Particle, a Triplex-Forming Long ncRNA, Regulates Locus-Specific Methylation in Response to Low-Dose Irradiation

Highlights

- Expression of tumor suppressor MAT2A is regulated by PARTICLE via triplex formation

- PARTICLE interacts with silencing complexes G9a and PRC2

- PARTICLE and MAT2A are candidate biomarkers in patient plasma post-radiotherapy

- Inverse dose response of PARTICLE challenges the radiation linear non-threshold rule

Accession Numbers

GSE67008

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In Brief

O’Leary et al. find a long non-coding RNA called PARTICLE that is overexpressed following irradiation. PARTICLE represses a tumor suppressor MAT2A via triplex formation and interaction with the polycomb repressor complex. PARTICLE also acts as a cytosolic scaffold for MAT2A in preparation for exosomal transport from the cell.
PARTICLE, a Triplex-Forming Long ncRNA, Regulates Locus-Specific Methylation in Response to Low-Dose Irradiation

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http://dx.doi.org/10.1016/j.celrep.2015.03.043
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SUMMARY

Exposure to low-dose irradiation causes transiently elevated expression of the long ncRNA PARTICLE (gene PARTICLE, promoter of MAT2A-antisense radiation-induced circulating IncRNA). PARTICLE affords both a cytosolic scaffold for the tumor suppressor methionine adenosyltransferase (MAT2A) and a nuclear genetic platform for transcriptional repression. In situ hybridization discloses that PARTICLE and MAT2A associate together following irradiation. Bromouridine tracing and presence in exosomes indicate intercellular transport, and this is supported by ex vivo data from radiotherapy-treated patients. Surface plasmon resonance indicates that PARTICLE forms a DNA-IncRNA triplex upstream of a MAT2A promoter CpG island. We show that PARTICLE represses MAT2A via methylation and demonstrate that the radiation-induced PARTICLE interacts with the transcription-repressive complex proteins G9a and SUZ12 (subunit of PRC2). The interplay of PARTICLE with MAT2A implicates this IncRNA in intercellular communication and as a recruitment platform for gene-silencing machineries through triplex formation in response to irradiation.

INTRODUCTION

The abundance of long non-coding RNAs (lncRNAs) in the genomes of higher organisms contributed to their initial relegation as irrelevant transcriptional noise emanating from RNA polymerase II infidelity (Struhl, 2007). Defined as RNA transcripts ranging in length from 200 up to ~100,000 bp lacking a significant open reading frame (Gertone et al., 2004), thousands of lncRNAs have been identified by computational transcriptome prediction (Amaral et al., 2011). Recognition of their functional importance arose from the demonstration of their participation in genomic transcriptional control, with influence ranging from a single locus (Wan et al., 2013) to entire chromosomal regions (Chaumeil et al., 2006; Zhang et al., 2007). Regulation may be through association with chromatin complexes in cis (Azzalin et al., 2007; Schoeftner and Blasco, 2008) or trans (Gupta et al., 2010; Rinn et al., 2007; Tsiol et al., 2010). However, for most lncRNAs, their physiological function remains elusive, with some indications of involvement in disease (Huarte and Rinn, 2010) or as responders to genotoxic insults (Ozgür et al., 2013).

Ionizing radiation instigates direct macromolecular damage as well as indirect (non-targeted) cellular and tissue stress (Morgan and Sowa, 2015; Pluder et al., 2011). While the DNA damage response is linear with the applied dose, ancillary effects may deviate significantly from linearity (Kadhim et al., 2013). Crucially, these non-targeted reactions to radiation may be more prominent than the direct damage responses at low doses, i.e., milligray range exposure typically encountered in the workplace, during medical imaging, and from natural sources. Thus, low doses may evoke alternative biological responses compared to those emanating from medium- or high-dose exposure (Mullenders et al., 2009; Waldren et al., 2004).

The molecular players known to be involved in the radiation response include lncRNAs and the polycomb repressor complex 2 (PRC2), as mediators coordinating cellular repair (Campbell et al., 2013; Wang et al., 2008). While the mechanistic role of PRC2 itself in DNA damage signaling must be elucidated,
In lncRNAs are becoming recognized as important participants in PRC2 recruitment with their tertiary structure key to specific target gene recognition (Margueron and Reinberg, 2011). LncRNAs such as lncRNA-p21 and PANDA are upregulated by DNA damage as direct targets of p53 (Hung et al., 2011). Regulation of Cyclin D1 (CCND1) following ionizing radiation has been demonstrated as a target for promoter-derived lncRNAs, e.g., ncrNA-CCND1s (Wang et al., 2008). ANRIL (anti-sense non-coding RNA in the INK4 locus) is overexpressed following high-dose irradiation (5 Gy exposure), which may be related to suppression of senescence by p16 (Ozgur et al., 2013). However, limited knowledge exists on the mechanistic response of lncRNAs to ionizing radiation and, in particular, the influence of dose and cellular context (Ozgur et al., 2013).

Here we introduce the lncRNA PARTICLE (HUGO gene nomenclature PARTICLE, promoter of MAT2A)-antisense radiation-induced circulating lncRNA, NCB1 reference sequence NR_038942.1), which demonstrates a substantial increase in expression after low-dose radiation. PARTICLE is located within the MAT2A gene that encodes the catalytic subunit of methionine adenosyltransferase (MAT), the enzyme responsible for the production of s-adenosylmethionine (SAM), which is the principal methyl donor of the cell (Mato et al., 1997). Co-localization of PARTICLE and MAT2A cytosolic transcripts potentially destined for export via exosomes in response to irradiation is demonstrated. We provide evidence that nuclear PARTICLE is a suppressor of MAT2A through triple helix formation and provides a methyltransferase and polycomb repressor complex recruitment platform. Broadly, our work identifies a potential mammalian riboswitch previously discovered in bacteria, and it suggests that lncRNAs subsequently have been exploited for roles in metabolomic regulation.

RESULTS

PARTICLE Expression Is Modulated by Exposure to Low-Dose Irradiation

Microarray analysis provided an initial platform for target discovery. While ~50 radiation-regulated candidates were identified (Table S1), four lncRNAs (NR_027405, BC036914, BX647881, and PARTICLE [NR_038942.1]) associated with genes (MTHFD2, GNA11, TFDP2, and MAT2A, respectively) were selected for further verification by Taqman qPCR assays (see Table S2 for primers and probes). Fold differences in the relative expression of these lncRNAs as well as PANDA (NR_109836.1, serving as a positive indicator of radiation exposure [Hung et al., 2011]) were initially assessed in T47D, a mammary cancer cell line (due to previous inclusion of this cell line in the high-throughput microarray analysis), at 4 and 24 hr after exposure to 2.5 Gy and in sham-irradiated controls (Figure 1A). No significant changes in expression were found for any of these lncRNAs at 4 hr after 2.5 Gy exposure (Figure 1A; p > 0.05). However, at 24 hr, significant increases were noted for the lncRNAs BX647881 and PARTICLE, BX647881, associated with TFDP2, increased by 3.48 ± 0.11 relative to sham-irradiated controls (p = 0.001). There was a more substantial increase of PARTICLE, associated with MAT2A, at 24 hr after 2.5 Gy irradiation (12.8 ± 2.6-fold compared to controls, p = 0.02, as tested in three independent experiments).

These lncRNA expression findings in the relatively radiation-resistant T47D cell line (Anastasov et al., 2012) prompted their further assessment in MDA-MB-361, a radiation-sensitive mammary cancer cell line (Anastasov et al., 2012; Figure 1B). This analysis highlighted different transcriptional responses for these lncRNAs between T47D and MDA-MB-361 and also may reflect alternative RNA steady-state levels. LncRNAs BC036914, PANDA, BX647881, and PARTICLE showed significantly enhanced transcription following irradiation (Figure 1B). A robust elevation in PARTICLE (27 ± 0.64-fold increase versus control values normalized to unity) was noted, especially 24 hr after 0.25 Gy low-dose irradiation exposure in MDA-MB-361 (p = 0.00028; Figure 1B). Increased levels for PARTICLE at 4 hr (12 ± 0.1-fold, p = 0.00056; Figure 1B) also were noted. While unchanged PARTICLE levels were found in sham-irradiated and 2.5 Gy-irradiated samples when analyzed at 4 hr, increased transcription occurred by 24 hr after 2.5 Gy in both T47D and MDA-MB-361 (p < 0.05; Figures 1A and 1B). To determine whether the increase in PARTICLE transcription due to irradiation was solely restricted to cancer cells, its expression also was assessed in three non-cancerous human cell lines (MCF10A, HUVEC, and HEK293). Enhanced levels of PARTICLE following low-dose irradiation were found in all three additional cell lines examined (Figures S1A–S1C), confirming PARTICLE to be a ubiquitous tissue responder to low-dose irradiation exposure, irrespective of malignant status or lineage.

A Relationship of PARTICLE with the MAT2A Gene Is Revealed by the Time Course of the Transcript Expression Pattern following Low-Dose Irradiation

Real-time qPCR analysis of PARTICLE, MAT2A, and GGCX transcription was undertaken at 4, 24, 48, and 72 hr after an acute exposure to 0.25 Gy or 2.5 Gy in MDA-MB-361 cells (Figures 1C–1E). Following 0.25 Gy irradiation, results showed increased PARTICLE transcript presence at 4 hr (8.79 ± 0.5-fold increase) in comparison with sham-irradiated experimental controls (p = 0.004). PARTICLE transcription continued to rise, peaking at 24 hr (20.17 ± 1.6-fold greater than controls, p = 0.006; Figure 1C). By 48 hr, PARTICLE transcript levels decreased back down toward control cell levels. Of note was the slightly different PARTICLE expression profile after the higher (2.5 Gy) dose of radiation. Levels were comparable to controls at 4 hr (p > 0.05), with a peak increase of 8.9 ± 1.2-fold at 24 hr (p = 0.03) that diminished 5 ± 0.38-fold by 48 hr (p = 0.018). The transcriptional decrease continued to 72 hr, reaching non-significance by that time (p = 0.2; Figure 1C).

The expression of MAT2A transcripts was analyzed due to the genomic co-localization of the PARTICLE and MAT2A genes. Dramatic elevation of MAT2A levels was determined at 4 hr post-irradiation (Figure 1D), while from 24 hr onward to 72 hr MAT2A transcripts in 0.25- and 2.5 Gy-irradiated samples showed levels comparable to sham-irradiated controls. There was a significant 6.2 ± 0.9 or 6.9 ± 2.3-fold increase over controls in MAT2A transcription following exposure to 0.25 Gy or 2.5 Gy irradiation (p = 0.001, p = 0.03, respectively) at this time point (Figure 1D). The proximity of GGCX to the PARTICLE/MAT2A locus prompted the assessment of potential variation in GGCX transcription during the radiation response.
Modest fluctuations in GGCX transcription were found after 0.25 Gy exposure (Figure 1E). As there was only a modest relationship with exposure or PARTICLE, GGCX was not followed further in this study.

**Increased PARTICLE Expression in Both the Nucleus and Cytoplasm Is Associated with MAT2A**

In situ hybridization utilizing fluorescently labeled FAM probes complementary to PARTICLE in MDA-MB-361 confirmed that low-dose irradiation induced elevated PARTICLE levels (Figures 2A–2C). Substantially increased intensity of PARTICLE staining was noted in both the nucleus and cytoplasm 24 hr following 0.25 Gy irradiation (Figures 2A–2C, middle). At the same time point, elevated levels of PARTICLE also were noted in these cells after 2.5 Gy exposure compared to controls (Figures 2A–2C, top), although transcripts appeared to be mainly restricted to the cytoplasm with levels not reaching those seen following the lower dose of irradiation. In sham-irradiated cells, PARTICLE (green fluorescence) was spatially discreet from MAT2A (red fluorescence) transcripts, with both the nucleus and cytoplasm having minimum co-localization (yellow) (Figures 2D–2F). Intriguingly, PARTICLE and MAT2A transcripts were seen to associate together as early as 4 hr after low-dose irradiation. Regions-of-interest analysis highlighted the increase in the co-localization coefficient in both nuclear and cytoplasmic zones (Figures 2D–2F).

Bromouridine RNA Tracing, Exosomes, and Radiotherapy Patient Plasma Analysis Support the Intercellular Transport of PARTICLE following Irradiation

Following transwell relocation, extraction, and north-western detection, RNA labeled with 5-bromouridine (BruU) was noted in recipient MDA-MB-361 exposed for 4 or 24 hr to BruU-labeled irradiated (0.25 Gy) donor cells (Figures 3A and 3B). An increase in the size range of Bru-labeled RNA with time was noted. 

*Figure 1. Characterization of IncRNA Transcripts following Low-Dose Irradiation*

(A) Histogram shows IncRNAs (NR_027405, BC036914, PANDA, BX647881, and PARTICLE) expression in T47D, 4 and 24 hr after exposure to 2.5 Gy. Genes associated with each IncRNA are indicated.

(B) Histogram shows IncRNAs (NR_027405, BC036914, PANDA, BX647881, and PARTICLE) expression plus associated genes (as above) in MDA-MB-361 (radiation-sensitive cell line) 4 and 24 hr after exposure to 0.25 or 2.5 Gy irradiation.

(C–E) Schematic diagram (top) and time course (bottom) of the relative expression of PARTICLE (red, C), MAT2A (red, D), and GGCX (red, E) 4–72 hr following either 0.25 or 2.5 Gy irradiation in MDA-MB-361 with values at 0 Gy taken as a value = 1 (dashed line) for comparative purposes. Data are represented as mean ± SEM with significance represented by asterisks (p < 0.05) where appropriate. See also Figure S1 and Tables S1 and S2.
PARTICLE was detected by northern blotting of Bru-labeled RNA in recipient cells (Figure 3B).

PARTICLE levels were significantly elevated in exosomes isolated from the media of MDA-MB-361 exposed to 0.25 Gy (1.8 ± 0.05-fold over sham-irradiated controls = 1, p = 0.0008) and to a lesser extent following 2.5 Gy exposure (0.6 ± 0.07-fold versus controls, p = 0.005; Figure 3C). A concomitant enhancement of MAT2A transcripts also was noted in extracellular exosomes after the low-dose (0.25 Gy) exposure (1.7 ± 0.2-fold, p = 0.012) with non-significant levels reached after 2.5 Gy irradiation for this gene (0.47 ± 0.28-fold, p = 0.23; Figure 3C). Substantial up-regulation of PARTICLE expression in T47D following exposure to exosomes isolated from irradiated MDA-MB-361 (24 hr after 0.25 Gy) was noted (35 ± 3.2-fold, p = 0.006; Figure 3D), but was not evident for MAT2A (Figure 3E). In vitro exposure of whole blood to 2 Gy with exosomal isolation revealed that, although PARTICLE levels varied among individuals, increases ranged from 4.4 ± 0.38- to 22.3 ± 1.8-fold over control levels (p < 0.005; Figure 3F). Likewise, MAT2A mRNA also was detected in these exosomes with significantly elevated levels, ranging from 3 ± 0.09- to 4.3 ± 0.6-fold over controls for all exosomal extracts (n = 3; p < 0.05; Figure 3F). Increases in PARTICLE also were noted in plasma samples obtained from three of four post-radiation therapy patients (Patients A–C, p < 0.05; Figure 3G). Concomitant increases in MAT2A over controls also were revealed (A, 5 ± 0.9-fold, p = 0.014; B, 1.85 ± 0.11-fold, p = 0.002; C, 2.2 ± 0.5-fold, p = 0.01; D, 1.3 ± 0.23-fold, p = 0.01; Figure 3G).

PARTICLE Silencing Enhances MAT2A Transcription

Knockdown of PARTICLE (Figure S3) caused overexpression of MAT2A in sham-irradiated MDA-MB-361 cells (Figure 4A). PARTICLE expression was decreased (86.3%, p = 0.002) when MDA-MB-361 were transduced with lentivirus encoding the silencing small hairpin RNA (shRNA). A compared to empty vector alone (similar to the effects of Cisplatin exposure, Figure S4). At the same time, a reverse trend toward a 2.2-fold elevation was detected in MAT2A (p = 0.06). In cells transfected with lentivirus encoding shRNA_B, PARTICLE expression was again strongly decreased (94.2%, p = 0.00001) and again coincided with a concomitant 3.3-fold increase in MAT2A transcript expression (p = 0.001; Figure 4A).

Expression of MAT2A Is Enhanced in PARTICLE Knockdown Cells following Low-Dose Radiation Exposure

Knockdown of PARTICLE with lentivirus encoding shRNA_B resulted in an overshoot of MAT2A expression (7.7-fold increase, p = 0.0038) following low-dose (0.25 Gy) irradiation in MDA-MB-361 cells (Figure 4B). Essentially similar results were seen with
shRNA_A, resulting in MAT2A elevation (5-fold, p = 0.017; Figure 4B, right). Cells exposed to 2.5 Gy showed similar effects to sham-irradiated cells for PARTICLE and MAT2A transcription response (Figure 4C). PARTICLE expression decreased (62.4%, p = 0.017) in MDA-MB-361 cells infected with lentivirus encoding shRNA_A. This again correlated with a 0.5-fold significant increase in MAT2A (p < 0.05; Figure 4C, right). Slight improvement in PARTICLE knockdown was seen (71.6%, p = 0.0004) in cells infected with lentivirus encoding shRNA_B, coinciding with a greater, 2.5-fold, significant increase in MAT2A (p = 0.0029; Figure 4C, right). The increase in MAT2A expression was far greater following 0.25 Gy low-dose irradiation compared to 2.5 Gy exposure (5.2-fold higher, p < 0.005). These findings highlight the different transcriptional response profiles emanating from exposure to either low or medium irradiation dosage, and they emphasize the homeostatic repressive influence of PARTICLE on MAT2A expression.

**PARTICLE Represses MAT2A Promoter Activity**
Enhanced MAT2A promoter activity was noted in MDA-MB-361 with PARTICLE knockdown (Figure 4D, left). While non-infected (NT) cells showed expectedly negligible luciferase reporter expression, the transcriptional activity was significantly elevated in sham-irradiated cells and following 4 or 24 hr irradiation (0.25 or 2.5 Gy) donor cells. (Lane 1) RNA ladder (LD, Ambion); (lane 2) RNA from control (C); (lanes 3 and 4) Bru-labeled RNA isolated from recipient cells (300 and 3,000 bp lower and upper arrowheads). (Right) Northern blot shows PARTICLE detected in isolated Bru-labeled RNA in recipient cells exposed to Bru RNA labeled and irradiated (24 hr after 0.25 Gy) donor cells. (Lane 1) RNA ladder (as above); (lane 2) band showing PARTICLE (size 1,432 bp, arrowhead).

**Low-Dose Irradiation Promotes the Expression of the Catalytic Subunit of MAT and SAM Production**
As previous results revealed elevated MAT2A transcription within 4 hr of low-dose irradiation exposure, an investigation was undertaken to determine whether this would have an influence on MAT2A and SAM (Figure 5A). MAT2A in irradiated and sham-irradiated MDA-MB-361 was monitored over time (4–48 hr), revealing an increased concentration of the catalytic subunit of MAT 4 hr post-exposure to 0.25 Gy irradiation.
The rapid decrease in MAT2A protein levels by 24 hr post-irradiation exposure (p < 0.05; Figure 5B) also reflects the transcriptional expression profile (Figures 1 and 2). Such effects were not evident after exposure to 2.5 Gy, although a trend toward a decrease was apparent at the 24 hr time point, consistent with RNA levels. No alteration in MAT2A was found after 48 hr (Figure 5B), indicative of the transient response of this catalytic subunit specifically to low-dose irradiation exposure.

Intracellular levels of SAM varied from 12 ± 0.15 to 6.5 ± 1.5 nM/10^6 MDA-MB-361 when measured between 4 and 24 hr after sham irradiation (Figure 5C, left). While no effects were noted in MDA-MB-361 up to 48 hr after the 2.5 Gy dose of irradiation, SAM concentrations were twice as high at 4 hr after the low-dose (0.25 Gy) exposure, with levels returning to normal values by 24 hr (Figure 5C).

**PARTICLE Knockdown and Irradiation Have an Additive Effect on Intracellular and Extracellular SAM Production**

**PARTICLE** knockdown and concomitant elevation of MAT2A affected intracellular SAM levels in sham-irradiated cells. Significantly elevated SAM levels were noted for both the 4- and 24-hr time points, with concentrations 4- to 5-fold higher than transduced control cells (p < 0.005; Figure 5D, left). Following low-dose irradiation, a further substantial increase in intracellular SAM (80 ± 2.76 nM/10^6 cells) was noted in **PARTICLE** knockdown MDA-MB-361 (4-fold SAM increase over NT cells, p < 0.0005; Figure 5D, middle). This was only noticeable by 4 hr post-irradiation, particularly following low-dose exposure.

The concentration of extracellular SAM was assessed in the tissue culture medium conditioned by non-transduced or **PARTICLE** knockdown MDA-MB-361 plus or minus irradiation exposure. SAM concentrations in growth medium obtained from sham-irradiated cells varied from 9 ± 0.5 to 7 ± 0.12 nmol/l when collected at 4 to 48 hr after sham irradiation (Figure 5E, left). Significantly increased extracellular SAM levels were identified in the media of MDA-MB-361 exposed 4 hr earlier to low-dose (0.25 Gy) irradiation (p < 0.05; Figure 5E, middle). No significant alteration in extracellular SAM levels was found in media taken from cells 4–48 hr previously exposed to 2.5 Gy (Figure 5E, right).

SAM concentrations also were examined in the growth media of MDA-MB-361 in which **PARTICLE** had been depleted by lentiviral shRNA expression (Figure 5F). While no significant difference was found for extracellular SAM levels in sham-irradiated cells (Figure 5F, left), dramatically elevated SAM concentrations (24.7 ± 3 nmol/l) were identified in media from MDA-MB-361 (with **PARTICLE** knockdown) exposed 4 hr earlier to 0.25 Gy. SAM levels subsequently declined to values close to the normal range (6 ± 0.2 nmol/l) by 24 hr, remaining relatively unchanged up to 48 hr post-exposure (5 ± 0.7 nmol/l; Figure 5F, middle). A comparable trend was noted following 2.5 Gy exposure under similar culture conditions. Elevated SAM (13 ± 2.6 nmol/l,

**Figure 4.** **PARTICLE** Knockdown through Stable Lentiviral Transfection Causes MAT2A Overexpression and Promoter Over-activity

(A–C) Plots show fold change in **PARTICLE** (black histograms) and MAT2A (white histograms) expression relative to NT (non-Trans.) control levels (dotted line) in MDA-MB-361 that were sham irradiated (A) or exposed to 0.25 Gy (B) or 2.5 Gy (C). **PARTICLE** and MAT2A levels in MDA-MB-361 infected with lentivirus expressing non-specific (scrambled) control sequence (slight irradiation effect noted following 2.5 Gy) designed from shRNA_B region (red arrow [Figure S2], Scr. ctl.), pGP vector control (pGP ct.), shRNA_A (lenti_shRNA_A), or shRNA_B (lenti-shRNA_B) are shown. Data are represented as mean ± SEM with significant levels (p < 0.05) indicated where appropriate (asterisks). Note the presence of puromycin in the media did not have an effect on **PARTICLE** or MAT2A expression.

(D) Histogram shows Gaussia luciferase (GLuc) secretion normalized to secreted alkaline phosphatase (SEAP) indicative of MAT2A promoter activity in MDA-MB-361 (±**PARTICLE** knockdown [KD]). Tissue culture media were analyzed 4 and 24 hr following 0.25 Gy or 2.5 Gy cellular irradiation or sham irradiation. Data are represented as mean ± SEM. See also Figures S3 and S4.
p ≤ 0.05) was found only at the 4 hr time point. These findings highlight the consequences of PARTICLE suppression of MAT2A on SAM availability and the buffering of the radiation effect by control of SAM over-production.

**PARTICLE Controls the Methylation Status of the MAT2A Promoter CpG Island**

The MAT2A promoter contains a CpG island (annotated 108368) of 1,288 bp located on chromosome 2: 85765695–85766983 (NCBI *Homo sapiens* build number 37 version 2). The transcription initiation site for MAT2A resides within this region at chromosome 2: 85766100 orientated in a forward direction (NCBI refseq NM_005911). The sequence for PARTICLE also overlaps this CpG island from position chromosome 2: 85765818 for 123 bp, orientated in the antisense complementary direction (Figure 6A, top).

CpG island 108368 was almost totally unmethylated (>99.7% ± 0.2%) in sham-irradiated MDA-MB-361 cells (Figure 6A, bottom left). Following exposure to 0.25 or 2.5 Gy irradiation, the CpG island 108368 became progressively methylated over the next 48 hr. Differing levels of methylation were revealed following the two irradiation doses. After low-dose exposure, CpG island 108368 remained unmethylated up to 4 hr (>99.9% ± 0.01%), but became methylated by 24 hr (62% ± 3%) increasing to 85 ± 4% by 48 hr (Figure 6A, bottom middle). Following 2.5 Gy irradiation, methylation occurred earlier (16% ± 2%), appearing by 4 hr (Figure 6A, bottom right). While methylation status increased to 49% ± 2% and 59% ± 3% for 24 and 48 hr, respectively, these levels were lower than those evoked after low-dose (0.25 Gy) irradiation exposure (Figure 6A, bottom right).

In MDA-MB-361 with PARTICLE knockdown (Figure 6B, top), the CpG island 108368 remained unmethylated under sham-irradiated conditions, similar to that described for cells with normal PARTICLE expression (Figure 6B, bottom left). However, following low-dose (0.25 Gy) irradiation, the CpG island 108368 stayed un-methylated (>99.5%) in the absence of PARTICLE up to 4 hr, similar to non-transduced cells, and remained un-methylated up to 24 hr (>99%); methylation began only at 48 hr, reaching 57% ± 3% (28% ± 2% lower level than in non-transduced cells; Figure 6B, bottom middle). After 2.5 Gy exposure,
PARTICLE were predicted to bind upstream of a CpG island identified in silico, where triplex-forming oligonucleotides within the stream of the CpG island 108368 (described above) was identified. A region (chromosome 2: 85765239–85765251) 456 bp upstream of the CpG island 24 hr after low-dose irradiation, a time when PARTICLE expression is elevated and MAT2A transcription is suppressed.

**PARTICLE and MAT2A Form a Triplex Structure**

A region (chromosome 2: 85765239–85765251) 456 bp upstream of the CpG island 108368 (described above) was identified in silico, where triplex-forming oligonucleotides within PARTICLE were predicted to bind upstream of a CpG island that is located in the promoter of MAT2A (Figures 6A [green triangle] and 6C). Computational modeling indicated that triplex formation between PARTICLE and this site was highly probable (score rate = 12, error rate = 0.007) (Buske et al., 2012). Surface plasmon resonance (SPR) was used to confirm that PARTICLE indeed interacts with MAT2A via a triplex mechanism. A panel of RNA oligonucleotides covering the predicted triplex-forming motif (TFO-RNA) within PARTICLE was tested (Table S3). A significant spectral shift occurred only with the TFO-RNA (i.e., 1.1 ± 0.05 nm, p = 0.0000377, t stat. = 115.2) (Figure 6E). Moreover, addition of the TFO-RNA to the receptor MAT2A without prior formation of the DNA duplex (Figure 6E) provided negligible signal (i.e., 0.09 ± 0.05 nm, p > 0.05). The resulting RNA-DNA triplex may govern the CpG island methylation to instigate transcriptional suppression of MAT2A.

**PARTICLE Associates with Myb, G9a Lysine Methyltransferase, and the Polycomb Repressor Complex Subunit SUZ12**

Chromatin immunoprecipitation (ChIP) of protein interaction partners with PARTICLE was performed using cross-linked chromatin isolated from irradiated MDA-MB-361 (24 hr after 0.25 Gy). Differential occupancy analysis of qPCR data revealed that the increase of PARTICLE at the Myb-, G9a lysine methyltransferase-, and SUZ12-binding sites was 1.3 ± 0.1, 4.1 ± 0.3, and 9.2 ± 0.7-fold, respectively (Figure 7A). Pull-down of in vitro transcribed biotinylated PARTICLE confirmed the association with these nuclear proteins (isolated from MDA-MB-361 24 hr after 0.25 Gy) (Figure 7B). An electrophoretic mobility shift was identified in the presence of PARTICLE (±biotinylation) and the polycomb repressor complex subunit SUZ12 peptide. Binding specificity was demonstrated by increasing concentrations of unlabeled PARTICLE (up to 1,000-fold molar excess) that significantly diminished the interaction with the MAT2A duplex monolayer (W-MAT2A, blue and brown combs) previously formed on the gold SPR chip.

(E) Representative sensorgram showing the specificity of triplex detection by virtue of the differential between the signal of the TFO-RNA (red line) versus the signal provided by the other RNAs tested within the context of the PARTICLE sequence. This TFO-RNA signal is also specific in that it requires prior formation of the duplex. Thus, only a minimal spectral shift is generated by the TFO-RNA oligonucleotide injected over the receptor without previous formation of the duplex.
CpG island methylation. Moreover, cytosolic PARTICLE levels become dramatically increased following cellular irradiation amid juxtaposition with MAT2A transcripts, which most likely precedes their extracellular export.

MAT is a critical cellular enzyme that catalyzes the formation of SAM, the principal methyl donor, with MAT2A encoding the catalytic (α2) subunit of the widely distributed MAT isozyme (MATII) (Alvarez et al., 1993; Kotb et al., 1997). PARTICLE is transcribed in an antisense orientation to the forward plus strand from the MAT2A promoter. Our findings show that MAT2A transcription is activated much earlier than PARTICLE, within 4 hr following low-dose radiation exposure. This is accompanied by concomitant increases in translation and production of both intracellular and extracellular SAM. In the artificial absence of PARTICLE, transcriptional repression of MAT2A becomes diminished, resulting in elevated intracellular SAM levels, including an overshoot in irradiated PARTICLE-deficient cells. These results imply that PARTICLE directly influences the methylation cycle responsible for the conversion of methionine via SAM and s-adenosylhomocysteine (SAH) into homocysteine. As SAM is the methyl donor for multiple detoxifying methylation reactions, it is not surprising that it becomes elevated following genotoxic exposure, such as irradiation, and that it must be removed once the radiation response is completed.

Accumulating evidence suggests that long antisense ncRNAs function as epigenetic regulators of balanced transcription (Morris, 2009). This may be the principal pathway evoked by PARTICLE for modulating MAT2A expression according to selective pressures, i.e., irradiation, placed on the cell. PARTICLE might serve to keep in check excess availability of methyl groups required for increased DNA damage repair activity following radiation exposure. This response is most likely instigated by the upregulation of PARTICLE operating as a silencer through triplex (RNA:DNA:DNA) formation within the CpG island shore region (456 bp upstream of the CpG island) of the MAT2A promoter. While a similar triplex riboswitch has been identified upstream of MAT2A across a taxonomically wide range of bacteria, no mammalian evidence exists to date (Conrad, 2014). It can be inferred that human PARTICLE offers such a platform for controlling SAM production.

PARTICLE serves as a protein-binding platform, thereby enabling cis regulation of MAT2A transcription. PARTICLE interacts with the myeloblastosis (Myb) proto-oncogene transcription factor that previously has been implicated in MAT2A upregulation in human hepatocarcinogenic carcinomas and tumorigenesis (Yang et al., 2001). The lysine DNA methyltransferase (G9a) methylates H3K9 and H3K27 to silence transcription (Tachibana et al., 2001), and we also found G9a associating with PARTICLE in this study. The PRC2 complex catalyzes trimethylation of H3K27 and mediates transcriptional repression (Surface et al., 2010). A number of IncRNAs have been shown to interact with PRC2 to regulate target gene expression (Wang and Chang, 2011). Moreover, this report shows direct interaction with PRC2 and a radiation-responsive triplex-forming IncRNA, PARTICLE. Interestingly, a recent report demonstrated that G9a and PRC2 interact and co-localize genome-wide (Mozzetta et al., 2014). It could be proposed that PARTICLE binds SUZ12 and competes for PRC2 binding at target sites following

**DISCUSSION**

This study introduces PARTICLE, an IncRNA tuner of cellular methylation following radiation exposure. An active silencing mechanism of MAT2A transcription by nuclear PARTICLE is observed, consisting of a physical crosstalk between chromosomal DNA and IncRNA leading to DNA duplex-IncRNA triplex formation, gene-silencing complex interaction, and promoter

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**Figure 7. PARTICLE Associates with Myb, Lysine Methyltransferase G9a, and the Polycomb Repressor Complex Subunit SUZ12**

(A) Histograms show fold increase in PARTICLE from qPCR expression analysis of chromatin immunoprecipitated with various antibodies, including anti-Myb, anti-G9a, and anti-SUZ12 (see Supplemental Experimental Procedures for details).

(B) Representative western blots of eluted fractions (streptavidin bead purified) probed with anti-Myb, anti-G9a, and anti-SUZ12. Biotinylated in vitro transcribed PARTICLE was used to pull down Myb, G9a, and SUZ12 from MDA-MB-361 (24 hr after 0.25 Gy irradiated) nuclear extracts. (Lane 1) Protein detection in nuclear lysate input; (lane 2) absence of signal in the absence of biotinylated PARTICLE; (lane 3) Myb, G9a, and SUZ12 detected in nuclear extracts following in vitro transcribed biotinylated PARTICLE pull-down and streptavidin bead elution.

(C) Representative nucleotide retardation gel (6%) of electrophoretic mobility shift assay involving binding reactions containing biotinylated PARTICLE (b-PART., 10 nM), a SUZ12 peptide (2.5 μM), and increasing concentrations of unlabeled PARTICLE (20 nM to 20 μM) in the presence of 6.25 mM KCl is shown.

(D) Hill slope depicts the percentage of biotinylated PARTICLE (b-PARTICLE) bound to the SUZ12 peptide in the presence of molar excess of unlabeled PARTICLE. Data are represented as mean ± SEM.

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amount of SUZ12-bound biotinylated PARTICLE (Figures 7C and 7D). These data show that PARTICLE interacts directly with PRC2 via the SUZ12 subunit. It suggests that PARTICLE functions, in part with epigenetic silencing complexes, to curb possible overexpression of MAT2A (and hence over-production of SAM and DNA methylation) following exposure to ionizing radiation.
Irradiation exposure. Alternatively, PARTICLE may aid in the recruitment of PRC2 to repress MAT2A, as loss of PARTICLE might fail to silence overactive MAT2A transcription in response to irradiation insult. Further studies will be necessary to fully elucidate this function, although our work suggest that PARTICLE mediates MAT2A suppression via potential epigenetic regulatory mechanisms.

It is recognized that location within the cell is an important determinant in understanding the functional roles of lncRNAs (van Heesch et al., 2014), suggesting they may play different roles depending on their sub-cellular compartment context. We have identified such a process as a possible second role for PARTICLE in response to low-dose irradiation. While nuclear PARTICLE actively represses MAT2A, cytosolic PARTICLE becomes enriched and appears to associate with cytosolic MAT2A transcripts, predominantly after irradiation exposure (4 hr). This could serve as an additional mechanism of MAT2A repression, with PARTICLE regulating its availability for translation.

While it is generally accepted that microRNA and mRNA can be transferred between mammalian cells by an exosome-based transport mechanism (Lässer et al., 2011), this study has identified an lncRNA in exosomes generated from in vitro, ex vivo, and in vivo irradiation. It is tempting to speculate that the detection of PARTICLE in plasma exosomes indicates that they serve as vectors for genetic communication between cells in distant organs. Although this suggest that PARTICLE may have the capacity to affect the phenotype of recipient cells, to date no mechanistic basis is known (Kadhim et al., 2013).

The roles of IncRNA in the DNA-damage/repair response are only beginning to be unraveled. Importantly, the inverse dose response exhibited by PARTICLE to radiation challenges the perception that all events following exposure show a linear escalation with increased doses. While their crucial existence is now undisputed, no doubt the long non-coding genome will continue to surprise and reveal unexpected layers of cellular regulatory complexity.

**EXPERIMENTAL PROCEDURES**

**Human IncRNA Microarray Analysis**

Differentially expressed IncRNAs were identified through fold change and volcano plot filtering between samples. Genomic location was used to select for regulated intragenic IncRNAs.

**Propagation and Maintenance of Cell Lines**

MDA-MB-361, HEK293, U2OS, T47D, and HUVEC were propagated as described in the Supplemental Experimental Procedures. All cells were genotyped to confirm identity.

**Irradiation**

All irradiations were performed using a closed HWM-D 2000 unit (Siemens, 10-cm height and 33-cm circumference) delivering Cesium 137 gamma rays at a dose rate of 0.0082 Gy/s. Calibration was performed by the Research Unit of Medical Radiation Physics and Diagnostics (Helmholtz Zentrum Munich).

**Ethical Approval**

Samples from human subjects were obtained with informed consent. The Ethikkommission der Medizinischen Fakultät der Ludwig-Maximilian-Universität München, Germany, approved the study.

**Isolation of Exosomes from Human Blood Plasma**

Exosomes were isolated from human plasma of healthy donors by differential centrifugation as described in the Supplemental Experimental Procedures and as previously reported (Théry et al., 2006).

**Head and Neck Squamous Cell Carcinoma Patients**

The details of patient treatment and plasma isolation have been reported previously (Summerer et al., 2013).

**Triplex SPR Assay**

All SPR experiments were performed using a previously reported (Carrascosa et al., 2014; Sina et al., 2014) custom-made SPR platform.

**ChIP**

Experiments were performed using an EpiTect ChIP kit (QIAGEN, 334471) per the manufacturer’s instructions. Immunoprecipitation involved the following ChIP-grade antibodies: anti-Myb, anti-KMT1C/G9a, anti-NF-κB p105/p50, anti-p53, anti-SUZ12, anti-histone H2B (positive control), and normal IgG (negative control). See the Supplemental Experimental Procedures for further details.

**RNA Pull-Down**

See the Supplemental Experimental Procedures for experimental details.

**Electrophoretic Mobility Shift Assay**

Experiments were carried out per the manufacturer’s instructions (Thermo Scientific, 2015B) with the inclusion of 6.25 mM KCl in the binding reactions and electrophoresis through a 6% pre-cast nucleotide retardation gel.

**Statistical Analysis**

Values in the text are expressed as the mean ± SEM and n refers to the number of independent data. Triplex t statistic was tested using hypothesis parameters (HA: µT−µG > 0) and Origin 7 software. Differences between means were tested using the Student’s t test with p values < 0.05 taken to indicate statistical significance.

**ACCESSION NUMBERS**

The Gene Expression Omnibus (GEO) accession number for the human IncRNA microarray data reported in this paper is GSE67008.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2015.03.043.

**AUTHOR CONTRIBUTIONS**

V.B.O. designed and performed experiments, interpreted results, and wrote the paper. S.V.O. performed microscopy, graphics, and data analysis. L.G.C. performed in vitro triplex experiments. F.A.B. designed and implemented the Triplexator software. V.R. interpreted data and corrected the manuscript. M.N. and S.M. provided valuable samples, interpreted data, and corrected the manuscript. M.T. supervised the triplex experiments. N.A. performed lentiviral preparations, interpreted results, corrected the manuscript, and supervised the project. M.J.A. designed the project, contributed to the experimental strategies, interpreted data, corrected the manuscript, coordinated the project, and directed the research.

**ACKNOWLEDGMENTS**

The authors thank Mr. Michael Salomon (Sirion Biotech GmbH, Germany) for helpful advice on shRNA interference and secondary structure prediction analysis. This work was supported by the EURATOM Fission, European Commission 7th Framework Programme, Dark.Risk project (contract number 323216).
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