

The circular RNome of primary breast cancer

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Circular RNAs (circRNAs) are a class of RNAs that is under increasing scrutiny, although their functional roles are debated. We analyzed RNA-seq data of 348 primary breast cancers and developed a method to identify circRNAs that does not rely on unmapped reads or known splice junctions. We identified 95,843 circRNAs, of which 20,441 were found recurrently. Of the circRNAs that match exon boundaries of the same gene, 668 showed a poor or even negative ($R < 0.2$) correlation with the expression level of the linear gene. In silico analysis showed only a minority (8.5%) of circRNAs could be explained by

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known splicing events. Both these observations suggest that specific regulatory processes for circRNAs exist. We confirmed the presence of circRNAs of *CNOT2*, *CREBBP*, and *RERE* in an independent pool of primary breast cancers. We identified circRNA profiles associated with subgroups of breast cancers and with biological and clinical features, such as amount of tumor lymphocytic infiltrate and proliferation index. siRNA-mediated knockdown of *circCNOT2* was shown to significantly reduce viability of the breast cancer cell lines MCF-7 and BT-474, further underlining the biological relevance of circRNAs. Furthermore, we found that circular, and not linear, *CNOT2* levels are predictive for progression-free survival time to aromatase inhibitor (AI) therapy in advanced breast cancer patients, and found that *circCNOT2* is detectable in cell-free RNA from plasma. We showed that circRNAs are abundantly present, show characteristics of being specifically regulated, are associated with clinical and biological properties, and thus are relevant in breast cancer.

[Supplemental material is available for this article.]

It is a sign of the times that the ubiquitous use of massively parallel sequencing data has delivered a parade of new insights in the cancer field and has enriched our genomic vocabulary with events like chromothripsis, kataegis, and mutational and rearrangement signatures (Stephens et al. 2011; Maher and Wilson 2012; Nik-Zainal et al. 2012, 2016; Alexandrov et al. 2013). Sequencing RNA has had less of an impact on this vocabulary, with many reports concerning traditional gene expression analysis. However, depending on the methodology of generating the sequencing library, RNA-seq has the potential to study the large variety of RNA species, including noncoding RNAs, fusion transcripts, known and novel isoforms, and, recently gaining attention, circular RNAs (circRNAs). This class of RNA was discovered many decades ago (Hsu and Coca-Prados 1979), and circRNAs were long considered idiosyncrasies of the splicing machinery processing precursor mRNA into mature mRNA. More recent studies showed an unanticipated abundance of circRNAs (Salzman et al. 2012; Memczak et al. 2013) in (normal and malignant) human cells and became particularly interesting for the cancer research field with the description (Hansen et al. 2013; Memczak et al. 2013) of a circRNA that functions as a highly potent miR-7 sponge. miR-7 has a well-described role in several malignancies, including breast cancer, and functions as a tumor suppressor in most cancers (for review, see Zhao et al. 2015) but has also been reported (Foekens et al. 2008) as a potential tumor promoter in breast cancer. Other circRNAs and additional regulatory transcriptional roles have subsequently been described in cancer (Salzman et al. 2013; Guo et al. 2014; Li et al. 2015b; Kristensen et al. 2018). Because circRNAs lack a free 5' or 3' end, such molecules escape exonucleic acid degrading enzymes, making them more stable (Memczak et al. 2013) than their linear counterparts. Therefore, circRNAs represent potentially useful biomarker candidates for diagnosis and therapy-monitoring; indeed, cell-free circRNAs are present in exosomes (Li et al. 2015a) and saliva (Bahn et al. 2015). In breast cancer, little has been described except for one study (Nair et al. 2016) using the The Cancer Genome Atlas (TCGA) data bank. However, this cohort has a huge limitation because the RNA-seq data were prepared using a poly(A) selection step, thereby omitting the majority of circRNAs (as these lack a poly(A) tail).

Here we describe the identification of an extensive catalog of circRNAs in a large cohort of 348 primary breast tumors, using RNA-seq data obtained via random-primed cDNA synthesis (Smid et al. 2016), likely preserving all the circRNAs. We developed a circRNA mapping algorithm that, in contrast to previous identification methods (Salzman et al. 2012; Memczak et al. 2013; Guo et al. 2014; Nair et al. 2016; Szabo and Salzman 2016), does not rely on unmapped reads or on known splice junctions and that was applied directly on transcriptome sequence BAM files, thereby allowing the identification of circRNAs in a genome-wide and annotation-independent (Szabo and Salzman 2016) fashion.

Results

Identification of a plethora of circRNAs in primary breast cancer

In total, 95,843 circRNAs were identified (Fig. 1), of which 27% ($n = 25,783$) had a start and end position exactly matching to an exon belonging to the same gene (Fig. 2A). The vast majority (79%) of all circRNAs were not recurrent (i.e., only found in one sample). The number of circRNAs per sample (Fig. 2B) ranged from 37 to 7105 (median, 966). For recurrent circRNAs, found in at least two samples, the range per sample was 33 to 5269 (median, 834.5). Figure 2B also shows that the number of (recurrent) circRNAs is significantly higher in estrogen receptor (ER)-negative compared with ER-positive breast cancers (Mann-Whitney U test, $P < 1 \times 10^{-5}$ for both all and recurrent circRNAs). Because of the extraordinary abundance of candidate circRNAs, we focused on the—still sizeable—number of recurrently found circRNAs ($n = 20,441$) (total number in second bar of Fig. 2A). The most frequent recurring region in our cohort was the well-characterized (Hansen et al. 2013; Memczak et al. 2013; Kristensen et al. 2018) circRNA of *CDR1*, which was found in 339 out of 348 cases. Other previously reported and validated (Salzman et al. 2012) circRNAs such as *CAMSAP1*, *FBXW4*, *MAN1A2*, *RNF220*, *ZBTB44*, and *XIST* were also identified

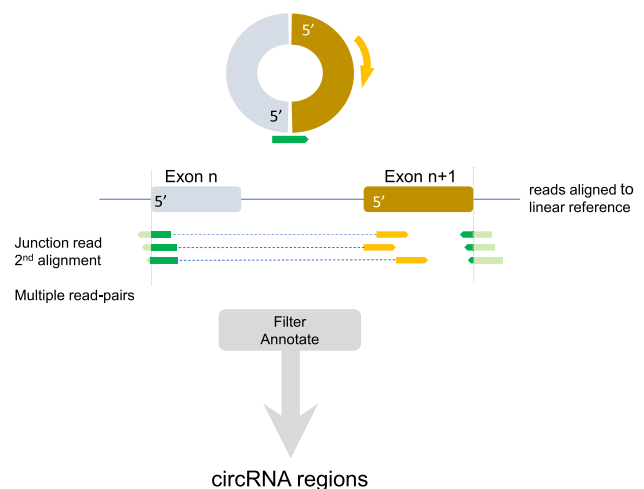


Figure 1. Schematic overview of identifying circular RNA (circRNA) regions. Assuming a circRNA molecule is present, a sequence read crossing the junction (green arrow) and its read-mate (gold arrow) would map to a linear reference in the manner depicted. The junction read would get multiple alignments, and the read-mate would be located in between the position of the junction read. Multiple read-pairs at the same junction strengthen the support for the circRNA. Subsequent additional filtering (details are in the Methods section) and annotation produced the list of circRNA regions.

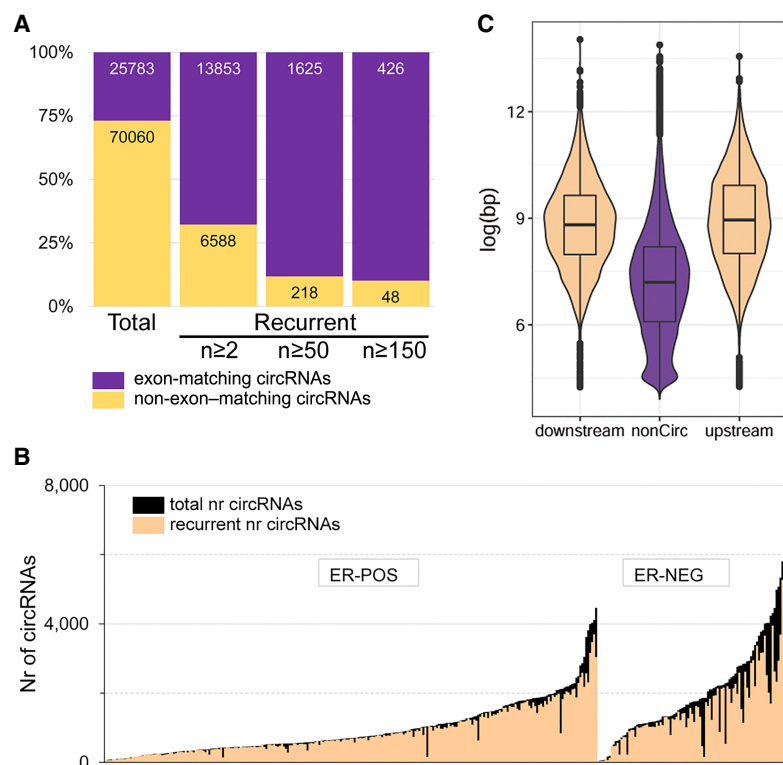


Figure 2. General characteristics of circRNAs in primary breast cancer. (A) Numbers of unique and recurrent circRNAs. Purple and gold indicate the number of circRNAs that, respectively, did or did not have a start and end position of a circRNA region exactly matching the start and end position of an exon of the same gene. (B) The number of circRNAs per sample, grouped by ER status. In black, the total number of circRNAs; in peach, the number of recurrent (identified in at least two samples) circRNAs. (C) Violin plots of the intron size (in log base pair) of noncircular regions and those located directly upstream of or downstream from a circRNA region.

in our cohort. A full list of identified circRNAs is provided in Supplemental Table S1.

General characteristics

Recurrent circRNAs were distributed across the genome, with one region on Chromosome 11 showing many closely spaced circRNAs (Supplemental Fig. S1). This region contains *MALAT1*, a highly abundant long noncoding RNA that is also frequently mutated in breast cancer (Nik-Zainal et al. 2016). Next, we evaluated the intron sizes upstream of and downstream from the circRNAs that match exon boundaries. Confirming previously reported results (Jeck et al. 2013; Zhang et al. 2014; Ivanov et al. 2015), Figure 2C shows that introns next to circRNA regions are significantly larger than introns not adjacent to circRNAs: on average 2.33 and 2.27 times larger, respectively, for introns upstream of and downstream from the circRNA (Mann-Whitney *U* test, both $P < 1 \times 10^{-5}$).

Next, we correlated the number of circRNA reads per circRNA with the expression of the respective full, linear gene, because a previous report in a limited cell line panel reported no genome-wide correlation between the circular and linear counterpart (Salzman et al. 2013). To avoid spurious correlations, only those circRNAs found in at least 50 samples were considered ($n = 1625$), and data were first normalized using the trimmed mean of *M*-values (TMM) (Robinson and Oshlack 2010). Correlations are listed in Supplemental Table S1 and ranged between -0.34 and 0.97 , with

210 circRNAs showing a negative correlation to the linear gene in which the circRNA is located. When considering the average and standard deviation of the distribution of all correlation coefficients, 30 circRNAs are at the low end of the distribution ($P < 0.05$), showing a correlation below $R = -0.182$. Finally, by using GENCODE information, the position of annotated start codons was matched to the circRNA positions. circRNAs recurring in at least 50 samples showed an almost threefold higher than expected presence of a start codon compared with circRNAs that were not recurring (χ^2 , $P < 0.0001$; 557 circRNAs; expected 207.3 circRNAs).

Are circRNAs distinct molecules, specifically regulated, or splicing residues?

Many of the circRNAs that are positively correlated with the expression level of the full-length linear transcript are thought to be a residue of splicing. Figure 3A shows an example; the circRNA of exons 3, 4, and 5 of the *WDR1* gene (Chr 4: 10,097,711–10,103,986, with a correlation coefficient of $R = 0.66$ to overall gene expression) exactly matches the difference between the known linear isoforms of this gene (Ensembl transcript ENST00000499869 includes exons 3, 4, and 5; ENST00000502702 lacks these ex-

ons). As current quantification methods do not take circRNAs into account, reads originating from the circular molecule are erroneously included in the read count of a linear isoform, resulting in an overestimation of the overall expression level. Another comprehensive example of a circRNA as splicing residue is shown in Figure 3B for *ESR1*. Full-length *ESR1* (ENST00000206249) has eight exons, whereas ENST00000406599 is a splice variant of *ESR1* that skips exons 2 to 5. circRNAs are found for several of these exons, indicating that they are likely splicing residues. We speculate that a single splice event from exons 1 to 6 generates an RNA molecule containing exons 2–5, from which a multitude of distinct circRNAs can be derived. In total, 23 patients show both a *circESR1* exon 2–3 and a *circESR1* of exon 4–5. If these circRNAs are derived from the same RNA molecule, the linear transcript would be ENST00000406599. A sequential model, in which first exons 2 and 3 are spliced out would prohibit the formation of a circRNA molecule of exons 3 to 4. However, we observed *circESR1* exon 2–3 in 110 patients and *circESR1* exon 3–4 in 64 patients, with 29 patients showing both these circRNAs. These must be derived from separate RNA molecules.

To investigate whether or not in general circRNAs should be considered splicing residues, we systematically evaluated how many of the identified circRNAs exactly match those exons that make up the difference between known linear isoforms of a gene. By using the GENCODE annotation for each gene, every possible known combination of spliced exons was matched to our circRNA catalog. Of the 25,783 circRNAs that matched to exons

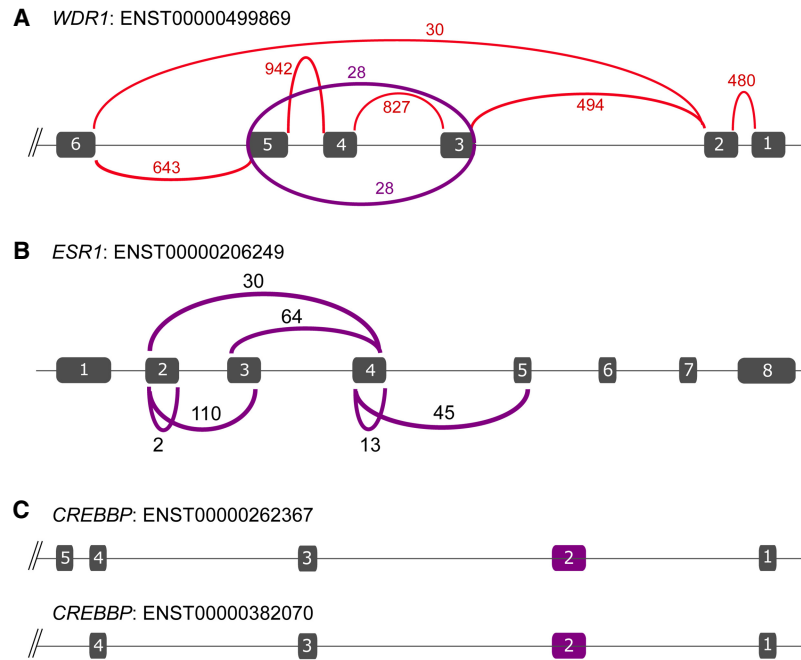


Figure 3. circRNAs are not just residues of splicing. (A) Sashimi plot of the number of reads that are aligned to *WDR1*, showing only the reads that span exons. In red are the normal exon-exon reads; in purple, the reads that span the circular junction. The line and boxes indicate the exons of the gene (the whole gene is not shown). (B) Isoform of *ESR1*. The arcs indicate the number of samples that have a particular circRNA. (C) Two isoforms of *CREBBP* that are known in the first five exons (other isoforms are described, but these start downstream from exon 5). Exon 2 (purple box) is an identified circRNA that is not a remainder of a splicing event.

of the same gene, only 2193 (8.5%) exactly matched exons known to differ between described isoforms of a gene. This was 16.9% for the circRNAs with a correlation coefficient of $R > 0.5$ to linear gene expression. This suggests that the vast majority of circRNAs that matched to exons of the same gene are generated by yet-unknown splicing events of the gene.

Because the majority of circRNAs did not match to known spliced exons, we manually inspected several highly recurrent circRNAs in the UCSC Genome Browser. For example, two isoforms of *CREBBP* are described (ENST00000262367 and ENST00000382070) that differ in the presence of exon 5 (of note, there are 10 additional known transcripts, but all of these transcripts start downstream from exon 5). However, we observed exon 2 as circRNA (Chr 16: 3,850,297–3,851,009) that was present in 160 patients (Fig. 3C), indicating that this circRNA is either specifically generated or is a splicing residue of a yet-undescribed isoform of *CREBBP* that skips this exon. Visual inspection of 10 samples that had high levels of *circCREBBP* exon 2 (at least 30 circular junction reads) showed two samples that each had one read that crossed the junction from exon 1 to exon 3, whereas the other samples showed no evidence of an isoform that skipped exon 2. This favors the notion that the circRNA of exon 2 is not a byproduct of splicing at this location.

Finally, we matched publicly available circRNA lists to gather (indirect) evidence of functional roles for circRNAs. Rybak-Wolf and colleagues reported (Rybak-Wolf et al. 2015) 4522 circRNAs in the mammalian brain that were evolutionarily conserved between human and mouse, which is considered an indication of function (Barbosa-Morais et al. 2012; Merkin et al. 2012). Of these, 2259 circRNAs (49.9%) were also present in our catalog. In addition, 3271 circRNAs were reported in an MCF-7 breast cancer

cell line panel (Tarrero et al. 2018). In total, 922 circRNAs showed (increased) expression in estrogen-stimulated MCF-7 cells compared with cells cultured in hormone-deprived medium, of which 733 circRNAs (79.5%) were present in our list. A poor correlation ($R < 0.2$) with the linear transcript was observed in our data for 78 of these circRNAs, suggesting independent regulation from their linear gene instead of ER-induced overall higher expression of all transcripts from that gene.

Validation of circRNAs

Besides the fact that we detected several already published circRNAs, thereby in part validating our method, we performed RT-PCR on a previously established independent cDNA pool of 100 primary breast tumors to confirm expression of three circRNAs, namely, *RERE* (circRNA Chr 1: 8,541,214–8,614,686), *CNOT2* (circRNA Chr 12: 70,278,132–70,311,017), and *CREBBP* (circRNA Chr 16: 3,850,297–3,851,009), all of which were poorly correlated with their linear counterpart. Figure 4 shows the PCR fragment sizes; expected and observed sizes were 89 and 155 bp for the small and bigger *CNOT2* fragment, 100 bp for *RERE*,

and 91 bp for *CREBBP*. The primer pair for *CNOT2* was able to amplify the circRNA of exon 2 to exon 3 of *CNOT2* but also the circRNA of exons 2, 3, and 4 of this gene (a circRNA that was also identified in the RNA-seq cohort).

A different primer pair to PCR *circCNOT2* showed additional fragments in addition to the expected fragments of 140 and 206 bp (Supplemental Fig. S2A). Sanger sequence analysis confirmed the circular junction sequence from exon 3 to exon 2 of *CNOT2* (Supplemental Fig. S2B) and from exon 4 to exon 2 (Supplemental Fig. S2C). After using BLAST to identify the sequence, the largest excised PCR fragment was found to contain an additional exon of *CNOT2* (Chr 12: 70,294,237–70,294,293) located between exon 2 and exon 3 that is not present in most isoforms of *CNOT2* (Supplemental Fig. S2D). The sequence showed exon 3, across the circular junction to exon 2, but reading through the location where the reverse primer was located, continuing the whole of exon 2, the additional exon, and ending in exon 3 again

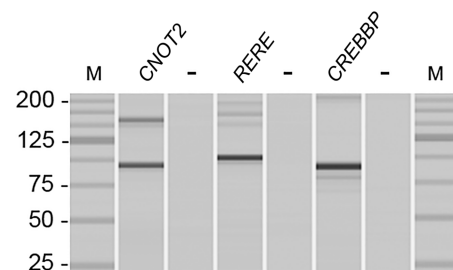


Figure 4. PCR products of circRNAs. PCR product sizes of circRNAs visualized using the MultiNA Microchip Electrophoresis System. (M) DNA size marker (25-bp fragment ladder); (-) the negative control (genomic DNA).

(Supplemental Fig. S2D). A likely explanation for this observation is that during cDNA generation, the RT polymerase generated a linear cDNA molecule containing multiple copies of the circular transcript (Supplemental Fig. S2E). In summary, the investigated circRNAs were all confirmed to be truly present in primary breast cancer.

Functional relevance of circRNAs in breast cancer cells

The potential functional relevance of the validated *circCNOT2* and *circCREBBP* transcripts, which were both poorly correlated with their corresponding linear transcript, was evaluated in breast cancer cell lines. First, expression levels of *circCNOT2* and *circCREBBP* were established in a panel of 55 cell lines, showing variable levels (Supplemental Fig. S3). Next, an siRNA was designed to specifically target the circular junction of *circCNOT2* in both MCF-7 (moderate expression level) and BT-474 (high expression level). This siRNA reduced expression of *circCNOT2* by 76% in MCF-7 and 71% in BT-474 breast cancer cells, relative to cells transfected with a nontargeting control (NTC), which resulted in significantly reduced viability of both MCF-7 and BT-474 cells (Student's *t*-test, $P < 1 \times 10^{-5}$ and $P = 4.94 \times 10^{-4}$, respectively) (Fig. 5).

circRNAs in driver genes

We matched our previously reported breast cancer driver gene list (Nik-Zainal et al. 2016) to our circRNA list. In total, 235 recurrent circRNAs were identified in 54 breast cancer driver genes. To integrate the data and obtain sufficient observations for analysis, we selected samples for which we had both RNA and genomic DNA sequencing results available and selected the genes with somatically acquired genetic events (mutations, copy number variants, and rearrangements) in at least 10 patients, yielding a list of 10 genes; *TP53*, *PIK3CA*, *PTEN*, *MAP3K1*, *CDH1*, *RB1*, *MAP2K4*, *ARID1B*, *ARID1A*, and *MLLT4*. For genes with multiple circRNAs, the circular region with the highest recurrence was chosen for analysis, with the exception of *TP53*, for which we only found two circRNAs in just one sample each (see Table 1). Only for *MAP2K4* was mutual exclusivity observed between the presence

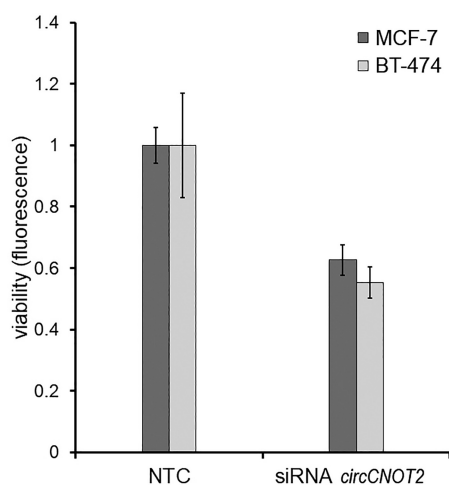


Figure 5. siRNA-mediated knockdown of *circCNOT2* affects viability in breast cancer cells. The effect of reduced *circCNOT2* expression on viability is shown in MCF-7 and BT-474 cells. Both cell lines show a significant decrease in viability ($P < 0.01$) following *circCNOT2* knockdown relative to cells transfected with nontargeting control (NTC). Error bars, SD of five wells.

Table 1. Number of samples with a DNA event and/or circRNA

circRNA region	Gene symbol	No. of samples		
		DNA events	circRNA	Both
Chr 17: 7,673,535–7,674,290	<i>TP53</i>	105	1	0
Chr 17: 7,674,859–7,676,622	<i>TP53</i>	105	1	1
Chr 3: 179,203,544–179,204,588	<i>PIK3CA</i>	84	77	25
Chr 10: 87,925,513–87,952,259	<i>PTEN</i>	47	13	4
Chr 5: 56,864,734–56,865,977	<i>MAP3K1</i>	30	210	22
Chr 16: 68,801,670–68,815,759	<i>CDH1</i>	22	7	0
Chr 13: 48,342,599–48,349,023	<i>RB1</i>	22	4	0
Chr 17: 12,054,889–12,113,360	<i>MAP2K4</i>	20	77	2
Chr 6: 156,829,227–156,935,576	<i>ARID1B</i>	14	28	1
Chr 1: 26,729,651–26,732,792	<i>ARID1A</i>	12	213	11
Chr 6: 167,870,383–167,889,326	<i>MLLT4</i>	10	28	0

of a somatic mutation or a circRNA in this cohort, in which 20 samples had a somatic mutation, 77 samples had a circRNA (Chr 17: 12,054,889–12,113,360), and only two samples had both a mutation and a circRNA ($P = 0.025$, CoMet exact test) (Leiserson et al. 2015). For *PIK3CA*, 25 patients showed a DNA event and a circRNA, three patients with a copy number aberration (amplification), and 22 patients with a base substitution in *PIK3CA*. These substitutions were located in four hotspots, p.H1047 (13 cases), p.E545 ($n = 4$), p.E542 ($n = 3$), and p.E726 ($n = 2$). None of these hotspots was located in the circRNA region that was found in these samples.

Breast cancer relevance

To investigate common biology in the samples, we used multiple correspondence analysis (MCA) to find naturally occurring subgroups. MCA is a generalized principle component analysis, suitable for categorical data. We used recurrent circRNAs (at least 50 cases) with the junction annotated to exons of the same gene ($n = 1625$) and labeled these per sample as circular or not circular, based on the presence or absence of junction reads in a sample. The main patient groups were, not unexpectedly, divided by ER status (Fig. 6A, left), whereas within circRNAs, the main division was whether or not the gene had a circRNA (Fig. 6A, right). Additional variation within the circRNAs was explained by the level of recurrence of the circRNAs (see Supplemental Fig. S4). Next, the presence/absence of circRNAs in a sample was used to cluster all samples into groups with distinct circRNA profiles. We used the gap statistic (Tibshirani et al. 2001) to determine the optimal number of sample groups, yielding six clusters (Fig. 6B; Supplemental Fig. S5). We evaluated these six sample groups on ER and tumor-infiltrating lymphocyte (TIL) status (Fig. 6C,D), number of circRNAs (Fig. 6E), and outcome for the patients in the clusters (Fig. 6F). Samples in cluster 1 and 3 were predominantly ER-negative, whereas ER positivity was predominantly present in groups 2, 4, 5, and 6. TIL status was established using a previously reported gene expression signature (Massink et al. 2015; Smid et al. 2016), labeling samples as high-TIL if the average expression of the TIL signature genes fell into the top quartile ($n = 87$, 45 ER-negative and 42 ER-positive) (Fig. 6D, respectively, labeled as red and orange). High-TIL cases were significantly ($\chi^2 P < 1 \times 10^{-5}$) more often present in clusters 1 (71% of cases) and 3 (45% of cases). Furthermore, the number of circRNAs per sample clearly distinguished the six clusters (Fig. 6E), showing a decreasing number of circRNAs from clusters 1 to 6. Finally, for a subset of 186 patients, relapse-free survival

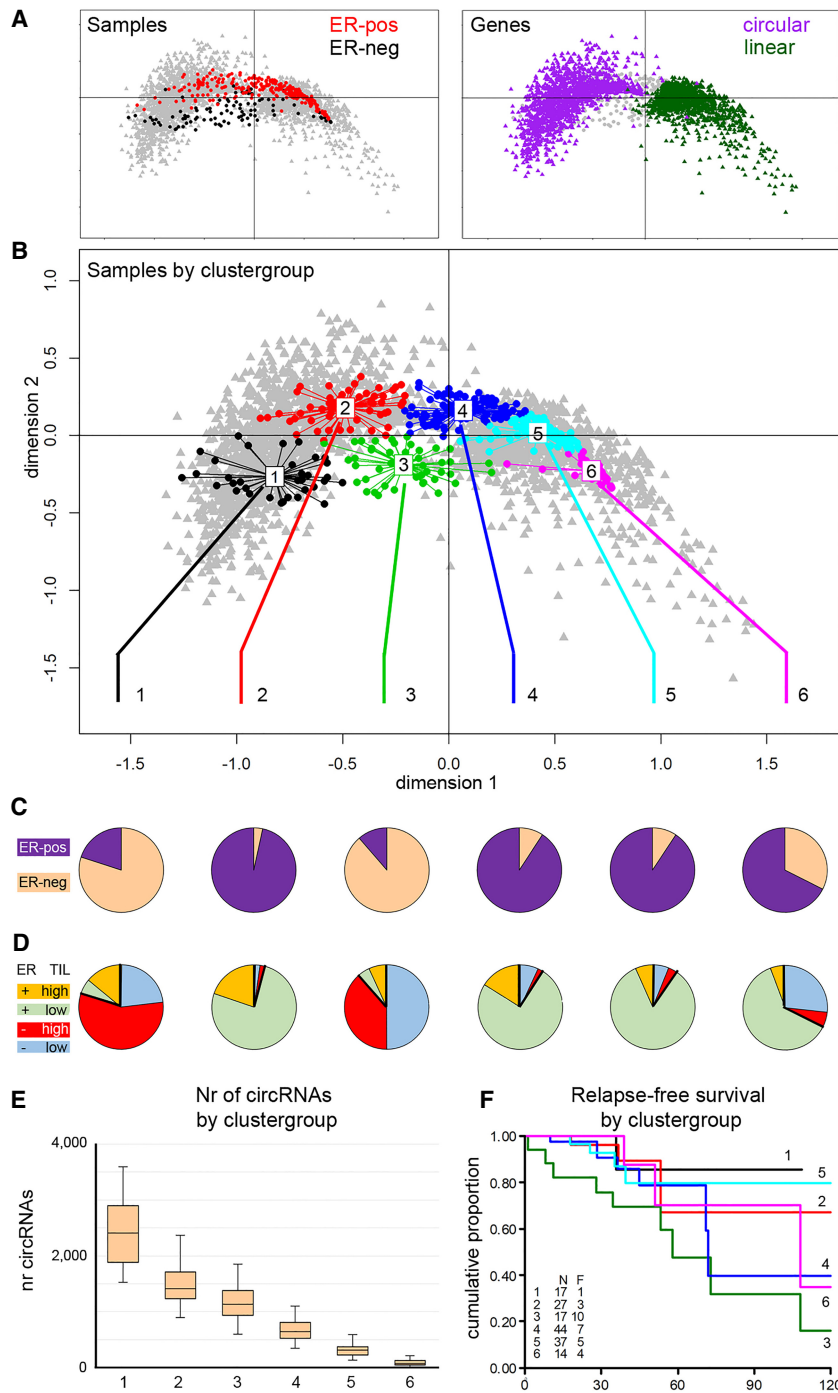


Figure 6. Analysis of sample groups according to circRNA presence. Multiple correspondence analysis (MCA) was used to find naturally occurring groups in the circRNA data. In an MCA plot, samples and circRNAs are projected onto the same plane, in which the relative distance to either the samples or the circRNAs is meaningful. The 0,0 point corresponds to a sample or circRNA with an average profile. (A, left) Samples are colored according to ER status: red, ER-positive; black, ER-negative. (Right) Purple and green indicate genes with or without circRNA expression, respectively. (B) Clustering identified samples with similar circRNA profiles; samples in the MCA plot are colored according to the cluster to which the sample belongs. (C) ER status (purple, ER-positive; peach, ER-negative) and (D) TIL status of the six sample groups: Red and orange are high-TIL cases and blue and green are low-TIL cases for ER-negative and ER-positive, respectively. (E) Number of circRNAs per sample group. (F) Relapse-free survival plot by sample group. (N) number of patients; (F) number of patients who relapse; (x-axis) months; (y-axis) the cumulative probability of relapse-free survival.

data were available; a survival plot for the six clusters (Fig. 6F) showed that the major difference was between clusters 1 and 3, which are both predominantly ER-negative. Although the number of events was low, direct comparison of cluster 1 with cluster 3 showed a significant difference in survival curves (log rank, $P=0.04$).

Differentially expressed circRNAs

We investigated if circRNA expression levels were associated with clinically relevant features of primary breast cancer, such as presence of TILs, the tumor's stromal content, proliferation, and hypoxia status. These features were inferred from generated (stroma; see Methods) or reported (Winter et al. 2007; Massink et al. 2015; Smid et al. 2016) gene expression signatures. By using these, we grouped our samples in a similar manner as explained earlier for the TIL status (Fig. 6D), labeling samples as high if the average expression of the signature genes fell into the top quartile. To identify significantly differentially expressed circRNAs, we compared the top quartile of samples to the remaining samples separately for the ER-positive and ER-negative cases. circRNAs with FDR-corrected P -values <0.05 and a fold-change greater than two were selected. Of these, the circRNAs that had a negative correlation with the linear gene expression were considered of particular interest and are listed in Table 2. Several of these circRNAs may thus potentially play a role in, or are at least connected to, the tumors that show hypoxic characteristics (e.g., *circKMT2C*) or accumulate in highly proliferative cells (e.g., *circRERE*, *circATXN2*), whereas, for example, *circASH1L* and *circPCH3* may be generated by surrounding stromal cells or infiltrating cells.

Clinical relevance

One of the reasons *CNOT2* was selected for validation was because of the poor correlation with expression of the linear gene ($R=-0.09$, $P=0.34$). This was more prominent in ER-positive ($R=-0.14$) compared with ER-negative cases ($R=0.097$). We validated this finding by making use of in-house array data (Smid et al. 2008) for linear *CNOT2* expression and a quantitative RT-PCR assay to measure *circCNOT2* (exon 2–3 Chr 12: 70,278, 132–70,311,017). The Spearman's correlation in ER-positive cases of *circCNOT2* with linear *CNOT2* was 0.079 ($n=187$, $P=0.28$) and in ER-negative cases

Table 2. Fold-change of circRNAs in top quartile of samples versus remaining samples in ER-subgroups of tumors

Regions	Gene symbol	Hypoxia		Proliferation		Stroma		TIL
		ER-negative	ER-positive	ER-negative	ER-positive	ER-negative	ER-positive	ER-negative
Chr 1: 155,438,327–155,459,898	<i>ASH1L</i>		2.0			2.1		2.0
Chr 1: 8,541,214–8,557,523	<i>RERE</i>	2.4		2.1				
Chr 1: 8,655,973–8,656,441	<i>RERE</i>				2.0			
Chr 2: 112,399,632–112,400,194	<i>RGPD8</i>	2.5	2.3		2.6			
Chr 3: 170,136,419–170,149,244	<i>PHC3</i>	3.4	2.1	2.6		2.8		2.5
Chr 5: 140,440,119–140,449,305	<i>ANKHD1</i>	2.2		2.2				
Chr 7: 152,309,966–152,315,338	<i>KMT2C</i>	2.1						
Chr 7: 1556,72,867–155,680,908	<i>RBM33</i>	2.4	2.4	2.6	2.5		2.3	
Chr 7: 17,868,407–17,875,790	<i>SNX13</i>	2.4		2.3				
Chr 10: 32,543,300–32,584,304	<i>CCDC7</i>		2.2		2.4			
Chr 12: 111,554,158–111,555,919	<i>ATXN2</i>				2			
Chr 13: 75,560,753–75,569,507	<i>UCHL3</i>							2.0
Chr 15: 25,405,461–25,411,971	<i>UBE3A</i>	2.7	2.2	2.4	2.1			
Chr 15: 92,996,957–92,998,621	<i>CHD2</i>		2.4		2.3			
Chr 18: 76,849,526–76,851,939	<i>ZNF236</i>		2.1		2.0			
Chr 20: 35,716,740–35,725,155	<i>RBM39</i>	2.5		2.2				

No significant circRNAs were identified in the TIL ER-positive group.

$R_s = 0.42$ ($n = 111$, $P = 2.9 \times 10^{-6}$), again showing absence of correlation in ER-positive cases. Thus, the role of *circCNOT2* apparently differs between ER-positive and ER-negative cases, and in the ER-negative cases, the significance of *circCNOT2* cannot be easily segregated from the linear counterpart. Therefore, we evaluated two different ER-positive breast cancer cohorts (Supplemental Table S3 shows clinical characteristics) for the potential clinical value of *circCNOT2*. We studied progression-free survival of aromatase inhibitor (AI) therapy in a multicenter cohort of 84 ER-positive patients who received this treatment for advanced disease. Quantitative reverse transcriptase-PCR (RT-qPCR) levels of *circCNOT2* showed a significant hazard ratio (HR) of 1.75 (95% confidence interval 1.32–2.33, $P = 1.06 \times 10^{-4}$), whereas RT-qPCR levels of linear *CNOT2* were not significant: HR 1.28, $P = 0.187$. Figure 7A shows a survival curve after grouping patients' *circCNOT2*-levels into three equally sized groups. A similar analysis in another cohort (Sieuwert et al. 2005), which included patients receiving first-line tamoxifen treatment ($n = 295$ patients), did not show a significant HR (0.97, $P = 0.57$) for *circCNOT2* or for linear *CNOT2* (HR 1.16, $P = 0.21$).

As circular molecules are expected to be more stable than their linear counterparts, we explored whether *circCNOT2* is a potential candidate as minimally invasive biomarker. To this end, we used cell-free RNA (cfRNA) from plasma samples of four breast cancer patients and amplified *circCNOT2* by RT-qPCR. All samples showed detectable and variable levels of *circCNOT2* (Fig. 7B), indicating that detection of circRNAs in plasma seems attainable in this exploratory setting.

Discussion

To our knowledge, we are the first to analyze RNA sequencing data using random-primed cDNA libraries from a large

primary breast cancer cohort for the presence of circRNAs, using a method that does not rely on unmapped reads. Previously, Nair and colleagues (Nair et al. 2016) analyzed TCGA RNA-seq data that were obtained using a poly(A)-based method. Although we identified 25,783 circRNAs that matched with an exon boundary of the same gene, Nair et al. (2016) reported 2146 circRNAs when we applied the same selection criteria as for our data set, and after transferring the hg37 coordinates (Nair et al. 2016) to hg38 (our data set), only 45 circRNAs were found that had the exact same start and end coordinates in both data sets. Thus, a random-primed method identifies many more circRNAs than when using poly(A)-selected material. On the other hand, there seem to be many uniquely identified circRNAs in these data sets. This could stem from differences in the methodology to detect or report the circRNAs but also could reflect the fact that many circRNAs are nonrecurrent.

We showed that circRNAs are found throughout the genome and have significantly larger-sized introns located directly

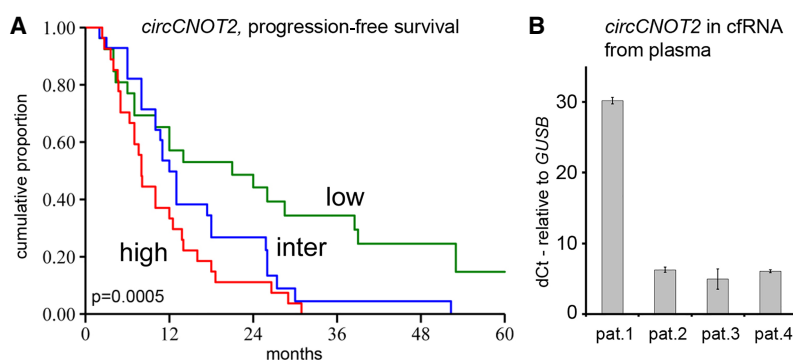


Figure 7. Clinical evaluation and presence in plasma samples. (A) Kaplan–Meier survival curve of progression-free survival for AI therapy in which patients were grouped in three equally sized groups based on their *circCNOT2* expression: red, blue, and green indicate the samples with high, intermediate, and low expression. The x-axis is in months; y-axis depicts the cumulative probability of progression-free survival on AI therapy. The P-value is the log-rank test for trend. (B) Expression levels of *circCNOT2* in plasma samples. Four metastatic breast cancer patients were evaluated. The y-axis depicts delta- C_t values of *circCNOT2* relative to *GUSB*. Error bars, SD of two measurements.

adjacent to the region on the genome that borders the circRNA, as also reported previously (Jeck et al. 2013; Zhang et al. 2014; Ivanov et al. 2015). Based on the presence/absence of circRNAs, six groups of samples were observed that differed in their ER status, TIL content, number of circRNAs, and prognosis. This indicates that there appears to be a functional biology associated with the biogenesis of circRNA molecules or at least a biology that cancerous cells can use to their advantage. Whether or not the circRNAs themselves serve that function or whether the process that generates differences in circRNA levels is the cause for the results presented here remains unknown at this time. The fact that several circRNAs were found differentially expressed in breast cancer subgroups, although these circRNAs are negatively correlated with the expression of the linear gene from which the circRNA is derived, corroborates the notion of functional circRNAs. Further experimentation is required to investigate the functional relationship of the differentially expressed circRNAs in the hypoxia, proliferation, stroma, and TIL phenotypes.

Although a synthetic circRNA construct including an internal ribosome entry site (IRES) can be translated (Wang and Wang 2015), current literature (Jeck et al. 2013; Guo et al. 2014; Szabo and Salzman 2016; Liang et al. 2017; Liu et al. 2018; Wang et al. 2018; Zeng et al. 2018) describes mostly noncoding functions for circRNAs, for example, as miRNA sponge *circCDR1* (Hansen et al. 2013; Memczak et al. 2013; Zhao et al. 2015; Kristensen et al. 2018) and *circZNF91* (Guo et al. 2014), whereas associative evidence was reported (Rybak-Wolf et al. 2015) of evolutionarily conserved circRNAs between human and mouse. Further support that circRNAs may be specifically generated and regulated was derived from a study using (estrogen-stimulated) MCF-7 cells (Tarrero et al. 2018), showing higher levels of H3K36me3 (posttranscriptional histone modification) and a higher number of Ago binding sites in circularizing exons.

Here, we contribute to the search for relevant circRNAs in three ways; first, expression levels of circRNAs that are not, or even negatively, correlated with the linear transcript of the gene may point to an intentional process. Although differences in degradation rates between the circular and linear isoforms may influence the correlation, we did not find systemic evidence for this (Supplemental Fig. S6). Furthermore, for genes that generate several distinct circRNAs, correlations can vary, indicating that degradation of the linear transcript cannot be the only explanation for the observed correlations. Second, circRNAs that include the start codon could potentially influence the expression of the linear gene, because the linear transcript from which the circRNA was spliced is now forced to use another start codon for its translation; if none is available, the transcript may be degraded. For example, *circCNOT2* (exon 2–3 Chr 12: 70,278,132–70,311,017) contains the start codon of the consensus (Pruitt et al. 2009) transcript (CCDS31857.1). Annotation shows that exon 1 is untranslated and both exon 2 and exon 4 start with the methionine codon (ATG). Thus, the linear transcript wherein exon 1 is ligated to exon 4 lacks the start codon from exon 2 and may use the ATG in exon 4 for its translation. Third, we observed that circRNAs matching exon boundaries of the same gene rarely (8.5%) overlap with known spliced exons. It could be that our analysis overlooked possible splice variants from the GENCODE annotation (both HAVANA and Ensembl exon annotations were included), but if the concordance is indeed this low, two scenarios may be applicable: Either circRNAs are still mostly a remnant of splicing, implying that many more transcript isoforms of genes exist, or otherwise circRNAs are specifically generated, implying that they do have a

biological role. The observations of variable expression levels of *circCNOT2* and *circCREBBP* in cell lines and especially the effect of *circCNOT2* knockdown on cell viability corroborate a biological role for circRNAs. Future studies are needed to systematically evaluate if the correlation between a circRNA and its linear transcript, the presence of a start codon, and/or known splice junctions are reliable criteria to prioritize circRNAs of interest.

Regardless, we were able to show clinical potential for *circCNOT2* by showing its association with the response to AI therapy. Knowing that circular molecules are not targeted by exonucleases, these molecules may be suitable candidates to be detected in cell-free environments (Li et al. 2015a), and in a pilot experiment, we showed that *circCNOT2* can indeed be detected in cfRNA from plasma samples of breast cancer patients. As such, *circCNOT2* could prove to be a useful biomarker to choose the right type of therapy or to monitor disease in a minimally invasive manner. Furthermore, we observed that very likely because of the strand displacement activity of the reverse transcriptase during cDNA generation, multiple concatemeric copies of a single circular molecule are made, contributing to the sensitive detection of circRNAs. In conclusion, we have demonstrated the abundance and potential roles of circRNAs in primary breast cancer. The methodology and selection criteria we used may help in making more sense of the seeming chaos and disorder existing in the flow from DNA to RNA to protein. circRNAs show the potential to function as relevant actors in the transcriptional regulation of RNA in addition to their promise as stable biomarkers that can be used for disease progression.

Methods

Sequencing

Internal review boards of each participating institution approved collection and use of samples of all patients in this study. RNA-seq data were generated by our consortium (Nik-Zainal et al. 2016; Smid et al. 2016) for 348 primary breast cancer tumors that are available through the European Genome-phenome Archive under accession number EGAS00001001178. Sequence protocols of the samples were previously described in detail (Nik-Zainal et al. 2016); in short, total RNA after gDNA removal, clean-up, and depletion of ribosomal RNA using duplex-specific nuclease (DSN) treatment was used as input for random-primed cDNA synthesis. Paired-end (75 bases) sequencing was performed on an Illumina HiSeq 2000. The resulting FASTQ files were mapped to GRCh38 using STAR (version 2.4.2a) (Dobin et al. 2013), and the resulting BAM files were sorted and indexed using Sambamba (version 0.6.6) (Tarasov et al. 2015; <https://github.com/lomereiter/sambamba/>). Gene annotation was derived from GENCODE Release 23 (<https://www.gencodegenes.org/>).

Identification of circRNAs

A detailed explanation of the methodology to identify circRNA reads, including the Perl script, is stated in the [Supplemental Methods](#). The script is also available at <https://bitbucket.org/snippets/MSmid/Le949d/identify-circularrna-reads>. In short, the method developed here uses sequence reads that have a “secondary alignment” (SA) tag. When using paired-end sequence data and assuming a circRNA molecule is present (Fig. 1, top), the sequence read that aligns over the crossing of the junction (green arrows) would “point toward” its read-mate (orange arrow) somewhere in the circle. Aligning these reads to the linear reference (Fig. 1, middle), the junction read will get an SA tag and will

be assigned to two locations if and only if this is the one and unique alignment configuration the STAR software can find. The read-mate aligns somewhere in between these two locations. Finding additional read-pairs showing this configuration, with a breakpoint at the exact same location, strengthens the evidence for circular transcripts. Only regions with at least five reads crossing the circular junction were included. After filtering (for details, see [Supplemental Methods](#)), GENCODE annotation was used to obtain the exon locations of genes that exactly matched to the circular region. For each sample, STAR also gives the raw read counts for all genes. These were normalized (TMM implemented in edgeR) (Robinson and Oshlack 2010) and used to correlate with the number of junction reads of the circular transcripts.

Multiple correspondence analysis

Because many genes only show a linear transcript in many samples, standard cluster analysis to identify groups of samples with similar circRNA-related biology is problematic because of the many missing values. Thus, the circRNA data were considered categorical using “circular” or “not circular” if a circRNA was present or absent in a sample. These categorical data are suitable for an MCA, a generalization of a principle component analysis. The MCA generates a combined plot that shows both patients and circRNAs such that patients/circRNAs that have similar patterns are closer together. R-packages “ade4,” “canceracm,” and “cluster” were used to perform the MCA and determine the optimum number of clusters. The latter was determined using the *clusGap* option (*k*-means to partition the samples) in the cluster package. R version 3.4.1 was used (R Core Team 2017).

Reverse transcription, PCR, and Sanger sequencing

Candidate circRNAs were selected and primers were designed such that a PCR would only yield a product when the RNA was circular, whereas in the linear situation, the primers would be divergent. Primer sequences are listed in [Supplemental Table S2](#).

First, total RNA, isolated with RNA-Bee according to the manufacturer’s instructions (CS105B, TEL TEST) was reverse transcribed into cDNA with the RevertAid H Minus first-strand cDNA synthesis kit (K1632; Thermo Fisher Scientific), followed by a RNase H step (AM2293; Ambion). Next, circRNAs were real-time PCR amplified at 10 ng input in a final volume of 25 μ L using 40 PCR cycles and an annealing temperature of 67°C with 330 nM of each primer and SensiFAST SYBR Lo-ROX mastermix (BIO-94050, Biorline), followed by a final 5-min extension at 72°C, in a MX3000P (Agilent Technologies). PCR products were visualized using a MultiNA microchip electrophoresis system (Shimadzu).

For sequencing, PCR fragments were separated on a standard agarose gel and were excised from gel using the QIAquick gel extraction kit from Qiagen according to manufacturer’s protocol. The sequencing reaction contained 2 μ L of gel-extracted PCR product, 1 μ L BigDye terminator v3.1 reaction mix (Thermo Fisher Scientific), 1 \times BigDye terminator sequencing buffer (Thermo Fisher Scientific), and 0.16 μ M of sequencing primer in a final volume of 10 μ L and was performed using an ABI2720 thermal cycler according to the following protocol: one step for 2 min at 96°C and 25 cycles of 30 sec at 96°C, 30 sec at 58°C, and 2 min at 72°C. Subsequently, the sequencing product was precipitated with absolute ethanol and 3 M of NaAc, resuspended in 20 μ L of Hi-Di formamide (Thermo Fisher Scientific), and ran on an ABI3130XL genetic analyzer (Thermo Fisher Scientific).

RT-qPCR

After RNA isolation and cDNA synthesis performed as described above, *circCNOT2* (Chr 12: 70,278,132–70,311,018) and *circCREBBP* (Chr 16: 3,850,297–3,851,009) transcripts were real-time PCR amplified at 10 ng input in a final volume of 25 μ L in 40 PCR cycles and an annealing temperature of 60°C with 200 nM of each primer and 100 nM Fam-labeled TaqMan MGB probe that covers the circular junction (Thermo Fisher Scientific) ([Supplemental Table S2](#)) in SensiFAST Probe Lo-ROX mastermix (BIO-84020, Biorline) using a MX3000P (Agilent Technologies). Levels were quantified relative to the average expression of three reference genes (*HPRT1*, *HMBS*, and *TBP*) ([Supplemental Table S2](#)) using the delta Cq method ($\Delta Cq = 2^{(\text{average Cq reference genes} - \text{Cq target gene})}$) (Schmittgen and Livak 2008). A serially diluted cDNA pool (Sieuwerts et al. 2014) of 100 independent breast tumor samples (containing both ER-positive/-negative and *ERBB2*-positive cases) was included in each experiment to evaluate the linear amplification and efficiencies for all genes and absence of amplification in the absence of reverse transcriptase. Samples in the cDNA pool were independent from the cases that were used for the RNA-seq cohort.

Detection of circRNAs in plasma

cfRNA was isolated with the Maxwell RSC miRNA tissue kit (Promega) adapted for plasma according to the manufacturer’s instructions. One milliliter of EDTA plasma of different metastatic breast cancer patients was used. These patients provided written informed consent. Six microliters of the resulting 50 μ L cfRNA (3.8–7 ng RNA/ μ L) was used to generate 20 μ L cDNA with the SuperScript IV VILO cDNA synthesis kit (Thermo Fisher Scientific). Next, 2 μ L of the cDNA was preamplified in the presence of 0.50 nM of the reverse primers of the hydrolysis probe assays for *circCNOT2* and *GUSB* as a reference marker during 15 cycles with TaqMan preamp mastermix (Thermo Fisher Scientific). Finally, 0.5 μ L of the preamplified product was measured real-time with the hydrolysis probe assays (200 nM forward primer, 200 nM reverse primer, and 100 nM FAM-labeled hydrolysis MGB probe) during 40 cycles with SensiFAST Probe Lo-ROX mastermix (BioLine) in a final qPCR volume of 25 μ L in a MX3000P qPCR machine (Agilent Technologies).

siRNA-mediated knockdown of circRNAs and cell viability assay

All cell lines in this study were established to be genetically unique and monoclonal and of correct identity by performing STR profiling using the PowerPlex 16 system (Promega). MCF-7 and BT-474 were plated at 60%–70% confluency in six-well plates and transfected with 50 nM ON-TARGETplus siRNA targeting *circCNOT2* (Horizon Discovery) using 4 μ L (MCF-7) or 8 μ L (BT-474) DharmaFECT 1 (Horizon Discovery) following the manufacturer’s instructions. Used sequences (5’-3’) were as follows: sense, AAA GAUAGGGAGACGUGGUUU; antisense, 5’-PACCACGUCUCCC UAUCUUUUU. The ON-TARGETplus nontargeting pool and ON-TARGETplus human UBB smart pool were included in each experiment as negative and positive controls, respectively (Horizon Discovery). After 24 h of transfection, cells were trypsinized, counted, and seeded in quintuplicate at 20,000 cells per well in 96-well plates. Cell viability was determined using the CellTiter-Blue cell viability assay (Promega) at day 0 and day 3. Viability measurements at day 3 were corrected for baseline viability values by subtracting the average measurement of day 0.

Gene expression signatures

We used several signatures: a TIL and proliferation signature (Smid et al. 2016), a hypoxia signature (Winter et al. 2007), and a stroma-specific signature using public data GSE5847 (Gene Expression Omnibus) (Boersma et al. 2008). We performed a paired *t*-test to obtain genes significantly higher expressed in microdissected stroma (FDR < 0.05 and fold-change > 1.7). For all signatures, genes that were up-regulated in the category of interest were matched to our data set, and the average expression of the signature genes was calculated per sample. Samples were labeled as high-TIL (or stroma, proliferation, hypoxia) if the average expression of the signature genes fell into the top quartile. To identify significantly differentially expressed circRNAs, we compared the top quartile of samples versus the rest, per ER-group. circRNAs were only included when detected in >50% of the samples and matched known exon locations of the same gene. Analyses were performed using BRB-ArrayTools developed by Dr. Richard Simon and the BRB-ArrayTools development team. circRNAs were considered significant when the FDR corrected *P*-value was below 0.05 and the fold-change greater than 2.

Breast cancer cohort treated with endocrine therapy

RT-qPCR was performed on a linear and circular isoform of *CNOT2* (Chr 12: 70,278,132–70,311,017) in a first-line TAM (Sieuwerts et al. 2005) and a first-line AI cohort to study the predictive value of *circCNOT2* on therapy response. The AI cohort was a multicenter cohort consisting of 30 patients from Erasmus MC, Rotterdam; 35 patients from The Netherlands Cancer Institute, Amsterdam; and 19 patients from the Translational Cancer Research Unit, Antwerp (Belgium). All 295 patients in the TAM cohort are patients of the Erasmus MC, Rotterdam. Patient characteristics are listed in Supplemental Table S3.

Statistical analyses

STATA version 14 was used to perform the statistical tests that are indicated in the text. *P*-values are two-sided, corrected for multiple testing when necessary, and considered significant below 0.05.

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