MicroRNA Analysis of ATM-Deficient Cells Indicate PTEN and CCDN1 as Potential Biomarkers of Radiation Response

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microRNA analysis of ATM-deficient cells indicate PTEN and CCDN1 as potential biomarkers of radiation response

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Genetic and epigenetic profile changes associated with individual radiation sensitivity are well documented and have led to an increase in our understanding of the mechanisms of the radiation-induced DNA damage response. However, the quest to identify reliable biomarkers of individual radiation sensitivity is on-going. Herein, we report a multi-biomarker approach using traditional cytogenetic biomarkers, DNA damage biomarkers and transcriptional microRNA (miR) biomarkers coupled with their potential gene targets to identify radiosensitivity in ATM (Ataxia-Telangectasia Mutated)-deficient lymphoblastoid cell lines (LCL) and ATM proficient cell lines that were used as controls.

Cells were irradiated with 0.05 Gy and 0.5 Gy using an Elekta Precise linac, with sham-irradiated cells as controls. At 1 hour post irradiation, cells were fixed for γH2AX analysis as a measurement of DNA damage, and cytogenetic analysis using the G2 chromosomal sensitivity assay, G-Banding and FISH techniques. RNA was also isolated for genetic profiling by microRNA (miR) and RT-PCR analysis. A panel of 752 miR were analysed, and potential target genes phosphatase and tensin homolog (PTEN) and cyclin D1 (CCND1) measured.

The cytogenetic assays revealed that although the control cell line had functional cell cycle checkpoints, the radiosensitivity of the control and AT cell lines were similar. Analysis of DNA damage in all cell lines, including an additional control cell line, showed elevated γH2AX levels.
for only one A-T cell line. Of the 752 miR panel analysed, 8 miR were found to be up-regulated, with 6 miR down-regulated in the AT cells compared to the control. Up-regulated miR-152-3p, miR-24-5p and miR-92-15p and all down-regulated miR were indicated as modulators of PTEN and CCND1. Further measurement of both genes validated their potential role as radiation response biomarkers. The multi-biomarker approach not only revealed potential candidates for radiation response but also additional mechanistic insights of response in AT deficient cells.

**INTRODUCTION**

In the last decade, the definition and classification of ionizing radiation biomarkers have been reported through several European Union Framework 7 multidisciplinary consortia such as Multibiodose (2010-2013), RENEB (Realizing the European Network in Biodosimetry (2012-2015)) and DoReMi (Low dose Research towards multidisciplinary Integration (2010-2015)) with the multipurpose use of biomarkers for epidemiological and biodosimetry investigations ([1-4]). These include biomarkers of low dose exposure and biological response, individual susceptibility and early detection of a radiation-induced health effect, of which considerations to the characteristics of a good biomarker and the useful in vitro approaches have been made.

Although the DoReMi project was completed in 2015, research has continued under Melodi (Multidisciplinary European Low Dose Initiative (5)) and the DoReMi multidisciplinary report (4) was later updated to include novel radiation biomarkers emerging from technical advances in metabolomics and transcriptomics, and to critique the current status of biomarkers ([6]). A roadmap for the development of biomarkers from discovery to implementation was presented for biomarkers of low dose exposure and early or late
radiation effects. The authors highlighted that the majority of potential biomarkers are in the development stage with only one biomarker that has progressed to the final stages of development with IR specific mRNA transcript profiles for FDXR. This gene has been reported in many proposed gene signature panels due to dose-dependent induction in different cell and tissue types (7-10). Furthermore, inter-comparison laboratory or biodosimetry studies have demonstrated that both single genes and gene panels can be used to estimate exposure of samples with the same accuracy and sensitivity of established and traditional cytogenetic assays (11-12).

The DNA damage response (DDR) pathways are potential targets for transcriptional biomarkers of cancer susceptibility and radiation exposure; in particular the ATM/chk2/p53 pathway, which responds to radiation-induced double strand breaks (DSB) leading to cell cycle arrest or DNA repair. The DSB are sensed by the MRN complex (MRE11-Rad50-NBS-1) leading to ATM activation, phosphorylation of serine 139 of γH2AX and extension around the DSB, initiating repair protein assembly (13-14). Consequently, γH2AX has been used as a biomarker of DNA damage and repair and for predicting radiosensitivity in individuals (15-17) and applied to a wide range of established cell lines, primary cell cultures and peripheral blood lymphocytes as well as 2-dimensional tissue models and tissue sections as reviewed by Rothkamm et al (18). The role of ATM, a PI3K-like kinase that is phosphorylated at specific serine/threonine sites when activated, is central to this pathway. Deficiencies in the ATM gene lead to phenotypic elevated radiosensitivity observed in clinical conditions such as Ataxia Telangiectasia (AT) and AT-like disorders (ATLD) (19-21). After DSB are sensed, the cell cycle must be halted to allow sufficient time for DNA repair processes, facilitated through ATM-activated Chk2. This leads to p53-mediated inhibition of cyclins and cyclin-dependent kinases, such as Cyclin D1 (CCDN1) and CDK4/6 at the G1 cell cycle checkpoint (22). Failure to
undergo DNA repair may result in permanent cell cycle arrest, enhanced apoptosis or cellular senescence. The PI3K/Akt pathway is also involved in the survival of cells after IR-induced DNA damage, through overriding the G2/M cell cycle arrest mechanism; conversely inhibition of PI3K or Akt, for example through the tumour suppressor PTEN, induces cell apoptosis and therefore elevates cellular radiosensitivity (23-25).

Further transcriptomic analyses have shown that microRNA (miR) are promising biomarkers of radiation oncology (26). They are small, non-coding RNA molecules of 19-22 nucleotides that regulate more than 50% of cell protein coding genes and regulate important processes of the DNA damage response such as DNA repair, cell cycle control and apoptosis. It has previously been shown that important genes of these processes (such as CDKN1, SESN1, ATF3, MDM2, PUMA and GADD45A) were upregulated in stimulated T cells in response to IR with a significant dose- and time-dependent modification of miR expression (specifically miR-34-5p and miR-182-5p) (27-28).

Given the current published evidence associating the ATM/Chk2/P53 pathway with elevated radiosensitivity and potentially regulated by miR, normal and AT radiosensitive lymphoblastoid cell lines were used to measure IR-induced DNA damage using the classic cytogenetic and DNA damage biomarkers followed by miR screening and identification of gene targets in a multi-biomarker approach. All biomarkers selected for this study were based on the DoReMi (Low dose Research towards multidisciplinary Integration) multidisciplinary biomarker reports by Pernot et al (4), and Hall et al (6), and the recent report which reviews the progress made in low dose health risk research by the DoReMi consortium (29).

MATERIALS AND METHODS
**Cell Lines and culture conditions**

Epstein-Barr immortalised lymphoblastoid cell lines (LCLs) coded; C1, 2139, AT2Bi and AT3Bi were used for this study. C1 and 2139 cell lines were derived from healthy donors and kindly gifted by the Queensland Institute of Medical Research, Australia and the Institut Curie, Paris, respectively. The AT2Bi and AT3Bi cell lines were derived from clinically established Ataxia-Telangiectasia patients and kindly gifted from the College of Medical and Dental Sciences, University of Birmingham, UK. Both AT2Bi and AT3Bi are known to have defective Ataxia-Telangiectasia- mutated (ATM) protein causing the typical clinical and cellular manifestations of AT including heightened radiosensitivity (30). C1, 2139, AT2Bi and AT3Bi lymphoblast cells were cultured in RPMI 1640 medium (Sigma Aldrich, Wexford, Ireland) supplemented with 12.5% FBS and 1% L-Glutamine (Sigma Aldrich), at 37 °C and 5 % CO₂. All cell lines were seeded at a density of 2x10⁵/ml and passaged once a density of 1x10⁶/ml cells had been reached. Cells were seeded 18 hours prior to irradiation, a T25 flasks at a density of 1x10⁶cells/ml (G2 chromosomal radiosensitivity assay), 2x10⁴cells in total (growth curves), or 2x10⁵/ml (γH2AX and molecular experiments) at a final volume of 5 ml per T25 flask (Sarstedt, Numbrecht, Germany).

**Irradiation Conditions**

Cells were irradiated using a 6MV photon beam produced by an Elekta Precise linear accelerator (LINAC) at St. Luke’s Hospital, Dublin, operating at a nominal dose rate of 6Gy/min. The LINAC was calibrated in accordance with the 1990 IPSM code of practice by the Medical Physics Department at St. Luke’s Hospital (31), with 100 Monitor Units (MU, a measure of ‘beam on’ time) delivered a dose of 1Gy at 1.4 cm deep in water positioned 100 cm from the source for a 10 X 10 cm² field. To achieve a uniform irradiation of flasks, the
irradiation conditions were altered from those at calibration. A 30 x 35 cm² field was used to deliver each dose. The flasks were also positioned 10 cm deep in a water equivalent phantom 90 cm from the source in which 100MU delivers a dose of 0.812Gy at 10 cm deep in water for a 10 x 10 cm² field. The number of MU required to deliver each of the doses outlined were corrected for the different scatter conditions present with the larger field size (30 x 35 cm²). Therefore, a correction factor of 1.1372 was applied, which is the ratio of the field area of a large field to a smaller one. Thus, at 90 cm from the source, 100MU delivers a dose of 0.9234 Gy (0.812 X 1.1372), and therefore the delivery of 0.05Gy required 6 MU and 0.5Gy required 55 MU (MU were rounded up to the nearest whole number as partial MU could not be delivered on the LINAC). The calculated doses were verified using Gafchromic EBT3 film (Ashland Inc., Bridgewater, NJ, USA) and the film was calibrated against a Farmer type ionization chamber using the triple channel dosimetry method (31). The film was scanned using the single scan protocol (32) on an Epson Expression 10000 XL scanner with the recommended scanning resolution of 72 dpi in a 48-bit RGB format (31, 33-34). Glass was placed over the calibration and test film during scanning to minimize ringing artifacts. The film was analyzed using FilmQA Pro (Ashland Inc., Bridgewater, NJ, USA).

**Cell Growth Assay**

To determine the effect of radiation on the growth potential of the cells, flasks were seeded and irradiated as described above. At 5-7 days post- irradiation, cells were isolated and counted in duplicate using a Coulter cell counter (Beckman Coulter, Co Clare, Ireland). Total cell numbers were calculated and analysed with reference to sham-irradiated controls.
**Gamma-H2AX analysis by Flow Cytometry**

DNA damage was determined by γ-H2AX analysis and measured by flow cytometry. Cells were fixed at 1-hour post irradiation in 2 % paraformaldehyde and stored in 70 % ethanol at -20 °C. To stain, cells were permeabilised using 0.25 % Triton X, followed by blocking with a 4 % FBS solution in PBS for 30 minutes. A primary antibody solution (anti-phospho-histone H2A.X (Ser139), clone JBW301, 1:500; Merck Millipore, Darmstadt, Germany) was added and incubated overnight at 4 °C, followed by a 1-hour incubation with the secondary antibody (F(ab’)- Goat anti-Mouse IgG (H+L), Alexa Fluor-488, 1:200; Thermo Fisher, Carlsbad, CA, USA) at room temperature. Cells were washed, counterstained with 1 % propidium iodide solution and analysed on an Accuri C6 flow cytometer (BD, Oxford, UK). The mean fluorescence of 10,000 cells was calculated using the Accuri C6 Sampler software, with cells stained only with the secondary antibody acting as a negative control for each sample.

**G2 chromosomal radiosensitivity assay**

The G2 Chromosomal radiosensitivity assay as previously reported for whole blood lymphocytes (35-37), was applied to all 2139, AT2Bi and AT3Bi cells to measure radiation-induced cell cycle checkpoint response by mitotic indices and G2 chromosomal radiosensitivity. The mitotic index (MI) was calculated by counting the ratio of cells in metaphase to all cells on the slide up to 1000 cells in total for each dose (0 Gy and 0.5 Gy) and cell line. Radiation-induced mitotic inhibition (RIMI) was calculated by subtracting the 0.5 Gy MI from the 0 Gy MI. A G2 radiosensitivity score was assigned to each of the cell lines and irradiation dose by calculating the total number of chromosomal aberrations per 100 metaphases scored for each cell line and dose. A radiation-induced G2 score (RIG2) was
calculated by subtracting the spontaneous aberrations in the G2 score at 0 Gy from those recorded at 0.5 Gy.

**Cytogenetic G-Banding and karyotyping**

Cytogenetic preparations were made from irradiated 2139 and AT (AT2Bi and AT3Bi) LCL according to the G2 chromosomal radiosensitivity assay. For G-Banding, the metaphase spreads on glass slides were covered with 30% hydrogen peroxide solution for one minute followed by a wash with 0.9% NaCl solution. The metaphase preparations were then placed in trypsin solution for 2 mins, washed with Gurr buffer (pH 8) and then stained in 1 ml of Leishmann: Gurr buffer (1:2) solution for 1 min. The slides were washed with Gurr buffer, then distilled water and dried before they were mounted with a coverslip using DPX. Each slide was evaluated under the microscope set up for bright-field use, noting conditions of under or over banding or staining. Twenty five metaphases were karyotyped under the microscope and analysed for chromosomal aberrations.

**Fluorescent In Situ Hybridisation (FISH)**

Cytogenetic preparations (metaphase spreads as above) obtained from radiation-exposed 2139, AT2Bi and AT3Bi LCL were soaked in sodium chloride and sodium citrate buffer (SCC) for 2mins at 37°C, before being applied to/ treated with protease solution for 30-40 seconds at 37°C. Slides were then washed in 1xPBS, dehydrated in an ethanol series (70%, 85% and 100%) for 2 minutes each at RT and air dried before hybridisation. Hybridisation FISH probes were used to identify deletions or rearrangements in ATM-TP53 particularly for the AT cells (AT2Bi and AT3Bi). Probes for ATM-TP53 were used to confirm the presence of ATM or TP53 gene in all LCL. Conditions such as B-cell chronic lymphocytic leukaemia (B-CLL), a malignancy often associated with Ataxia-Telangectasia has shown deletions in the genes of ATM (38, 39)
and P53 (40). Probes were mixed according to the manufacturer’s instructions and the required amount was added to each slide. Slides were transferred to a Hybrite machine with the selected Hybridisation program of 75°C for 2 min and 37°C for 20 hours. When hybridised samples were removed, the slides were immersed in wash solution (0.4xSSC/0.3% NP 40) for 2 minutes and then transferred into a solution of 2xSSC/0.1% NP40 for a minimum of 1 minute. DAPI (20μl) was added as a counterstain and slides were mounted in coverslips. For FISH microscopy, 100 Interphase cells were recorded.

**MicroRNA expression**

An expression panel of 752 miR was performed on 2139, AT2Bi and AT3Bi cell lines (Exiqon, Vedbaek, Denmark), in accordance with company protocols. Briefly, RNA (50ng) was reverse transcribed and cDNA assayed in 10μl PCR reactions (miRCURY LNA™ universal RT microRNA PCR, Polyadenylation and cDNA synthesis kit, ExiLENT SYBR® Green master mix). The amplification was performed in a LightCycler® 480 Real Time PCR System (Roche) in 384 well plates. Melting curve and Cq values were analysed using Roche LC software. Cq values were calculated as the second derivative, with values greater than 37 omitted from further analysis. All data was normalized to the average of assays detected in all samples (average – assay Cq).

**Gene Expression**

Irradiated LCL were analysed for selected PTEN and CCND1 gene expression by Real Time PCR (RT-PCR). RNA was extracted from cells using the phenol-chloroform method and concentration measured using the Nanodrop (Maestrogen, Las Vegas, NV, USA). CDNA was synthesised using the q-script cDNA kit (Quanta Bio, Beverly, MA, USA), according to manufacturer’s instructions. Primers for Tubulin, PTEN and CCND1 were designed (Table 1) and synthesised (Sigma Aldrich), and reactions were performed in duplicate in 96 well plates.
Each reaction was composed of 10 µl SYBR Green with low ROX, (Kapa Biosystems, London, UK), 1 µl of forward and reverse primers, 6 µl PCR grade water, and 2 µl cDNA. Non template controls replaced cDNA with 2 µl PCR grade water. Reactions were run for 45 cycles on AB 7500 fast PCR cycler (Applied Biosystems).

Table 1 Forward and reverse primer sequences for housekeeping gene Tubulin, and for targets PTEN and CCND1

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward Sequence</th>
<th>Reverse Sequence</th>
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<tr>
<td>Tubulin</td>
<td>5’GCTTCTTGGTTTCCACAGC'3</td>
<td>3’CTCCAGCTTGGACTTCTTGC'5</td>
</tr>
<tr>
<td>PTEN</td>
<td>AGACAAATTCCCAGCTTACTTCTGC</td>
<td>ACCAGGTGCTTCATAGAGTAGG</td>
</tr>
<tr>
<td>CCND1</td>
<td>GACAGGTCACATCAGAAGAGC</td>
<td>CCTTCAGAGTAATTTGCCCAGG</td>
</tr>
</tbody>
</table>

Statistical Analysis

All statistical analysis was performed using Microsoft Excel, versions 2010-2016. Mean and standard deviations were calculated, and significance was determined using paired or unpaired t-tests of each radiation dose relative to its 0 Gy control, for each individual cell line, as appropriate.

RESULTS

Cell Growth assay for monitoring cellular viability

All cell lines were irradiated to 0.05 and 0.5 Gy and cultured for 5 days to measure growth potential. Percentage growth was calculated relative to the sham-irradiated control after 5
days in culture, and counted using a Coulter Counter and displayed in Figure 1(A). After 5 days in culture the control 2139 cells indicated a linear dose response for each low dose (0.05 and 0.5Gy) compared to the 0Gy control (Figure 1A). Similarly the AT cells (AT2Bi and AT3Bi) indicated a dose response for 0.5Gy but not 0.05Gy. This was expected because we previously reported differential molecular mechanisms of Apoptosis for 0.05Gy compared to 0.5Gy between 1hr and 24hr direct irradiation (41). The additional control cell line C1 did not show a radiation dose response comparative to the 2139 control cells.

γH2AX Biomarker of DNA damage response

All cell lines were irradiated to 0.05 and 0.5 Gy and fixed for Gamma-H2AX (γH2AX) analysis through flow cytometry as shown in Figure 1 (B). % positive cells were calculated, and normalised to the sham-irradiated control of each cell line. Since the cytogenetic biomarker of radiosensitivity (G2 chromosomal radiosensitivity) did not discriminate G2 radiosensitivity between the control 2139 and AT cells (AT2Bi and AT3Bi), the γH2AX assay was employed to measure double strand breaks (DSBs) induced by radiation in all cells. An additional control cell (C1) with functional ATM similar to 2139 was also analysed. Fluorescent foci are equal to the number of DSB induced by IR. Figure 1B presents γH2AX positive cells in the 4 LCLs at 1 hour post-irradiation. A modest increase in γH2AX positive cells was evident in the AT3Bi cell line to 1.5 fold of the 0 Gy control, however this was not significant (p>0.05). Irradiation of the 2139 and AT2Bi cell line decreased γH2AX levels below that of the sham-irradiated cells, however this was not significant (p>0.1). There was no dose dependence of response in any cell line tested (Figure 1B). This assay was also performed at later timepoints with no observable trends between the cell lines and doses (data not shown).
Figure 1: Control (C1 and 2139) and AT (AT2Bi and AT3Bi) LCLs exposed to 0Gy, 0.05Gy and 0.5Gy IR for measuring (A) Cell growth and (B) γH2AX as a biomarker for DNA double strand breaks induced by IR. Data shown are representative of 4 independent experiments, mean +/- SD

Cytogenetic Biomarkers of radiation response
The G2 chromosomal radiosensitivity assay was used as cytogenetic biomarker of low-dose radiation-induced effects in the control 2139 and AT (AT2Bi and AT3Bi) lymphoblastoid cell lines. Assessment of mitotic indices (MI) through the G2 chromosomal radiosensitivity assay is a good indicator of cell cycle checkpoint response to ionising radiation, whereby radiation-induced mitotic inhibition (RIMI) is the calculated difference between the 0.5 Gy and 0 Gy MI. The normal expected MI for the G2 chromosomal radiosensitivity varies between 2-5%, whereas the RIMI can be varied depending on cellular response to IR. All cell lines presented MI within the expected ranges for 0 Gy as presented in Figure 2A, however RIMI was more pronounced in 2139 (1.2) compared to AT2Bi (-0.3) and AT3Bi (0.6). This indicated that the control cells 2139 had superior cell cycle checkpoint efficacy compared to the AT cells, probably due to functional ATM. All cell lines presented elevated G2 chromosomal aberrations when irradiated to 0.5 Gy compared to their non-irradiated counterpart as presented in Figure 2B. Interestingly, the control 2139 cell line had similar radiation-induced G2 chromosomal radiosensitivity RIG2 (203 aberrations/100 metaphases) as the two AT cell lines AT2Bi and AT3Bi (134 and 183 aberrations/100 metaphases respectively), which indicated that although checkpoint response by MI appeared to be functional compared to the AT cells, radiation-induced chromosomal damage was similar to the AT cells. This finding merited further cytogenetic investigation, performed in collaboration with the Genetics Department, Our Lady’s Children’s Hospital, Crumlin, Dublin. Cytogenetic karyotyping using the G-Banding Technique was performed on the 2139 and AT cells (AT2Bi and AT3Bi) and followed up with Fluorescent In-Situ Hybridisation (FISH) using an ATM/TP53 probe. The cytogenetic analysis on 2139 cells surprisingly showed a loss of a sex chromosome in all of the cells analysed (Figure 3), with no other single cell or recurrent aberrations detected. The loss of a sex chromosome is associated with the constitutional diagnosis of Turner’s syndrome in
females. FISH analysis using ATM (11q22)/TP53 (17p13.1) probe set presented two copies of each ATM and P53 in each cell line with no detectable deletions, numerical aberrations or translocations at these loci in the 100 Interphase cells analysed.

Figure 2: Control (2139) and AT (AT2Bi and AT3Bi) LCLs exposed to 0Gy (grey bars) and 0.5Gy (black bars) in G2 chromosomal radiosensitivity assay for (A) Mitotic Index and (B) G2 score. Data shown are representative of 3 independent experiments, mean +/- SD, **p<0.01, ***p<0.005.
MicroRNA biomarkers of radiation response

MicroRNA (miR) analysis was performed on the control (2139) and two AT (AT2Bi, AT3Bi) cell lines, to generate miR expression profiles and elucidate the efficacy of miR as a biomarker of radiation response, compared to the cytogenetic and DNA damage biomarkers shown in Figures 1-3. Figure 4A illustrates a heatmap presenting the most highly expressed miR in the cell profiles, which were then further analysed to determine their increase or decrease in cells deficient in ATM relative to the mean of all cell lines (Figures 4B, 4C).

While all three cells lines showed differences in overall miR expression profiles, there were common patterns between the two AT cell lines, which differed from normally responding
MicroRNA biomarkers of radiation response

MicroRNA (miR) analysis was performed on the control (2139) and two AT (AT2Bi, AT3Bi) cell lines, to generate miR expression profiles and elucidate the efficacy of miR as a biomarker of radiation response, compared to the cytogenetic and DNA damage biomarkers shown in Figures 1-3. Figure 4A illustrates a heatmap presenting the most highly expressed miR in the cell profiles, which were then further analysed to determine their increase or decrease in cells deficient in ATM relative to the mean of all cell lines (Figures 4B, 4C).

While all three cells lines showed differences in overall miR expression profiles, there were common patterns between the two AT cell lines, which differed from normally responding cells ATM-expressing 2139 cells (Figure 4A). MiR424-5p presented the most marked differential expression between 2139 and both AT cell lines, with a 3.8-fold decrease in normally responding cells, and a 1.9-fold increase in ATM-deficient cells (Figure 4B). MiR618 also decreased in normally responding cells by 2.5-fold relative to the mean, while expression increased in both AT cell lines (Figure 4B). Conversely, miR335-3p increased in normally responding cells by 3.4-fold, with a decrease in both AT cell lines by an average of 1.7 fold relative to the mean (Figure 4C).
Figure 4: microRNA expression profiles for control (2139) and AT (AT2Bi, AT3Bi) cell lines as analysed by Exiqon, Denmark. (A) An unsupervised heatmap analysis of the 50 most highly expressed miR in all three cell lines, ranging from green to red to reflect the level of decrease or increase from the mean. Increased (B) and decreased (C) miR expression in AT cells relative to the mean of all cell lines. Data shown are representative of one independent experiment.

Analysis of differentially expressed microRNA reveals common gene targets.

A panel of targets for the most differentially expressed miR was compiled through a systematic literature search, with emphasis on genes with roles in DNA damage response and repair. ATM is an integral part of this machinery and it was hypothesised that its deficiency in AT cell lines would be reflected in an increase or decrease in expression of a panel of miR. The mean expression of miR in all three cell lines (2139, AT2Bi and AT3Bi) was calculated and each individual cell line subtracted from the mean. MiR that were consistent in expression between both AT cell lines and different from the control cells were included, with the targets for those miR also detailed. As shown in Table 2, the predominant DNA repair-associated genes identified as targets of miR increased or decreased in AT cells included the tumour suppressor phosphatase and tensin homolog (PTEN) and the G1/S cell cycle checkpoint gene cyclin D1 (CCND1). These genes were both found to be directly and indirectly regulated by the miR of interest.
**Gene biomarkers of radiation response**

Gene expression analysis of PTEN and CCND1 was carried out in normal (C1 and 2139) and AT (AT2Bi and AT3Bi) cell lines. RT-PCR was performed on cDNA isolated from all cell lines to investigate the expression of miR target genes exposed to 0, 0.05 and 0.5 Gy IR. Fold increase of genes was calculated using the $2^{-\Delta\Delta Ct}$ method, relative to 0 Gy controls and an expression was recorded over a value of 1 (Y-axis). In Figure 5, it is evident that the expression of PTEN (Figure 5A) and CCDN1 (Figure 5B) was elevated after irradiation to 0.05 Gy relative to 0 Gy in 2139 and AT cells. Normally responding C1 cells showed a modest increase in expression of both genes in response to irradiation, however the relative increase did not exceed 2.2 fold (CCND1, 0.5 Gy). The highest increase in PTEN expression was observed in AT3Bi cells, with a 35-fold increase relative to sham-irradiated cells, although this was not significant (Figure 5A). The largest increase observed in CCND1 expression was seen in 2139 cells, with a 6.4-fold increase over sham-irradiated cells. The AT cell lines showed a more modest increase of 2.6 (AT2Bi) and 4.7-fold (AT3B) (Figure 5B). However, due to inter-experimental variation, these fold changes were not significant.

**Table 2:** Expression panel of miR upregulated or downregulated in both AT cell lines compared to control cells, relative to the mean of all cell lines.

<table>
<thead>
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<th>Expression in AT cells relative to control cells</th>
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Figure 5: Expression of miR target genes *PTEN* and *CCND1*. (A) PTEN gene expression upregulated in both AT cells and 2139 compared to C1 LCL at 0.05Gy compared to 0 and 0.5Gy IR. (B) CCND1 gene expression also upregulated in AT cells and 2139 compared to C1.
at 0.05Gy. old increase of genes was calculated using the $2^{\Delta \Delta \text{Ct}}$ method, relative to sham-irradiated cells. Data shown are representative of 3 independent experiments, mean +/- SD.

**DISCUSSION**

Many advances have been made in low dose radiation research throughout this decade and through the multidisciplinary European Union DoReMi consortium (2010-2015), which arose from the original recommendations made by the High Level Expert Group (HLEG) on low dose radiation risk research (29). In particular, it was recognised that there was an urgent need for biomarkers of low dose radiation exposure, individual susceptibility and the effects of radiation damage (early and late) which have been since characterised by members of the consortium (4, 6). The authors of this manuscript were also involved in a part of DoReMi for investigating the use of Raman Spectroscopy as a novel tool and biomarker of individual radiation sensitivity. Raman Spectra can be generated from patient samples to produce a unique low dose IR-induced biochemical profile (57, 58). To validate and consolidate Raman Spectral analyses, the G2 Chromosomal radiosensitivity assay was used as a cytogenetic Biomarker of radiosensitivity because it was routinely carried out in our laboratory for different cohorts of patient lymphocytes and cell lines (35-37). In more recent years, our group has employed the use of γH2AX as a biomarker of DNA damage and individual radiosensitivity because it can yield quantitative results through flow cytometry with parallel qualitative confocal imaging and of which is more time-efficient than cytogenetics. Furthermore, previous reports show increased γH2AX foci increased with increasing radiation dose in lymphoblastoid cell lines (59). Herein we applied both cytogenetic and γH2AX biomarkers to assess the radiation sensitivity of normal (C1 and 2139) and clinically characteristed AT (AT2Bi and AT3Bi) lymphoblastoid cell lines. Lymphoblastoid cell lines (LCLs)
are T-lymphocytes immortalised with Epstein-barr virus and they were selected because parallel studies on whole blood lymphocytes from cohorts of patients were being carried out at the same time and therefore biomarker studies were limited. Although LCLs are not directly comparable to responses recorded in whole blood lymphocytes, they were advantageous for conducting the additional biomarker studies reported within. Similarly, the low doses selected for the experiments were based on the parallel blood studies that were carried out. It was surprising that the G2 chromosomal radiosensitivity scores in the AT cell lines were not significantly elevated compared to the control 2139 cells, although cell cycle checkpoint efficacy observed by mitotic indices (MI) and the calculated radiation-induced mitotic inhibition (RIMI) appeared to be superior in the 2139 cells compared to both AT cells. This would be expected if ATM is functional in the normal 2139 cells as ATM transduces the IR-induced DNA damage signal through a serine/threonine phosphorylation cascade. AT2Bi and AT3Bi cells were derived from clinically characterised AT patients and cellular features of radiosensitivity was previously established through the colony forming cell survival and chromosomal assays in which both AT cell lines showed similar spontaneous chromosomal aberration rates. However clinical and cellular heterogeneity was reported between the cell types (30). Given this reported heterogeneity between AT2Bi and AT3Bi, and the unexpected G2 chromosomal radiosensitivity response between the AT cells and 2139, a further cytogenetic analysis incorporating G-banding with karyotyping and Fluorescent In-Situ Hybridisation (FISH) using a dual ATM/TP53 probe set was performed. FISH was included in the analysis as TP53 is directly signalled by ATM phosphorylation and deletions of TP53 has been previously recored in 17% of B-cell leucocytic leukaemia (B-CLL) (40). Deletions in ATM in Ataxia-telangectasia patients have been long associated with malignancies such as leukameia and lymphomas (38, 39), and in particular older AT patients. Since both AT2Bi and
AT3Bi were derived from a 36 and 15 year old AT patient respectively, the cytogenetic FISH analysis of ATM and TP53 was warranted. Two copies each of ATM and TP53 were detected in the control 2139 and AT cells (AT2Bi and AT3Bi) in the specific cells that were analysed and therefore no specific deletion was detected. There is well-documented evidence of the heterogeneity in AT mutation types which lead to defective ATM (60-62), and a significant proportion are attributed to missense mutations which would not be detectable at the cytogenetic level and would require molecular characterisation. However, given the established presence of both copies of ATM by FISH in all cell lines, knowledge of the mutation type was not required. However, the G-banding karyotyping analysis led to a surprising incidental finding in the control 2139 cells. The absence of an X-chromosome was evident and is characteristic of Turner’s syndrome. There are conflicting reports of chromosomal radiosensitivity levels in Turner syndrome cells. In one report, 5 patients with the 45, X karyotype compared to 9 controls irradiated with X-ray (200 rads) demonstrated chromosomal aberrations similar to the controls, indicating the X-monosomy does not influence IR-induced chromosomal aberrations (63). However, another report demonstrated elevated levels of chromosomal radiosensitivity after 3 Gy IR in two comparative Turner’s syndrome variants variants (45 X complement and 46 XX gonadal dysgenesis) that were compared to age- and sex- matched controls (64). There is limited evidence in the literature to support either hypothesis. In light of this cytogenetic incidental finding, an additional control lymphoblastoid cell line (C1) was later incorporated as an additional control to 2139 where possible.

The γH2AX biomarker was utilised to measure the IR-induced DNA damage response in all cell lines (C1, 2139, AT2Bi and AT3Bi). ATM phosphorylation of the variant histone H2AX on serine 139 (γH2AX) localises as discrete nuclear foci quantifiable by immunofluorescence of which
a one to one correlation between radiation-induced DSBs and γH2AX foci can be recorded. The formation of these foci has been shown to be the recognition step for the non-homologous end joining (NHEJ) DNA repair pathway (15-17). No significant differences between the cell lines in γH2AX positivity was observed. A study on 40 human cell lines representing 8 different syndromes to detect a quantitative correlation of cellular radiosensitivity with various biomarkers; including γH2AX, reported that the IR-induced γH2AX foci did not predict moderate radiation sensitivities (65). Similarly, γH2AX foci in T-lymphocytes derived from radiotherapy-treated gynecological cancer patients did not correlate with late radiotoxicity, however the same authors reported a linear dose response with gamma radiation for whole blood and isolated T-lymphocytes (66). A recent critical review of the functional assays for individual radiosensitivity determined that γH2AX immunofluorescence alone was not sufficient to predict radiosensitive cases and that other cytogenetic biomarkers or cell survival bioassays are too time consuming to predict radiosensitivity in routine clinical use (67). This further necessitates the requirement for further molecular biomarkers.

Given the overall poor correlation of radiosensitivity with the cytogenetic and γH2AX biomarkers in our lymphoblastoid cell lines, a genetic approach was favoured but with complementarity to the previous chromosome and DNA damage biomarkers, with a focus on the ATM/chk2/P53 pathway with other DNA damage and repair mechanisms. A microRNA (miR) expression panel of 752 miR was performed on the control (2139) and AT (AT2Bi and AT3Bi) cell lines and a panel of gene targets for the most differentially expressed miR was compiled, with an emphasis on DNA damage response genes to align with our chromosome and DNA damage biomarkers related to the ATM/chk2/P53 signalling pathway. One of the limitations of this study was the reliance of only one control (2139) cell line, which was due
to the high cost associated with the microRNA experiment. Upregulated microRNAs of miR-152-3p (43), miR4-24-5p (46) and miR-92-15p (48) indicated that PTEN (phosphatase and tensin homolog) was a potential target and all downregulated miR indicated both PTEN and CCDN1 genes as potential targets. The expression of both PTEN and CCNDN1 genes were analysed in all cell lines and were shown to be upregulated expressed at the lower IR dose of 0.05 Gy. Interestingly, the C1 control showed no significant expression of PTEN compared to 2139, AT2Bi and AT3Bi, with a dose-dependent expression profile for CCND1. PTEN negatively regulates the PI3-Kinase/Akt pathway and has been associated with radiosensitivity and impaired double strand break repair in lung and prostate cancer cells (68, 69). Other studies have reported that PTEN mutations lead to radioreistant phenotypes in glioblastoma (GBM) (68) with resistance mechanisms mediated by phosphorylation of PTEN on Tyrosine240 (pY240-PTEN,) leading to DNA repair through Rad51 (70). CCND1 is the regulatory subunit of cyclin dependent kinases (CDK) which phosphorylates and inactivates retinoblastoma (RB) protein to promote cell cycle progression in the G1/S stage, and is directly signalled through the ATM/Chk2/P53 pathway. Both potential biomarkers are related to the DNA damage and repair mechanisms induced by ionising radiation and warrant further investigation and validation with more radiation doses, cell lines or biological models.

CONCLUSION

There was an unexpectedly poor correlation observed between the control 2139 cell line with the AT (AT2Bi and AT3Bi) cell lines using cytogenetic and γH2AX biomarkers, most likely due to the underlying cytogenetic abnormality identified in the control 2139 cells. However, this is not withstanding the fact that these biomarkers have proved invaluable for other associated studies carried out at our Institute (35-37, 58). When a genetic approach analysing
miR and their gene targets was taken, a better comparison could be made between the control 2139 and AT cells. This miR analysis indicated potential genetic biomarkers of radiosensitivity as well as providing mechanistic insights into the low dose radiation response particularly for 0.05 Gy. Although the speed at which molecular work can be conducted with the provision of additional mechanistic information of radiation response, it is also important that the traditional more time-consuming methods of cytogenetics and cell survival should not be overlooked. These assays are nonetheless hugely informative and reliable, and they are supported by decades of work in radiation research and in contrast, molecular technologies are advancing at a rapid rate with far less validation. When undertaking a molecular study on radiosensitivity biomarkers, we suggest a multi-biomarker approach to include optimised traditional methods with considerations for the biological model, dose-dependance and the scale of the study.

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this assay be used for rapid biodosimetry in a large scale radiation accident?


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8. Abend M, Badie C, Quintens R, Kriehuber R, Manning G, Macaeva E, et al. Examining Radiation-Induced In Vivo and In Vitro Gene Expression Changes of the Peripheral...


64. Garcia Heras J, Coco R. Chromosomal sensitivity to X-rays in lymphocytes from patients with Turner syndrome. Mutat Research 1986; 160(1):33-38


