Abstract. The long non coding RNA deleted in leukemia 2 gene (Dleu2) has recently been demonstrated to be an active player in the progression of several types of cancer, including hepatocellular carcinoma. Dleu2 may serve a role in modulating the downstream effects mediated by alternative splicing of its multiple exons. However, the proportional expression of these alternative splicing populations of the Dleu2 exons is currently unknown. To determine how Dleu2 could be affected by alternative splicing, a series of alternative splicing primer sets were designed to investigate which transcripts were preferentially activated when Dleu2 was targeted for down-regulation or upregulation. A specific Dleu2 small interfering RNA that targeted an exon upstream of the tumor suppressor microRNA site significantly knocked down Dleu2 expression across all the primer sets used, which targeted 13 different alternative splicing transcripts over 5 different promoter sites in the mouse liver cell line, AML-12. Similarly, 50 µM Resveratrol led to significant upregulation of Dleu2 in 11 alternative splicing transcripts. These results show that Dleu2 is capable of successful modulation across alternative splicing transcripts that can be screened, and also that Resveratrol can be a potential nutraceutical, which may potentially lead to novel approaches in the use of IncRNA Dleu2 for diagnostics and regulation.

Introduction

Deleted in leukemia 2 gene (DLEU2) is a long noncoding RNA (IncRNA), which has been shown to serve a role as a tumor suppressor gene in several types of blood cancer (1,2). Chronic lymphocytic leukemia is characterized by DLEU2 deletion as well as its tumor suppressor microRNAs (miRNAs/miRs) miR-15a/16-1 region in ~55% of cases (3,4). As a tumor suppressor gene, the DLEU2/miR-15a/miR-16-1 locus has been extensively characterized (5-7). Since the DLEU2/miR15a/16-1 locus is present in several somatic tissues, it also has shown to serve a role in regulating cell proliferation in various cell types including, vascular endothelial cells (8), various lymphomas and leukemias (9), as well as hepatocytes (10). Physiologically, miR-15a and miR-16-1 are upregulated as a miRNA cluster from the intronic 13q14.3 region of the DLEU2 IncRNA (10). DLEU2 itself has also been implicated in the sponging of several miRNAs, including miR-30a (11) and miR-455, which can regulate several downstream functions (12). However, to the best of our knowledge, there are no studies that have assessed the effects of IncRNA DLEU2 alternative splicing in the liver.

Large-scale sequencing projects over the last decade have demonstrated the extent of alternative splicing of mammalian transcripts. Studies have shown that >95% of human genes generate transcripts that are alternatively spliced (13-15). However, not all alternatively spliced transcripts produce functional proteins (16-18). In humans, numerous studies have demonstrated an association between dysregulation of RNA splicing and the development/progression of several diseases (19,20). Certain IncRNAs have been shown to serve a crucial role in the regulation of alternative splicing in response to several stimuli or during disease (21-23). Interestingly, Dleu2 IncRNA in mice is differentially regulated, with 15 different transcripts available over five different promoter sites (Fig. S1; from ENSEMBL database: uswest.ensembl.org/Mus_musculus/Gene/Summary?db=core;g=ENSMUSG0000097589;r=13q14.3.22q13.3;dbxref=GEX:ENSMUSG00000008879) (24). MiR-15a/15b and miR-16 are encoded by introns downstream of exons present on some, but not all, of the Dleu2 transcripts (denoted by a yellow star in Fig. S1). It has previously been shown that the DLEU2/miR-15a/miR-16-1 locus contributed to cell apoptosis and progression in liver fibrosis (10), modulated miR-30a sponging in clear cell renal cell carcinoma (11), and miR-15a/16 suppression was implicated in the development of pleural mesothelioma (25).

Dependent on the size of the introns spliced out, and the exons that are retained, Dleu2 alternative splicing can have different functions on IncRNA/RNA/miRNA regulation inside the cell. In an in silico study by Ma et al (26), RNA sequencing (RNAseq) platforms showed that exon 9 of DLEU2 was a better marker than total DLEU2 expression for...
predicting unfavorable overall survival rates in patients with esophageal adenocarcinoma. Though methylation analysis of the DNA region encoding Dleu2 lncRNA with regard to several cancer types, including esophageal adenocarcinoma (26), and pediatric acute myeloid leukemia (27) have been performed, no specific studies assessing the alternative splicing populations of Dleu2 in the liver or any other tissues were found. In the present study, it was shown that Dleu2 alternative splicing transcripts were affected by silencing or overexpression, and this may improve our understanding of how Dleu2 dysregulation and modulation affect progression of various diseases.

Materials and methods

Experimental sample used for comparison of Dleu2 splicing primer sets. In vitro experiments were performed on the mouse liver cell line AML-12 using small interfering (si) RNA knockdown for downregulation of Dleu2 expression, and upregulation was achieved by treating cells with trans Resveratrol antioxidant.

Cell lines and reagents. AML-12 (ATCC® CRL-2254), the α-mouse liver cell line 12, was cultured in complete media (DMEM/F12 supplemented with 10% FBS, 1% penicillin/streptomycin, 1X Insulin/Transferrin/Selenium solution and 40 ng/ml Dexamethasone) as recommended by the supplier (all form ATCC). siRNA transfections were performed using OptiMEM media, Lipofectamine® RNAiMAX, Silencer Select Negative Control #1 (cat. no. 4390843 5 nm) and mouse Silencer Select #1 (cat. no. #4390771; sense, UGCUCUUAUAAGCAUUAAtt; antisense, UUAUACGUUAUAAGACGAcg; 5 nm) (all from Thermo Fisher Scientific, Inc.). Trans Resveratrol exposure.

RNA extraction and reverse transcription-quantitative (RT)q-PCR. RNA was extracted using the Qiagen RNeasy Mini Prep kit (Qiagen, GmbH), according to the manufacturer's protocols, and RT-qPCR was performed as previously described, using the primer sequences designed for alternative splicing and the other sequences shown below (28).

### Alternative splicing primer design and primer sets.

SiRNAs were developed to specifically target the mouse Dleu2 exon (Mouse chromosome 14, Exon 4, 61,632,437-61,632,483 bp) directly upstream of the miR-15a intron site (Thermo Fisher Scientific, Inc.). Alt1 (Dleu2 alias) primers for mouse (m) mDleu2 were designed using NCBI BLAST, targeting the largest cDNA sequences in their database (National Institutes of Health). Alt1 was designed to bind to 4 of the Dleu2 transcripts at the same time in ENSEMBL; however, this primer set did not target several of the other alternative Dleu2 splicing sets in the mouse ENSEMBL database (ensembl.org).

Therefore, 15 different spliced Dleu2 sequences were investigated, with spliced exon-exon cDNA sequences retrieved from the ENSEMBL website, and specific primer sets designed to target different Dleu2 transcripts.

These cDNA sequences were sequentially run on NCBI BLAST to assess the similarity, with only outlier exon regions not matching any other mDleu2 cDNA sequences selected. However, several primer sets still had matching regions on the larger -202 and -207 cDNA sequences yet still targeted their main splicing sequence; these primer sets were kept. These available selected areas were then screened using Primer3Plus (primer3plus.com) to find regions capable of producing standard PCR products (20-25 bp primer length, Tm 60°C, 60% GC content, 50-200 bp product size). These targeted primer sets were then screened against the BLAST alignment tool in ENSEMBL to check for other possible DNA matches in the mouse genome. Primer sets with low E values (below 0.05) that were specific for the mDleu2 gene were kept while others were discarded. Primers that matched all these criteria (Table SI) were developed and purchased from Sigma-Aldrich; Merck KGaA for use in the qPCR. Ineffective primer pairs that resulted in multiple melt curve peaks during qPCR amplification were discarded post-testing. Primers targeting specific Dleu2 transcripts are shown outlined in various colors on the ENSEMBL in Fig. 1B.

Expression of genes associated with proliferation. RT-qPCR was also used to measure expression of genes associated with proliferation, including proliferating cell nuclear antigen, transforming growth factor-β and epidermal growth factor receptor (EGFR) in Dleu2-silenced and control AML-12 cells.

Statistical analysis. All data are presented as the mean ± the standard error of the mean. Comparisons between groups were performed using a Student's t-test (two groups). P<0.05 was considered to indicate a statistically significant difference. Statistical analysis was performed using GraphPad Prism version 6.0 (GraphPad Software, Inc.).

### Results

#### Alternative splicing modulates Dleu2 for targeted primers.

The Alt1 primer set targets mDleu2-208, -203, -210 and -211,
four of the mDleu2 transcripts produced in the Dleu2 transcript map with overlapping regions (Fig. 1Ba). As such, this primer set provides a general outlook on how mDleu2 is expressed overall in the mouse under downregulating/upregulating conditions.
experimental conditions. Resveratrol treatment for 20 h resulted in significant upregulation (P<0.01) in the mDleu2 Alt1 subset, and transfection of mDleu2 siRNA significantly downregulated its expression (P<0.001; Fig. 1Aa). The mDleu2-208-209 primer set only targets mDleu2-208 in the Dleu2 transcript map (Fig. 1Ab and Bb). After 20 h Resveratrol treatment significantly upregulated (P<0.05) in the mDleu2-208 subset was observed, whereas siRNA-mediated knockdown resulted in significant downregulation (P<0.05), although the effect was less potent than that on the Alt1 set. The mDleu2-203 primer set only targets mDleu2-203 in the Dleu2 transcript map (Fig. 1Ac and Bc). The Resveratrol treatment had no significant effect on the mDleu2-203 subset, whereas mDleu2 siRNA resulted in knockdown (P<0.05), although the effect was less potent than that on the Alt1 set. The mDleu2-201 primer set targets mDleu2-210, and overlaps -202 and -207 in the Dleu2 transcript map (Fig. 1Ad and Bd). Resveratrol treatment resulted in significant upregulation (P<0.05) in the mDleu2-210 subset, and mDleu2 siRNA knockdown resulted in significant downregulation (P<0.05), although it was less significantly associated than that of the Alt1 set.

The mDleu2-201 primer set only targets mDleu2-201 in the Dleu2 transcript map (Fig. 1Ae and Bc). Resveratrol treatment resulted in significant upregulation (P<0.05) in the mDleu2-201 subset, and mDleu2 siRNA knockdown resulted in significant downregulation (P<0.05). The mDleu2-204 primer set targets mDleu2-204, and overlaps part of -202 and -207 in the Dleu2 transcript map (Fig. 1Af and Bf). Resveratrol exposure resulted in significant upregulation (P<0.01) of the mDleu2-204 subset, and mDleu2 siRNA knockdown significantly downregulated expression of this subset (P<0.01).

The mDleu2-215 TEC primer set, mDleu2-215 TEC primer set, and mDleu2-202+207 primer set targets mDleu2-215 TEC, mDleu2-213 and overlaps -215 TEC, and mDleu2-202 and -207 in the Dleu2 transcript map, respectively (Fig. 1Ag and Bg; Fig. 1Ah and Bb; Fig. 1Ai and Bi). Resveratrol treatment resulted in significant upregulation (P<0.05) of the mDleu2-215 TEC subset, mDleu2-213 subset, and mDleu2-202+207 subset; and mDleu2 siRNA knockdown resulted in significant downregulation of all three sets (P<0.05).

The mDleu2-209 primer set targets mDleu2-209 in the Dleu2 transcript map (Fig. 1Aj and Bj). Resveratrol exposure did not significantly affect the mDleu2-209 subset; however, the mDleu2 siRNA did significantly downregulate its expression (P<0.05). The mDleu2-Gm27010-201 primer set only targets mDleu2-Gm27010-201 in the Dleu2 transcript map (Fig. 1Ak and Bk). Resveratrol treatment did not have a significant effect on the mDleu2-201 subset, whereas mDleu2 siRNA did significantly downregulate its expression (P<0.05).

Expression of genes associated with proliferation. Expression of proliferating cell nuclear antigen, transforming growth factor-β and epidermal growth factor receptor was not altered significantly when Dleu2 expression was knocked down (Fig. S2).

Discussion

DLEU2 is an lncRNA that was only discovered a decade ago, and has more recently been implicated in several types of cancer, including multiple types of blood cancer, as well as hepatocellular carcinoma. As such, it is possible that changes to the expression patterns of one or several of its multiple transcripts, and thus dysregulation of the downstream lncRNAs, miRNAs and/or RNAs interactions, may lead to abnormalities. Dleu2 has been shown to regulate several physiological pathways, including proliferation (29-32), although, it does not alter proliferation of the non-cancerous AML-12 cells (Fig. S2). Interestingly, Dleu2 has a unique alternative splicing transcriptome, with multiple patterns for exon-exon splicing and intron removal (2,26). How these populations of various transcriptions arise may be important in interpreting the pathological outcomes. For example, changes in the expression levels of various exons within Dleu2 have been shown to be correlated with unfavorable overall survival rates in patients with esophageal adenocarcinoma (26). Therefore, the ability to monitor and investigate changes to these expression patterns or alternative transcription motifs may be of use in identifying dysregulated cellular biology and cancer progression.

Based on the results of the individual alternative splicing primer sets, a repeating pattern for all lncRNA Dleu2 primers used to measure Dleu2 transcript expression was observed. The mDleu2-Alt1 primer set targeted the mDleu2-208, -203, -210 and -211 transcripts, and the expression of these transcripts increased when treated with Resveratrol, and decreased when expression was knocked down using siRNAs. Most individual primer sets displayed this same pattern with significant upregulation of mDleu2 upon exposure to Resveratrol, and significant downregulation in the siRNA knockdown experiments, although there were some exceptions; specifically Resveratrol did not result in significant downregulation of mDleu2-203, -209 and -Gm27010-201, a downward trend was observed in these cases as well. Whereas the Resveratrol results showed some inconsistencies, what was noteworthy about these results was that in the Dleu2 transcripts which lacked the Exon 4 sequence, the target of the siRNAs, they were still knocked down significantly overall. Specifically, mDleu2-210, -204, -202+207, -209 and -Gm27010-201 transcripts lacked Exon 4 from the ENSEMBL alternative transcription map (Fig. S1), yet were still significantly knocked in the cells transfected with the siRNAs. It is possible that there are interactions present between the various alternative transcripts that modulate the expression of the entire Dleu2 transcript family, or perhaps the targeted siRNAs also targeted other areas similar to Exon 4 on the other transcripts. Future experiments using other targeted siRNAs that specifically target other exons on Dleu2 may provide additional information as to how this principle within Dleu2 is regulated.

Overall, the results found evidence of Dleu2 modulation across the spectrum of transcripts, with upregulation due to Resveratrol exposure and downregulation due to siRNA treatment. Upregulation of Dleu2 occurred across all transcripts generally uniformly (though not always significantly), with increased lncRNA expression from the 5'-promoter sites available (Fig. S1, bottom panel, red bars). c-Myb and PPAR are known to be positive regulators of these promoter binding sites in Dleu2 (33). Furthermore, noted shifts in splice patterns are present in diseased states as compared with the global splicing map of the human genome, which can modulate the frequency of Poly-A choices on transcripts (21).
It is possible that the limited changes in Dleu2 expression from 20 h treatment with Resveratrol and 24 h siRNA silencing treatments using the individual transcript primer sets may be due to the limited time span of the AML-12 cell culture experiments, and that longer exposures may have resulted in more significant results. Some transcripts, particularly mDleu2-203, -209 and -Gm27010-201, were not as significantly affected by Resveratrol treatment relative to the others, perhaps indicating more resistance to changes in expression compared with the other transcripts. As specific changes to DLEU2 exon expression have been confirmed in esophageal adenocarcinoma (26) and pediatric acute myeloid leukemia (27), exon expression changes are definitely a possibility in altering liver Dleu2 transcript expression. However, more specific testing is required to determine these alternative splicing changes in future experiments, perhaps using RNASeq to sidestep the limitations of overlapping targeted primer sets on the nested mouse Dleu2 transcripts. Additionally, several of these initial results need be assessed and confirmed in vivo or even using human liver tissues to identify the effects of alternative splicing.

A previous study looked at a novel alternative splicing subset within an overlapping DLEU2/LEU5/RFP2 cluster, containing multiple alternative splicing sites to produce monocistronic transcripts or a bicistronic transcript (34). Primers were designed to individually target exons of the LEU5/RFP2 cluster as well of that of DLEU2 and the overlapping sequences between the two genes. Whilst this previous study involved artificial constructs of areas surrounding exons from DLEU2/LEU5/RFP2, they were able to confirm specific intronic sequences between exons using RT-qPCR. The present study only investigated Dleu2 sequences within specific unique exons, whereas the previous study was searching for sequences spanning multiple exons, although RT-qPCR was used in both studies to build a larger exon-exon map of alternative splicing regions within and surrounding lncRNA Dleu2.

In other studies involving pediatric AML (27) and esophageal adenocarcinoma (26), alternative splicing of DLEU2 was investigated, and hypermethylation or expression of specific exons was shown to be associated with cancer progression. As such, there may be numerous other mechanisms available, in addition to simple alternative splicing, by which Dleu2 exons was shown to be associated with cancer progression. Furthermore, a nutraceutical, Resveratrol, was shown to increase the expression of Dleu2. This may assist in identifying a means of cancer prevention for certain types of cancer. Recently, Soreq et al (22) used wide annotations of alternate promoters, splicing and alternative poly-A sites to identify and quantify both disease- and treatment-induced splicing shifts, miRNA binding site modifications, putatively changed protein-protein interactions and other transcript structural changes in Parkinson's leukocytes. Future studies in in vivo mouse and human samples should expand in the above directions to investigate Dleu2 and alternative splicing in depth.

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Availability of data and materials
The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions
MC and MKK designed the study and wrote the manuscript. MKK performed the experiments. JZ analyzed and interpreted the data. All authors read and approved the final manuscript. MKK, JZ and MC confirmed the authenticity of all the raw data.

Ethics approval and consent to participate
Not applicable

Patient consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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