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Expression of microRNAs in HPV negative tonsil cancers and their regulation of PDCD4

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ABSTRACT

Global rates of tonsil cancer have been increasing since the turn of the millennia, however we still have a limited understanding of the genes and pathways which control this disease. This array dataset which is linked to our publication (Zhang et al., 2015) describes the profiling of human miRNAs in tonsil and normal adjacent tissues. With this dataset, we identified a list of microRNA (miRNA) which were highly over represented in tonsil cancers and showed that several miRNAs were able to regulate the tumour suppressor PDCD4 in a temporal manner. The dataset has been deposited into Gene Expression Omnibus (GSE75630).

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Specifications Human tonsil SCC and normal adjacent tissues Organism/cell line/tissue Male and female Sex Sequencer or array type LNA Exigon array Data format Raw: GenePix Results GPR File Experimental factors Tumour vs. normal Experimental features Extraction of total RNA from tumour and normal tissues followed by miRNA profiling using custom LNA oligonucleotide arrays Consent Level of consent allowed for reuse if applicable Sample source location Svdnev. Australia

1. Direct link to deposited data

http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE75630.

2. Experimental design, materials and methods

2.1. Patient cohort

The cohort consisted of 43 patients (39 males, 4 females) treated for tonsillar cancer at Royal Prince Alfred Hospital Sydney, Australia between 2002 and 2006. The mean age was 57 years (range 39–80). Seventeen Tonsil Squamous Cell Carcinomas (SCCs) and matched microscopically normal adjacent (2 cm outside the surgical margin) tissues proved suitable for the profiling analyses (see Fig. 1 for experimental overview).

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2.2. Isolation of RNA from fresh tissue or cultured cells

Approximately 100 mg of fresh frozen tissue was diced and homogenized with a scalpel blade and then mixed with 1 mL of Trizol reagent (Invitrogen, USA). Total RNA was then extracted using isopropanol precipitation and quantified using a NanoDrop ND 1000 (Thermo Fisher Scientific, USA). Samples with ratios of 260/ 280 in the range of 1.71 to 2.1 were used for the downstream arrays.

2.3. Total RNA labeling

MicroRNAs were labeled at the 3'-end with a P-CU-C3-Cv3 RNA linker by RNA ligation as described [2,3]. 60 µL ligation reaction was prepared with 6 µg of total RNA, 0.1 mM ATP, 20 mM MgCl₂, 3.5 mM DTT, 10 mg/mL BSA, 10% DMSO, 50 mM HEPES, pH 7.8, 250 ng of P-CU-C3-Cy3 (GeneLink, USA) and 20 units of T4 RNA ligase (NEB, USA). The reaction was incubated on ice for 2 h followed by precipitation at -70 °C for 20 min with 0.3 M sodium acetate, 0.5 mg/mL glycogen (Life Technologies, USA) and 2 volumes of 100% ethanol to remove any unbound RNA-linkers. Each labeled sample was dissolved in 30 µL of 400-fold diluted ULS labeled reference set, then mixed with 300 µL Church and Gilbert hybridization buffer. This mixture was denatured at 95 °C for 2 min before hybridization. A mixture of 371 synthetic DNA reference oligonucleotides (Sigma-Genosys, Australia) containing complementary sequences to all LNA probes, was randomly labeled using the ULYSIS labeling kit (Invitrogen, USA) and then filtered using a MicroSpin™ G-25 column (Amersham, USA). Aliquots of a 400-fold dilution of labeled reference set were stored at -20 °C until needed.

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Data in Brief





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Fig. 1. Experimental overview of the project. Total RNA was extracted from cancer and normal tissues and the miRNA population was assessed using a custom printed LNA oligonucleotide array.

2.4. LNA Microarrays

A commercial LNA-modified oligonucleotides library (Exiqon, Denmark) based on miRBase release 7.1, covering 371 human and

mouse miRNA was utilized for expression profiling. Features were deposited onto GAPS II slides (Amersham, USA) at a concentration of 10 μ M (Australian Genome Research Facility, Australia). Individual miRNA LNA probes were printed four times on each array. In addition,

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