostate cancer miRNAome (40–42).

**PBMC.** PMBC are leukocytes that have a round nucleus such as lymphocytes (T cells, B cells), natural killer cells and monocytes. Circulating miR-31 was the top-up-regulated species (1.8 fold) in PBMC from ZD versus ZS rats. Because miR-31 was also overexpressed in ZD esophagus, tongue and skin, this result suggests an association between circulating and tissue miR-31. miR-150 was the most down-regulated species in PBMC of ZD rats (–5.1 fold). In patients with sepsis, miR-150 levels in PBMC and plasma were significantly reduced (43). In this regard, patients with sepsis are frequently ZD (44).

To summarize, dietary ZD induces dysregulation of key oncogenic and tumor-suppressor miRNAs in the esophagus and across all tissues profiled. The most shared miRNAs in signatures of the ZD tissues were oncogenic miR-21 and miR-31 (see Supplementary Table III online), with miR-21 up-regulated in six (esophagus, pancreas, liver, skin, lung, tongue) and miR-31 in four tissues (esophagus, skin, PBMC, tongue).

**Validation of nanoString profiling data**

To validate the nanoString results, we performed Taqman miRNA assays on 14 selected miRNAs (n = 6 rats/tissue/dietary group). In all 14 cases, the same pattern and fold changes response to dietary ZD was observed using Taqman as with the nanoString assay (Figure 3A).
miR-21 and miR-21 overexpression correlates with down-regulation of their tumor-suppressor targets

In human tongue SCC, a correlation of miR-21 overexpression with loss of its tumor-suppressor targets PDCD4 (30) and TPM1 (45) mRNA and protein expression is documented. We determined if up-regulation of oncogenic miR-21 in ZD tissues is accompanied by down-regulation of these two known tumor-suppressor targets. Using IHC, PDCD4 was strongly expressed in the nuclei of epithelial cells of ZS esophagus and tongue. This nuclear staining of PDCD4 was reduced or absent in the hyperplastic ZD esophagus and tongue (Figure 3B). In contrast, miR-21 ISH signals were frequent and found mostly in stromal calls of ZD tissues and were generally scarce in ZS tissues (Figure 3B). These data establish that miR-31 and miR-21 expression is cell-type specific with miR-31 localized to epithelial and miR-21 mostly in stromal cells.

Overexpression of miR-31 and miR-21 is associated with ESCC in ZD rats

ZD induced an inflammatory gene-expression profile that fueled ESCC in rats with repeated exposure to NMBA (7). Using qRT-PCR on archived RNA samples (7), miR-31 and miR-21 exhibited 5.4 and 3.2-fold up-regulation in ESCC-bearing ZD rats versus cancer-free ZS controls (Figure 4A). ZR, a regimen that prevented ESCC development (7) reduced their expression to ZS levels (Figure 4A). ISH on archived FFPE tissues (7) showed reduced expression of miR-21, miR-31, miR-150, miR-223 and down-regulation of miR-375, miR-204, skin (up-regulation of miR-21, miR-31, miR-203; down-regulation of miR-150), lung (up-regulation of miR-21, miR-31, miR-203), pancreas (up-regulation of miR-21 and down-regulation of miR-217, miR-216a, miR-216b-5p) and prostate (down-regulation of miR-330, miR-148, miR-184, miR-147). Thus, Taqman miRNA data confirm the accuracy of the nanoString data.

Cellular localization of miR-31 and miR-21 in ZD esophageal and tongue epithelia

We focused our study on the top-up-regulated miR-31 and miR-21 because of their roles in human ESCC and OSCC (27,29–32). Their cellular localizations in esophageal and tongue preneoplastic tissues after chronic dietary ZD (23 weeks) was defined by performing ISH on FFPE tissues using high affinity double Dig-labeled LNA probes (Exiqon, Vedbaek, Denmark; n = 10 diets/group). The hyperplastic ZD esophagus and tongue epithelial cells showed abundant miR-31 ISH signal, as compared with nonproliferative ZS tissues (Figure 3B). In contrast, miR-21 ISH signals were infrequent and found mostly in stromal calls of ZD tissues and were generally scarce in ZS tissues (Figure 3B). These data establish that miR-31 and miR-21 expression is cell-type specific with miR-31 localized to epithelial and miR-21 mostly in stromal cells.

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Expression of miR-31 and miR-21 in human tongue SCC

The cellular localization of miR-31 and miR-21 in human tongue SCC was determined using ISH (n = 5 cases). As in rat tongue SCC, all five cases of human tongue SCC showed prevalent expression of miR-31 in tumor cells and miR-21 expression in stromal cells (Figure 5, SCC human tongue). The ISH data in human and rat tongue SCC (Figure 5) and rat ESCC (Figure 4B) establish that miR-31 and miR-21 expression in SCC is cell-type specific, with miR-31 localized to tumor cells and miR-21 to tumor stroma.

GO analysis

To gain an understanding of the molecular pathways that are governed by the predicted targets of dysregulated miRNAs in ZD tissues, we performed biological processes enrichment using GO (48). Using target genes found in common in more than two databases (RNAhybrid, TargetScan, miRDB and microRNA.org), we performed GO analysis using DAVID (16) for each of the seven tissues (cutoff expression in esophageal cancer development and prevention; and their dysregulation by ZD is associated with ESCC.

Zn therapy reduces miR-31 and miR-21 expression in rat tongue SCC

Zn supplementation in rats decreased 4-nitroquinoline1-oxide (NQO)-induced tongue SCC incidence and elicited a reduced proliferative/inflammatory cancer phenotype (8). We asked if Zn supplementation that suppresses tongue cancer development also attenuates miR-31 and miR-21 expression. ISH analysis was performed on FFPE tissues from NQO-induced rat tongue SCCs with or without Zn therapy (8) in ZR (n = 10 rats and Zn treatment group). In near serial sections of these tumor samples (Figure 5, SCC rat tongue), miR-31 expression was localized to tumor cells, whereas miR-21 was predominantly found in stromal cells. Importantly, ISH signals of both oncomiRs were reduced in intensity and scope following Zn therapy (Figure 5, ZnSup vs ZS), thereby showing that their expression is responsive to Zn therapy.

Expression of miR-31 and miR-21 in human tongue SCC

The cellular localization of miR-31 and miR-21 in human tongue SCC was determined using ISH (n = 5 cases). As in rat tongue SCC, all five cases of human tongue SCC showed prevalent expression of miR-31 in tumor cells and miR-21 expression in stromal cells (Figure 5, SCC human tongue). The ISH data in human and rat tongue SCC (Figure 5) and rat ESCC (Figure 4B) establish that miR-31 and miR-21 expression in SCC is cell-type specific, with miR-31 localized to tumor cells and miR-21 to tumor stroma.

GO analysis

To gain an understanding of the molecular pathways that are governed by the predicted targets of dysregulated miRNAs in ZD tissues, we performed biological processes enrichment using GO (48). Using target genes found in common in more than two databases (RNAhybrid, TargetScan, miRDB and microRNA.org), we performed GO analysis using DAVID (16) for each of the seven tissues (cutoff...
expressed miRNAs are down-regulated in the public ESCC/OSCC
SCC; mm-miR-31 and mm-miR-21); 50 µm (miR-31 and miR-21; H&E,
chloro-3′
No ISH signal was seen with mm-miR-31 or mm-miR-21 (2 mismatches).
tumor cells (blue, arrow heads), and miR-21 signal in stroma (blue, arrows).
but reduced/infrequent in stromal cells of ZnSuppl tongue SCC. Two cases of
proliferative tongue SCC from ZS rats but moderate/infrequent in tumor cells
value of ≥1.3 fold-change and

Fig. 5. ISH localization of miR-31 and miR-21 in archived rat and human
tongue SCCs. Rat tongue SCC; miR-31 signal (arrow heads; blue, NBT/BCIP; counterstain, nuclear fast red) was strong/abundant in tumor cells of
proliferative tongue SCC from ZS rats but moderate/infrequent in tumor cells
of regressing tongue SCC from Zn-supplemented (ZnSuppl) rats. miR-21
signal (blue, arrows) was strong/abundant in stromal cells of ZS tongue SCC
but reduced/infrequent in stromal cells of ZnSuppl tongue SCC. Two cases of
human tongue SCC (H&E) are shown: miR-31 signal was predominantly in
tumor cells (blue, arrow heads), and miR-21 signal in stroma (blue, arrows).
No ISH signal was seen with mm-miR-31 or mm-miR-21 (2 mismatches).
miRNA was localized by 4-nitro-blue tetrazolium (NBT) and 5-brom-4-
chloro-3′-indolylophosphate (BCIP). Scale bars: 100 µm (H&E, rat tongue
SCC; mm-miR-31 and mm-miR-21), 50 µm (miR-31 and miR-21; H&E,
human tongue SCC).

of ±1.3 fold-change and P-value < 0.05). A survey of overrepresented
GO terms documented a bias toward genes involved in Zn binding
and gene regulatory processes (Table I). In particular, ZD esophagus
showed overrepresented processes important in inflammation and cel-
lular proliferation, including serine/threonine-protein kinase activity,
MAPK and JNK cascade and epithelial cell proliferation.

In addition, we performed functional classification of miRNA
targets of up- and down-regulated miRNAs in ZD esophagus (see
Supplementary Table IV online). The biological processes targeted by
the 23 up-regulated miRNAs include processes in inflammation and
cell proliferation, including serine/threonine-protein kinase activity,
PPP2R2A and MAP2K4 [target of 

miR-31
]; P = 2.27E-08); regulation of cell
motion (27 genes, P = 3.53E-07); and inflammatory response (seven
genes, P = 0.028); negative regulation of gene expression (41 genes,
including tumor suppressor PDCD4; P = 1.10E-04). Importantly, our
present study documented a correlation between overexpression of
miR-21 ISH signal (Figure 3B) in the inflammatory ZD esophagus/
tongue and loss of expression of its tumor-suppressor target PDCD4 in the
same tissues (Figure 3C). In addition, functional classification of
targets of the seven down-regulated miRNAs showed overpresen-
tation of biological processes such as transcription regulation, nega-
itive regulation of gene expression and Zn ion binding.

To determine if these predicted targets of dysregulated miRNAs
are relevant to human ESCC development, we retrieved differen-
tially expressed genes (tumor versus normal tissue, P-value < 0.05)
from human ESCC/OSCC gene-expression data using ArrayExpress
and Gene Expression Omnibus databases. All the selected genes
(112 down-regulated and 38 up-regulated, see Supplementary Table
V online) belong to the functional families of the predicted targets
of dysregulated miRNAs (see Supplementary Table IV online).
Interestingly, 47.5% (112 of 236) of our predicted targets for the over-
expressed miRNAs are down-regulated in the public ESCC/OSCC
gene expression database as compared to 29% (38 of 133) of our pre-
predicted targets for the down-regulated miRNAs. Together, our bioin-
formatics analysis provides a repertoire of putative target genes (see
Supplementary Table V online) related to the altered miRNAome of
the inflammatory ZD esophagus that may prove to be highly relevant
to ESCC development.

Discussion
Dietary ZD increases the risk of ESCC. In a well-established rat model,
prolonged ZD causes overexpression of numerous cancer-associated
inflammation genes in the esophagus, thus establishing an inflam-
matory microenvironment that fuels ESCC (7). In the present study,
we show that this inflammatory ZD esophagus exhibits a protu-
morogenic miRNA signature resembling the miRNAomes of human
ESCC or OSCC (25–33), with overexpression of five oncogenic
miRNAs (miR-31, miR-21, miR-223, miR-142-3p, miR-221) and
down-regulation of two tumor suppressors (miR-204, miR-375). The
data suggest a connection between dysregulation of specific miRNAs
and ZD-induced chronic inflammation in esophageal preneoplasia.

In particular, miR-31 and miR-21, the top-up-regulated species in
the ZD esophagus, are reported to be elevated in many cancers and
inflammation-associated diseases. miR-21 is one of the most con-
sistently up-regulated oncomiRs in human cancers, and is thought
to contribute to inflammation-associated diseases, including cancers
(12). miR-31 is a pleiotropically acting miRNA that is up-regulated
in many types of human cancers, including ESCC, tongue SCC, liver
cancer and colorectal cancer, but down-regulated in breast metastasis
(49). Interestingly, miR-31 expression levels are reported to increase
during inflammatory bowel disease-associated neoplastic transforma-
tion (50). Our data show that miR-21 overexpression in inflamma-
tory ZD esophagus and tongue is correlated with down-regulation of
its tumor-suppressor targets PDCD4 and TPM1 (Figure 3, B-D). In
human ESCC, miR-21 negatively regulates PDCD4 and knockdown
of miR-21 inhibits cellular proliferation and invasion (51). In ESCC
tissues, loss of PDCD4 protein expression is correlated with cancer
aggressiveness (tumor stage, nodal metastasis) (52). In tongue SCC
tissues, miR-21 may act as an antiapoptotic factor by blocking TPM1
expression (45). Although miR-31 overexpression in ZD esophagus
and tongue (Figure 3B) is accompanied by down-regulation of its
known tumor-suppressor target PPP2R2A (46) (Figure 3D), the
role of PPP2R2A down-regulation has not been reported in human
ESCC or OSCC. We speculate that the down-regulation of such
tumor-suppressor targets may relate to the increased cell proliferation
and inflammation associated with ZD, as suggested by our GO enrich-
ment analysis of targets of up-regulated miRNAs (see Supplementary
Table IV online). The biological role of miR-21 and miR-31 in
ZD-induced inflammation-associated esophageal carcinogenesis
remains to be determined.

Significantly, our present Taqman and ISH data show that miR-31
and miR-21 levels were substantially higher in ZD esophagus bearing
ESCC versus cancer-free ZS controls (Figure 4 versus Supplementary
Figure 1 online). Replenishing Zn reversed the inflammatory gene
signature (7) and reduced miR-31 and miR-21 levels to ZS levels
(Figure 4A, see Supplementary Figure 1 online). Thus, the different
ESCC outcome in Zn-modulated animals are directly associated with
their elevated and reduced miR-31 and miR-21 levels. The data estab-
lish for the first time that dietary Zn regulates miR-31 and miR-21
expression in ESCC development and prevention.

The tumor-suppressive miR-375 is known to circulate at low levels
in ESCC patients (25), and Ryu et al. (53) showed its acute rever-
sal in serum of Zn-depleted human subjects following a Zn-repletion
regimen. The present ISH data that miR-31 and miR-21 expres-
sion was concomitantly reduced in regressing tongue SCCs from
Zn-supplemented rats (8) indicate that these oncomiRs are responsive
to Zn therapy (Figure 5). Thus, Zn supplementation may be thera-
peutic in combating ESCC or OSCC by correcting aberrant miRNA
expression. Although Zn therapy inhibits tongue cancer develop-
ment by suppressing inflammation and cellular proliferation (8), the

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mechanism whereby Zn decreases miR-31 and miR-21 expression in tongue cancer inhibition needs to be further investigated.

The cellular origins of individual miRNAs are of critical importance in the understanding of the mechanistic roles of miRNAs in tumor development. Whereas miR-21 presence in stromal cells of human colorectal (17) and breast cancer (54) was previously reported, the localization of miR-31 to esophageal/tongue preneoplastic epithelia (Figure 3B) and tumor cells (Figure 4B and Figure 5) is reported here for the first time. Thus, miR-31 and miR-21 expression is cell-type specific and may serve as markers in tumor diagnosis and prognosis. In addition, the profiling result that miR-31 was up-regulated in ZD esophagus and tongue, as well as in PBMC from ZD animals, indicates that miR-31 may serve as a noninvasive diagnostic marker for a pre-malignant esophageal/tongue condition. In fact, patients with ESCC or OSCC display high levels of miR-31 (31,32) as well as miR-21 (25,30) in tumor tissues and in serum/plasma. Conversely, miR-21 that was overexpressed in most ZD tissues was not up-regulated in PBMC from ZD animals. This result requires further exploration to determine whether the miR-21 overexpression in PBMC could arise at later stages of ESCC or OSCC development due to elevation of miR-21 expression in the tumor stroma (Figure 4B and Figure 5), as compared to stroma of hyperplastic tissues (Figure 3B).

The miRNAomes of the other six inflammatory ZD tissues (Figure 1, B-D) showed dysregulation of specific oncogenic and tumor-suppressor miRNAs (Figure 2C). These changes resemble the hallmarks of the miRNAomes of various human conditions, such as pancreatitis and pancreatic ductal adenocarcinoma (22,38), liver cancer (39), inflammatory COPD and lung cancer (37,38), prostate cancer (40–42), inflammatory human psoriatic skin (34) and PBMC of patients with sepsis (43). Interestingly, patients with these inflammation conditions, namely psoriasis (35), COPD and sepsis (44) are frequently ZD. Thus, ZD modulation of expression of specific miRNAs has implications with regard to the role of ZD in the pathogenesis of these diseases.

With the exception of red meat and seafood, the Zn content in most foods is low. Individuals subsisting largely on a cereal diet are likely to be ZD. Global dietary ZD is estimated to affect 30% of the population, or 4% to 73% across subregions, with higher frequencies occurring in developing countries (55). The mechanistic link to ZD in most tumor conditions, namely psoriasis (35), COPD and sepsis (44) are frequently ZD. Thus, ZD modulation of expression of specific miRNAs has implications with regard to the role of ZD in the pathogenesis of these diseases.

In summary, our present data show that prolonged dietary ZD causes aberrant miRNA expression in a wide variety of tissues associated with inflammation, suggesting a likely mechanism contributing to the burden of human diseases associated with dietary ZD. Importantly, the demonstration of the dysregulation of miR-31 and miR-21 by ZD in inflammatory esophageal/tongue neoplasia provides new insight into the mechanisms whereby ZD promotes human ESCC and OSCC.

Supplementary material
Supplementary Tables I, II, III, IV, V and Supplementary Figure 1 can be found at http://carcin.oxfordjournals.org/

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