N-BLR, a primate-specific non-coding transcript, modulates the epithelial-to-mesenchymal transition and leads to colorectal cancer invasion and migration

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Abstract

Non-coding RNAs have been commanding increasingly greater attention in recent years as the few that have been functionalized to date play important roles in key cellular processes. Here we show that N-BLR, a ~900 bp non-coding RNA, modulates the epithelial-to-mesenchymal transition, increases colorectal cancer invasion, and functions as a migration enabler by affecting the expression of ZEB1 and E-cadherin. In patients with colorectal cancer, N-BLR expression associates with tumor stage and invasion potential. As N-BLR contains several instances of a category of DNA motifs known as pyknons, we also designed a custom-made array to investigate the possibility that other pyknon loci may be transcribed. For several of the loci probed by the array we found that the corresponding pyknons are differentially expressed between cancer and normal tissue samples. Taken together the data suggest that a systematic study of other pyknon-containing non-coding RNAs like N-BLR may be warranted in the context of colorectal cancer.

Novel experimental methods and recent technological advances have established that, in addition to the protein coding regions, significant parts of the human and other genomes give rise to ncRNAs (Bertone et al. 2004; Carninci et al. 2005; Cheng et al. 2005; Khaitovich et al. 2006). In terms of diversity, ncRNAs handily outnumber protein coding transcripts complicating functional investigations (Louro et al. 2009). Indeed, many classes of experimentally identified ncRNAs have been reported in the literature including miRNAs (Bartel 2004), piRNAs (Kim 2006), lincRNAs (Guttman et al. 2009), tiRNAs (Taft et al. 2009), moRNAs (Langenberger et al. 2009), sdRNAs (Taft et al. 2009), or long enhancer ncRNAs (Orom et al. 2010), and others. However, the full complement of ncRNAs and their functional involvement in the regulation of cellular processes and, by extension, in the onset and progression of human disorders continues to remain largely unknown (Mattick 2009).
Among ncRNAs, the best-studied group is that of microRNAs (miRNAs). MiRNAs are short RNAs, 19-23 nucleotides (nts) in length that bind their target mRNAs in a sequence-dependent manner thereby regulating the target’s expression (Ambros 2008; Bartel 2009; Rigoutsos 2009). During the last decade, miRNAs have been found to be implicated in many disease settings (Mendell 2008; Vasilescu et al. 2009; Williams et al. 2009; Small and Olson 2011) whereas more recently they were also shown to act as mediators of molecular interactions that obviate direct molecular contact (Poliseno et al. 2010). In the latter context, co-expressed transcripts (protein-coding as well as non-coding) that are targeted by the same collection of miRNAs “compete” for a finite amount of available miRNA molecules (ceRNAs) (Poliseno et al. 2010; Karreth et al. 2011; Tay et al. 2011). Changes in the abundance of any one ceRNA can affect those among the remaining ceRNAs that have expression near an underlying transition threshold (Poliseno et al. 2010; Cesana et al. 2011; Mukherji et al. 2011; Sumazin et al. 2011; Tay et al. 2011; Ala et al. 2013).

Long non-coding RNA (lncRNAs) burst onto the scene a few years ago and many are currently known in the public domain (Guttman et al. 2009; Djebali et al. 2012). Even though the full spectrum of ncRNAs remains unclear (Mattick and Makunin 2006) several lncRNAs have been shown to be important in diverse contexts such as chromatin modification and remodeling (Gupta et al. 2010a; Kogo et al. 2011), X chromosome inactivation (Lee and Lu 1999; Ogawa et al. 2008; Zhao et al. 2008; Tian et al. 2010), lineage-specific transcriptional silencing (Pandey et al. 2008), regulation of mRNA export (Chen and Carmichael 2009), activation of a growth-control gene program (Hutchinson et al. 2007; Yang et al. 2011) or of homeobox genes (Wang et al. 2011), lineage-specific silencing (Pandey et al. 2008; Latos et al. 2012), somatic tissue differentiation (Flynn et al. 2012), etc. LncRNAs have also been linked to human conditions such as brachydactyly (Maass et al. 2012) and the Prader-Willi syndrome (Bischof et al. 2007), and to cancers such as melanoma (Khaitan et al. 2011), colon (Calin et al. 2007a; Kogo et al. 2011) and prostate cancer (Prensner et al. 2011).

Pyknons (‘peak-non-s’) are a class of DNA motifs that were initially identified computationally using an unsupervised pattern discovery process (Rigoutsos and Floratos 1998; Rigoutsos et al. 2006). The motifs are ≥16 nts long and have ≥30 exact copies in the intergenic and intronic space of the genome;
what makes these motifs intriguing is that nearly all messenger RNAs (mRNAs) contain one or more pyknons in them in orientation that is sense to the mRNA suggesting the possibility that mRNAs and ncRNAs sharing the same pyknons act as ceRNAs for one another. Pyknon sequences are organism-specific: in fact, across human and mouse, we have found that the respective sets of pyknon motifs are neither syntenic nor conserved at the sequence level (Rigoutsos et al. 2006; Tsirigos and Rigoutsos 2008). Despite this absence of synteny and sequence conservation, as we showed previously, the intronic instances of human and mouse pyknons capture widespread functional conservation (Tsirigos and Rigoutsos 2008). Virtually all pyknon sequences are sense to the spliced mRNAs of some protein-coding genes and antisense to the introns of other protein-coding genes: these sense/antisense relationships between mRNAs and introns suggest intriguing possibilities for trans regulatory control (Tsirigos and Rigoutsos 2008) a possibility that received experimental support (Robine et al. 2009; Saito et al. 2009; Rigoutsos 2010). Pyknons have also been reported in plants where they are found to have the same properties as their animal counterparts (Feng et al. 2009). More recently, it was reported that the DNA methyltransferase DNMT1 binds RNA at pyknon loci and that the corresponding regions are hypomethylated (Di Ruscio et al. 2013).

Here we describe our discovery and characterization of N-BLR (pronounced: eNaBLeR), a pyknon-containing lncRNA. We examine N-BLR’s expression between normal colon and colorectal cancer (CRC) and describe its role in enabling invasion and migration by modulating the epithelial-to-mesenchymal transition (EMT). The organism-specific aspect of the pyknon motifs, the realization that some of the corresponding genomic loci are transcribed into RNAs, and the fact that a pyknon-containing transcript (N-BLR) is intimately linked to disease, prompted us to explore further the possibility of transcription at other pyknon loci as well as to investigate the possibility of a connection to disease that extends beyond N-BLR. To this end, we designed a custom microarray aimed at probing the genome at more than 1,000 pyknon loci and used it to examine evidence of transcription at those loci for several tissues, and for both normal and disease samples. It is important to emphasize that working with sequences which repeat multiple times in the human genome has its own inherent complications. We remained mindful of this fact throughout the duration of our project: in the case of the used primers, we
ensured that they are unique; in the case of the probes that we designed for our array, we intentionally mixed probes that are unique to the genome with probes that are not and discuss the rationale for this choice in more detail in the Results and Discussion sections.

Results

Transcription of genomic regions containing pyknons correlates with clinical parameters and overall survival in colorectal cancer

When we embarked on this project it was not clear whether pyknons capture “passive” DNA motifs (e.g. genomic locations to which molecules bound) or “active” sources of novel transcripts. Thus, we sought to examine the possibility of transcription by designing qRT-PCR assays for 11 genomic pyknon instances selected within regions associated with loss of heterozygozity (LOH) or within known fragile sites (Calin et al. 2004). In what follows, we denote these regions of interest as pyk-reg-14, pyk-reg-17, pyk-reg-26, pyk-reg-27, pyk-reg-40, pyk-reg-41, pyk-reg-42, pyk-reg-43, pyk-reg-44, pyk-reg-83, and pyk-reg-90 (Supp. Table 1 and Supp. Table 2). Owing to our long-standing interest in colorectal cancer (CRC), we used the 11 assays to explore the possibility of transcription across the following seven cell lines: Colo320, SW480, HCT116, IS174, HT-29, Colo205, and SW620. We observed transcription from all 11 genomic pyknon locations with expression levels that varied among the seven cell lines (Supp. Fig. 1).

These initial findings spurred us to focus our investigations on tissue samples from human normal colon and CRC. We used qRT-PCR to evaluate the 11 regions of interest in a set of 82 tumor (“sample set A” – Supp. Table 3A) and 28 adjacent normal mucosa samples of Caucasian ancestry. In this set of samples, we found significant expression differences for six of the 11 loci in cancer compared to normal tissue: pyk-reg-14, pyk-reg-40, pyk-reg-41, pyk-reg-42, pyk-reg-44, and pyk-reg-90 (Fig. 1a). Additionally, for five loci we detected significant differences between microsatellite stable (MSS) and instable (MSI) CRCs (Fig. 1b). One of the loci in particular, pyk-reg-90, stood apart from the rest: multivariate logistic regression analysis revealed a significant correlation between high expression of pyk-reg-90 and high tumor stage. The result was arrived at by analyzing 127 patient samples (we combined
sample sets A and B in order to reach the higher number of patients needed for the statistical analysis) and had an associated odds ratio=3.682, $p=0.010$ (see Supp. Table 4). Moreover, we found that high pyk-reg-90 expression was also associated with poor overall survival in the combined cohort (sample sets A and B) of 127 patients ($p=0.016$, Fig. 1c). We examined a third independent set (“sample set C”) consisting exclusively of 21 metastatic CRC patient-derived xenografts and found pyk-reg-90 to be highly expressed compared to U6 and to be present in 15 of the 21 samples ($p=0.0258$ when compared to the probability of observing this frequency accidentally, Supp. Figure 2).

The above findings in cell lines and in paired normal/CRC samples indicated that the transcript corresponding to the pyk-reg-90 pyknon instance warranted further study. In what follows, and for reasons that will become evident shortly, we will be referring to the transcript captured by the pyk-reg-90 qRT-PCR assay as “N-BLR.”

**Cloning of the N-BLR IncRNA and in situ hybridization**

The N-BLR transcript has its source in the 3p21.1-3p21.2 locus on the forward strand of chromosome 3. Targeted sequencing allowed us to zoom in on a 20 Kbp region from which we were able to clone N-BLR, an 845-nt transcript, originally from HCT116 and Colo320 cells as well as from normal colon. Subsequent Sanger sequencing confirmed that the same exact sequence, in terms of nucleotide content and length, was cloned from all three sources. This IncRNA is transcribed from the forward strand of chromosome 3 in the intergenic space between the WDR51A locus and the ALAS locus and corresponds to a contiguous block of genomic DNA (i.e. it is not spliced). WDR51A is located on the reverse strand, i.e. on the opposite strand from N-BLR and its transcription start site (TSS) is approximately 1.2 kb upstream from N-BLR (Supp. Figure 3). ALAS is on the same strand as N-BLR but more than 40 Kb downstream from it. Notably, N-BLR does not harbor any open reading frames (Hackanson et al. 2008) of significant length suggesting lack of protein-coding potential. Moreover, we verified that at the pyk-reg-90 locus transcription is preferentially derived from the forward strand i.e. it is sense to pyk-reg-90, and, thus to N-BLR (Supp. Figure 4A). We also screened for additional transcripts using primers targeting flanking regions 1, 2.5 and 5kb beyond N-BLR and in both genomic orientations: in nearly all
cases, the qRT-PCR-identified transcripts were expressed at levels lower than those of N-BLR; the only exception was in the immediate 5’ region of N-BLR, where WDR51A gene is located (Supp. Figure 4B). Lastly, we performed in situ hybridization (ISH) in CRC and in adjacent normal colon samples, using a custom-designed locked-nucleic-acid probe: ISH reveals a 2–fold higher expression of N-BLR in tumor tissue versus matched normal colon sections (Fig. 2a), in agreement with qRT-PCR findings on N-BLR expression.

**N-BLR is a novel component of the apoptotic pathway**

We selected to experiment with Colo320 cells where N-BLR is up-regulated compared to SW620 cells (see Supp. Figure 1). The expression of N-BLR in SW620 is comparable to that in normal colon so we used SW620 as control. We designed four siRNAs (labeled as N-BLR siRNA1, siRNA2, siRNA3 and siRNA4) each of which efficiently targeted N-BLR. We combined two of the siRNAs (N-BLR siRNA1 and siRNA3) in a siRNA pool that reduced the clone’s expression to less than 30%, in a dose dependent manner (Supp. Figure 5A and 5B). A non-targeting siRNA comprising a random sequence was used as a control. After treatment with the siRNA pool, N-BLR expression began decreasing at 24 hrs reaching minimum at 96 hrs in both cell lines. Cell counts of Colo320, but not of SW620, were significantly decreased at 96 hrs after treatment with either one of the single N-BLR siRNA1 and siRNA3 or the N-BLR siRNA1+3 pool (Supp. Figure 5C).

Apoptotic profiling of Colo320 cells at 96 and 120 hrs revealed significantly increased expression of cleaved PARP-1, a substrate of Caspase-3 and Caspase-7 compared to control following siRNA treatment with the N-BLR siRNA1+3 pool (Fig. 2b). Expression of the X-linked inhibitor of apoptosis (XIAP), an inhibitor of cell-death proteases Caspase-3 and Caspase-7, was abolished in Colo320 cells treated with siRNAs against N-BLR (p<0.001), but not in SW620 cells (Fig. 2c). The levels of Caspase-3, Caspase-7, Caspase-8, and Caspase-9 were significantly increased after siRNA targeting of N-BLR in Colo320 cells, but not in SW620 (Fig. 2d, 2e and 2f). The higher apoptosis in Colo320, but not SW620, was further confirmed by cell cycle analyses (Supp. Figure 5D and 5E). Thus, N-BLR is a previously unknown component of the apoptotic pathway.
N-BLR promotes invasion and migration

By transient down-regulation of N-BLR with two siRNAs, siRNA1+3 pool (Supp. Figure 6A), we found that the number of migrated HCT116 cells in a transwell assay is significantly lower when compared to control (30-40% decrease – see Supp. Figure 6B). Similarly, we found that transient transfection with two other siRNAs, N-BLR siRNA3 and siRNA4 – Supp. Figure 6C, reduced the invasion potential of HCT116 cells (Supp. Figure 6D showing E-cadherin expression). To investigate further the effect of N-BLR down-regulation during tumorigenesis, we established stable N-BLR shRNA expressing clones (labeled as clone #3-1 and clone #4-7) and used them with HCT116 cells – the qRT-PCR assays indicate that the HCT116 cells are concordant with Colo320 cells with regard to the expression of the corresponding genomic transcripts – where they significantly inhibited the expression of N-BLR (Fig. 3a). We independently tested these clones versus stable clones that carried an empty vector in transwell based motility assays and found that down-regulation of N-BLR led to a concomitant decrease by more than 50% in invasion and an over 60% reduction in migration (Fig. 3b and 3c). Following transfection with each stable clone (vs. stable transfection with empty vectors) we carried out microarray-based analysis of gene expression: we found E-cadherin to be among the most up-regulated genes, and Vimentin among the most down-regulated ones (Fig. 3d). This is notable since E-cadherin and Vimentin are involved in the EMT and cell motility control in human colon carcinoma (Greco et al. 2010). We confirmed these findings by qRT-PCR (Fig. 3e), immunofluorescence (Fig. 3f) and Western blotting (Fig. 3g). Furthermore, the down-regulation of Vimentin was paralleled by down-regulation of ZEB1 (Fig. 3g and 3h), the latter being a known master regulator of EMT in human cancers.

N-BLR is regulated by endogenous miRNAs

We next examined the possibility that N-BLR is regulated by miRNAs. In particular, we focused on the miR-200 family of miRNAs, a family with known involvement in the EMT (Gregory et al. 2008; Song et al. 2013). Of the five family members two, miR-141 and miR-200c, are endogenous and over-expressed
after transient down-regulation of N-BLR in Colo320 cells (Fig. 4a). MiR-200a is expressed at low levels and increased only in clone#3-1 but not in clone #4-7 (data not shown) and therefore not further considered for the study. These two miRNAs are also significantly over-expressed in the stable clones #3-1 and #4-7 (Fig. 4b). Furthermore, the levels of N-BLR were decreased about 70% in HCT116 cells after miR-141 transfection and about 40% after miR-200c transfection (Supp. Figure 7). To determine whether the interaction between the two miRNAs and N-BLR is direct we used the RNA22 algorithm (Miranda et al. 2006) to look for target sites in the full 845 nt sequence of N-BLR. The algorithm predicted several potential binding sites for miR-141 and miR-200c (Fig. 4c), which were confirmed by luciferase assays (Supp Fig. 8). We also measured the expression of miR-141 in CRC samples that over-expressed N-BLR and found a significantly lower expression compared to matched normal tissue samples (Fig. 4d). Finally, using ISH we identified miR-200c in matched tumor and normal samples and measured an ~50% reduction of expression in tumors versus normal, and an epithelium-specific expression (Fig. 4e).

N-BLR expression modulates the EMT

Recent work revealed a novel modality in molecular interactions where two or more transcripts ‘regulate’ one another by competing for binding to the same miRNA (or group of miRNAs); the participating transcripts are called “competing endogenous RNAs” or ceRNAs (Poliseno et al. 2010; Rigoutsos and Furnari 2010; Cesana et al. 2011; Karreth et al. 2011; Sumazin et al. 2011; Tay et al. 2011). As mentioned, the miR-200 family has been linked to the EMT: indeed its members target, among other genes, the ZEB1/ZEB2 transcription factors that are repressors of E-cadherin expression (Gregory et al. 2008). At the same time, we have shown that two of the family’s members, miR-141 and miR-200c, target and down-regulate N-BLR, which in turn has a inverse relationship with E-cadherin. Importantly, the inverse correlation between the expressions of N-BLR and E-cadherin extends to and has been observed in our cohort of CRC patients (Fig. 4f). The inverse correlation continues to hold when we compared the lymph-node positive (i.e. metastases to the lymph nodes) versus negative tumors (Fig. 4g). Together, these findings support a model whereby N-BLR decoys ZEB1 by competing with them for
binding to miR-141 and miR-200c. Consequently, an increase in the expression of N-BLR as the one we observe in the cell lines and in the CRC samples will attract the attention of a concomitantly higher number of copies from the available endogenous miR-141/miR-200c relieving pressure from and thereby up-regulating ZEB1. This will lead to a subsequent decrease of E-cadherin conferring the cells a mesenchymal phenotype that is correlated with increased invasiveness and migratory potential.

**Genome-wide profiling of pyknon transcripts**

In light of the above many and diverse observations, we conjectured that the instances of the pyknon DNA motifs might serve as ‘homing beacons’ of sorts for genomic loci from which non-coding transcripts, potentially with functional relevance, could be transcribed. We set out to investigate this possibility by designing a custom microarray. There are more than 209,000 pyknons in the human genome (Rigoutsos et al. 2006; Tsirigos and Rigoutsos 2008) that have, by design, a minimum of 30 intergenic/intronic instances (=exact copies) and at least one more exonic instance. A standardized list of all known human pyknons together with a complete list of their coordinates across the span of the human genome is available at [http://cm.jefferson.edu/pyknons.html](http://cm.jefferson.edu/pyknons.html). Among these more than 6 million instances, we randomly sub-selected a little over one thousand locations for our microarray by focusing on intergenic and intronic (i.e. non-protein-coding) instances that occur in the previously reported cancer associated genomic regions (Calin et al. 2004) (CAGRs). There are 1,292 such locations that correspond to 300 pyknons and are distributed across all chromosomes ([Supp. Figure 9](#)).

At each of the selected 1,292 locations, we centered a window of 100 nts and designed a 40-nt probe within it that overlapped with the corresponding pyknon; within the same window, we also designed a second 40-nt probe on the opposite strand that overlapped with the pyknon’s reverse complement. This choice was dictated by our earlier published observation that for several thousand pyknons their reverse complement is also a pyknon (Rigoutsos et al. 2006). Of the 1,292 chosen locations 115 did not satisfy the probe design quality control criteria. The remaining 1,177 locations are probed in both the forward and reverse orientations for a grand total of 2,354 array probes. The microarray was augmented to also include probes for all known human miRNAs.
The 40-nt probes that we used to construct the array fall into two categories: those that have unique instances on the genome and those that have multiple instances. It is worth emphasizing that both the unique and non-unique array probes are important. Indeed, the incorporation of both types was by design. Each one of the unique probes allows to unambiguously interrogate a single location and to determine whether there is a transcriptional product from that region in one or more of the examined tissues. On the other hand, non-unique probes also have merit as they permit us to interrogate multiple genomic locations simultaneously, at the expense of losing localization information. Inclusion of non-unique probes was meant to help provide information along the lines of recent findings whereby protein-coding and non-protein-coding transcripts can modulate without direct molecular contact by competing for the same miRNA(s) that target them. Each such transcript can be thought of “decoying” the remaining transcript with which it shares a miRNA-targeting site.

**Unique and non-unique intergenic probes reveal tissue-specific expression profiles**

We collected 15 normal samples from different individuals that spanned nine different tissues (4 colon, 2 breast, 1 lung, 1 heart, 1 skeletal muscle, 1 testicle, 1 liver, 2 mononuclear cells and 2 B-lymphocytes). We used our microarray to examine potential expression detected by the probes while ensuring that we distinguished among intergenic unique probes, intergenic non-unique probes, intronic unique probes and intronic non-unique probes. For the purposes of investigating the possibility of tissue-specific expression we focused solely on intergenic unique probes: we found several pyknon profiles that cluster according to the tissue of origin, which in turn suggests the existence of tissue-specific pyknon signatures (**Fig. 5a**). Not surprisingly, extending the analysis to include non-unique intergenic probes revealed similar tissue-specific expression (data not shown). This indicates that a sizable and wide-ranging spectrum of ceRNAs that compete for the attention of the same molecule, e.g. miRNA (Poliseno et al. 2010; Karreth et al. 2011; Tay et al. 2011; Ala et al. 2013; Song et al. 2013), DNMT1 (Di Ruscio et al. 2013), etc. is at play in different tissues. By using the same set of non-unique intergenic probes, compared to miRNAs, the pyknon-region transcripts exhibit higher tissue specificity in normal tissues as gauged by the Spearman
correlation (Supp. Figure 10). Moreover, across all studied normal samples, the pyknon-regions exhibit significantly higher expression levels than miRNAs: Supp. Figure 11A shows the probability density functions of the normalized miRNA and pyknon probe intensities. The higher expression levels for the pyknon-regions were also observed in multiple additional normal samples (normal colon, B-lymphocytes, and mononuclear cells) from independent individuals (data not shown). Using an independent modality (qRT-PCR), we confirmed the data obtained from the array for select pyknon-regions comparing leukemia samples with normal B cell counterparts (see for example pyk-reg-14 in Supp. Figure 11B).

Unique intergenic probes reveal differential expression between normal and disease samples

We also used our array to examine potential expression of the pyknons in disease samples. In Fig. 5b we show an example pyknon signature comprising unique intergenic probes that can distinguish healthy colon samples from microsatellite-unstable (MSI) samples. Interestingly, a subset of unique intergenic probes can correctly distinguish the two main categories of CRC, the microsatellite stable (MSS) and the MSI among the tested samples (Fig. 5c).

In addition to CRC, which is relevant for the work we presented above, we also examined CLL and normal B-lymphocyte samples, in order to explore the potential that our findings extend to other disease contexts. As with the normal samples, the expression profiles of the unique intergenic probes clustered according to the tissue of origin. Fig. 5d shows an example pyknon signature that can distinguish CLL samples from normal B-lymphocytes.

Discussion

In this work, we presented our findings on N-BLR, a non-coding RNA that is a novel member of the apoptotic pathway where it acts as an inhibitor of apoptosis. In particular, we were able to show that inhibition of N-BLR led to down-regulation of XIAP and a subsequent up-regulation of the cell-death proteases Caspase-3 and Caspase-7, both of which are inhibited by XIAP. This suggests that those CRC settings where N-BLR is up-regulated are also associated with an increased resistance to apoptosis.
Additionally, N-BLR is able to modulate the EMT. This modulation is achieved mechanistically through N-BLR’s direct interactions with at least two miRNAs (miR-141 and miR-200c) and indirect interactions with at least three mRNAs (ZEB1, E-cadherin, Vimentin). These six transcripts (N-BLR, miR-141, miR-200c, ZEB1, E-cadherin, Vimentin) represent a new signaling pathway in EMT. Our findings are supported by a recent article reporting that low expression of miR-200c on the invasive front of the primary CRCs and overexpression of miR-200c in CRC cells result in increased E-cadherin and reduced Vimentin expression (Hur et al. 2012). As we were preparing to submit our findings, another group reported analogous results for a different lncRNA in a different disease context (Yuan et al, 2014). In particular, they reported on an lncRNA, lncRNA-ATB, that promotes invasion and metastasis in hepatocellular carcinoma through interactions with members of the miR-200 family and with ZEB1/ZEB2. LncRNA-ATB is located on the reverse strand of chr14 between location 19858732 and 19861177 (hg19 assembly) and is a composite of three LINE-1 retrotransposon fragments and one full-length SINE retrotransposon. Each of these four component-fragments appears at numerous other locations on the human genome which in turn raises the possibility that very complex interactions such as those that we described in (Rigoutsos et al. 2006; Tsirigos and Rigoutsos 2008) and more recently in (Rigoutsos 2010) may be at play.

The second half of our presentation focuses on this very last point. Recall that what made us embark on the work with N-BLR was our earlier findings of a category of DNA motifs, the pyknons, together with a hypothesis that pyknon instances may pinpoint the locations of non-coding transcripts worth exploring. Indeed, the very sequence of N-BLR contains several instances of pyknon motifs. To pursue our hypothesis and investigate whether other pyknon loci on the genome show evidence of transcription we designed and built a custom microarray. The array comprised two sets of 40-nt probes: probes that were unique to the human genome, and probes that had multiple genomic instances. The rationale for the first category of probes is clear. Our choice of non-unique probes was informed by recent findings whereby protein-coding and non-protein-coding transcripts that are targeted by the same miRNA(s) can modulate one another in the absence of direct molecular interaction, a process that has been alternatively known as “decoying,” “quenching,” or “sponging” (Ebert and Sharp 2010b; Ebert and Sharp 2010a;
Poliseno et al. 2010; Karreth et al. 2011; Tay et al. 2011; Ala et al. 2013; Song et al. 2013). Even though the second group of probes cannot, without additional effort, uniquely identify the genomic source of the ncRNA(s) hybridizing to them, it does demonstrate that at least one and perhaps multiple such transcripts exist. This finding and the fact that the corresponding underlying pyknon motif is identically present in at least one mRNA (this being the very definition of pyknon motifs) raise the distinct possibility that these ncRNAs can act as molecular decoys for the corresponding mRNA. With regard to the second goal, our analyses of unique intergenic probes revealed several pyknon-containing regions whose expression profiles are tissue- and cell-state (disease/normal) dependent. This in turn indicates that there are several non-coding transcripts that are involved in specific contexts and which currently remain uncharacterized.

In closing, it is worth emphasizing that additional examination of the N-BLR locus revealed that it is primate-specific and thus not conserved in rodents. As such, N-BLR’s activity cannot be captured by mouse models of colon cancer. This represents another intriguing dimension of the intricacies of human disease and of N-BLR’s regulatory activity of the EMT. In that regard, N-BLR and other similar molecules are unlike miRNAs (Esquela-Kerscher and Slack 2006; Hammond 2007; Mendell 2008; Slack and Weidhaas 2008), transcribed UCRs (Calin et al. 2007b; Scaruffi et al. 2009; Mestdagh et al. 2010), and lincRNAs (Gupta et al. 2010b). Organism-specific transcripts can be thought of as representing a paradigm shift that is in agreement with the increasing realization that human cancers differ from animal models involving the same gene and the specific human mutation (Van Dyke and Jacks 2002). Naturally, their sequence properties make N-BLR and molecules like it promising as novel prognostic indicators.

METHODS

Methods and any associated references are available in the online version of the paper.
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Author contributions

G.A.C. and C.G.L. participated in the custom array design. G.A.C., I.R., S.K.L and S.Y.N. wrote the manuscript. All authors discussed the results and read and commented on the manuscript’s contents during the initial submission and the subsequent revisions.

**Competing financial interests**

The authors declare no competing financial interests.
**FIGURE CAPTIONS**

**Figure 1. Pyknon locus expression in CRC samples by qRT-PCR.** a, Expression and distribution of pyknon-regions were analyzed between CRC and paired normal samples (set A – see text) by qRT-PCR. b, Expression and distribution of pyknon-regions were analyzed between microsatellite stable (MSS) and microsatellite instable (MSI-H) colorectal cancer by qRT-PCR. The number of samples with measurable expression values (under Ct of 35) is presented in parentheses. The numbers of cancer and normal samples in some cases differ from one another because patients with no expression values for the U6 or for pyknon-regions were excluded. Two-sided t-test has been used to evaluate differences between two groups. Y-axis values represent ratio of each pyknon-region to U6: ratios were calculated with the $2^{-\Delta Ct}$ method using U6 levels for normalization. c, Kaplan Meier curve reveals a poor clinical prognosis for patients with high pyk-reg-90 expression ($p=0.016$, log-rank test).

**Figure 2. Properties of N-BLR.** a, *In situ hybridization* shows differential expression of N-BLR in colon cancer and normal colon: hybridization for cytokeratin 19 (CK19) and RNA for N-BLR in colon cancer and in paired normal tissue showed epithelial expression and increased N-BLR expression in tumor comparing to paired normal tissue. b, PARP-1 expression following transfection of Colo320 and SW620 cells with siRNAs (N-BLR siRNA1+3 pool) against N-BLR. Profiling was carried out at 96 and 120 hrs. c: XIAP expression following transfection of Colo320 and SW620 cells with siRNAs (N-BLR siRNA1+3 pool) against N-BLR. d, e, f: expression of Casp3/7, Casp8 and Casp9 respectively following transfection of Colo320 and SW620 cells with siRNAs (N-BLR siRNA1+3 pool) against N-BLR. Profiling was carried out after 96 and 120 hrs.

**Figure 3. The effects on invasion of N-BLR down-regulation by specific siRNAs.** a, N-BLR expression in stable clones. N-BLR expression is decreased in stably silenced clones. b, Invasion assays show significant reduction after 48 hrs for cells with stably silenced N-BLR. c, Migration assay at 24 hrs identified also significant reduction in migration of N-BLR stably silenced clones. d, Significantly differential expression of E-cadherin and vimentin identified by 44K Agilent microarray in HCT116 stable shRNA N-BLR clones #3-1 and #4-7 compared to control HCT 116 empty vector clone. e, Confirmation of
microarray data by using real time PCR shows that E-cadherin is increased and vimentin is markedly decreased in both clones (#3-1 and #4-7). f, E-cadherin and vimentin were identified in vivo by immunofluorescence with specific antibodies. Immunofluorescence signal of E-cadherin (red color) was markedly increased in both clones. The vimentin signal was present in cells with Empty vector (red color) but not in clones #3-1 and #4-7. Blue color dots indicates nucleus. g, Western blotting for vinculin (loading control), E-cadherin and ZEB1 measured in the same clones. h, ZEB1 mRNA down-regulation in HCT116 stable shRNA N-BLR clones #3-1 and #4-7 compared to control HCT 116 empty vector clone.

Figure 4. Interaction between N-BLR and miR-200 family members. a, The effect of transient transfection of N-BLR siRNA3 and siRNA4 on the miR-200 family in Colo320 cells. MiR-141 and miR-200c are increased in both N-BLR siRNAs transfected cells compared to scrambled control. b, miRNA expression in stable shRNA clones. MiR-141 and miR-200c are increased in both clones (#3-1 and #4-7). c, Schematic of the location of interaction sites between N-BLR and the miRNAs. d, miR-141 was down-regulated in CRC samples with respect to normal controls when measured by qRT-PCR. e, In situ hybridization for miR-200c in colon cancer and paired normal tissue. CK19 showed epithelial expression and U6 normalizer expression are as in Fig. 2. f, N-BLR and E-cadherin expression according to tumor and lymph node metastasis. N-BLR is increased and E-cadherin is decreased in CRC when compared to normal colon. g, The same is true when CRC with lymph node invasion (LN+) were compared with cases without lymph node involvement (LN-). Asterisks mark cases with statistically significant difference compared with scrambled (*p \leq 0.05; **p \leq 0.01).

Figure 5. a, Pyknon clusters showing tissue specificity: the clusters were formed by considering solely unique intergenic probes. b, An example of a pyknon signature formed solely of intergenic unique probes that can distinguish healthy colon samples from microsatellite-unstable (MSI) samples. c, An example of a pyknon signature formed solely of intergenic unique probes that can distinguish MSI from microsatellite-stable (MSS) samples. d, A pyknon signature formed solely of intergenic unique probes that can distinguish CLL samples from healthy B-cell samples.
REFERENCES


**a** Cytokeratin staining

- Tumor
- Normal

**b** 96h 120h

- Colo320 SW620
- N-BLR siRNA1+3 Control Scrambled
- PARP cleaved

**c** 96h

- Colo320 SW620
- N-BLR siRNA1+3 Control Scrambled
- XIAP

**d** Caspase 3/7 (96hr)

- Colo320 SW620
- N-BLR siRNA1+3 Scrambled

**e** Caspase 8 (96hr)

- Colo320 SW620
- N-BLR siRNA1+3 Scrambled

**f** Caspase 9 (96hr)

- Colo320 SW620
- N-BLR siRNA1+3 Scrambled

**Ratio of N-BLR/U6**

- Tumor (T1) Normal (M1) Tumor (T11) Normal (M11)

- 0.89 0.32 0.93 0.5
The image contains several graphs and illustrations discussing the expression of miR-141 and miR-200c in normal and tumor tissues. The graphs show the relative expression levels of these miRNAs, with comparisons between scrambled, N-BLR siRNA3, and N-BLR siRNA4 treatments.

- **Graph a**: Bar charts illustrating the relative expression of miR-141 and miR-200c in response to different treatments.
- **Graph b**: Similar bar charts for miR-141 and miR-200c expression in clone control, Clone3-1, and Clone4-7 conditions.
- **Graph c**: A ladder diagram depicting the cloned N-BLR sequence with annotations for Luc, pyk90, miR-control, miR-141, and miR-200c.
- **Graph d**: A scatter plot showing the ratio of miR-141 to U6 in normal and tumor tissues with statistical significance (p=0.0001).
- **Graph e**: Images of immunohistochemical staining for miR-200c in normal and tumor tissues, with ratios of miR-200c/U6 indicated:
  - M1, miR-200c: 0.99
  - T1, miR-200c: 0.56
  - M11, miR-200c: 0.84
  - T11, miR-200c: 0.44
- **Graph f**: Bar charts comparing N-BLR and E-cadherin expression in normal and cancer tissues.
- **Graph g**: Bar charts showing N-BLR and E-cadherin expression in LN(-) and LN(+) conditions.