Next Generation Sequencing of Serum Circulating Nucleic Acids from Patients with Invasive Ductal Breast Cancer Reveals Differences to Healthy and Nonmalignant Controls

Julia Beck1, Howard B. Umovitz1, William M. Mitchell2, and Ekkehard Schütz1

Abstract

Circulating nucleic acids (CNA) isolated from serum or plasma are increasingly recognized as biomarkers for cancers. Recently developed next generation sequencing provides high numbers of DNA sequences to detect the trace amounts of unique serum biomarkers associated with breast carcinoma. Serum CNA of 38 women with ductal carcinoma was extracted and sequenced on a 454/Roche high-throughput GS-FLX platform and compared with healthy controls and patients with other medical conditions. Repetitive elements present in CNA were detected and classified, and each repetitive element was normalized based on total sequence count or repeat count. Multivariate regression models were calculated using an information-theoretical approach and multimodel inference. A total of 423,150 and 953,545 sequences for the cancer patients and controls, respectively, were obtained. Data from 26 patients with stages II to IV tumors and from 67 apparently healthy female controls were used as the training data set. Using a bootstrap method to avoid sampling bias, a five-parameter model was developed. When this model was applied to a validation data set consisting of patients with tumor stage I (n = 10) compared with healthy and nonmalignant disease controls (n = 87; 1,261,561 sequences) a sensitivity of 70% at a specificity of 100% was obtained. At a diagnostic specificity level of 95%, a sensitivity of 90% was calculated. Identification of specific breast cancer–related CNA sequences provides the basis for the development of a serum-based routine laboratory test for breast cancer screening and monitoring.

Introduction

Breast cancer is one of the most frequent malignancies among women and accounts for one in four cancer occurrences in adult women (1). Screening methods and adjuvant therapy following surgery led to a remarkable decrease in mortality between 1975 and 2000 (2). Nevertheless, ~185,000 new cases and 41,000 deaths from breast cancer are estimated to have occurred in 2008 (3). In addition to physical examination, detection of breast cancer has relied on imaging technologies for screening that have reached the apparent limits of their capacity and/or have low societal cost-benefit ratios (i.e., magnetic resonance imaging). The prime example is the decreased sensitivity of standard mammography in young women and women with dense parenchyma (4). Results from seven population-based community screening programs in the United States revealed an overall sensitivity of 75% and a specificity of 92.3%, although sensitivity in women with extremely dense breasts dropped to 63% (5). This is further influenced by a decrease in the use of mammography from 2000 to 2005 (6), with pain caused by breast compression during the procedure cited as a common complaint (7, 8). Development of minimally invasive molecular techniques to supplement imaging procedures for the early diagnosis and treatment of breast cancer and the consequent reduced mortality has remained elusive despite the recent remarkable advances in molecular biology.

The risk for breast cancer in humans has been associated with a limited number of gene mutations and gene expression signatures. High-risk single-gene alleles of BRCA1, BRCA2, PTEN, and TP53 are relatively rare in the general population and account for less than 25% of the familial clustering observed in breast cancer (9). Other reported breast cancer–associated single nucleotide polymorphisms include FGFR2, TNRC9, MAP3K1, LSP1, and CASP8, which in combination might distinguish women at high risk for the development of breast cancer (10). Tissue samples of established breast carcinomas can be evaluated for metastatic potential by gene expression signatures (11). Unfortunately, none of these

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**Authors' Affiliations:** 1Chronix Biomedical GmbH, Goettingen, Germany and 2Department of Pathology, Vanderbilt University, Nashville, Tennessee

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**Corresponding Author:** Ekkehard Schütz, Chronix Biomedical GmbH, Goetheallee 8, 37073 Goettingen, Germany. Phone: 49-551-3707-5722; Fax: 49-551-3913-968. E-mail: esc@chronixbiomedical.de
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known genetic risk associations is of sufficient diagnostic value for clinical use.

Cell-free circulating nucleic acids (CNA) isolated from blood plasma or serum have become increasingly recognized as potential biomarkers for the early detection and clinical monitoring of various human cancers. The first report that DNA could be detected in the blood of cancer patients occurred in 1948 (12), although efforts to explore this observation for diagnostic purposes did not occur until the last decade. Measurements of absolute levels of CNAs have been suggested for the diagnosis (13) and prognosis (14) of breast and lung cancer (15). The general diagnostic value of simple quantitative measures of CNA, however, is controversial because nonspecific CNA elevations are seen in patients with benign diseases (16–19). Cancer-specific DNA perturbations such as microsatellite instability, mutations, sequence length, and promoter methylation patterns detected in serum/plasma have been proposed for the diagnosis and clinical assessment of cancer treatment (20).

In this report, we show that the quantities of specific cell-free transposable elements and endogenous retroviral DNA sequences in blood could distinguish all stages of invasive ductal breast cancer from normal controls and nonmalignant controls with sensitivities and specificities that are promising as a useful clinical tool.

Materials and Methods

Specimen

Serum samples were obtained from 38 women with breast cancer and 67 healthy female donors. All patients were histologically diagnosed with infiltrating ductal carcinoma. The group was comprised of 10, 20, 2, and 6 patients with clinical stages of I, II (including IIA and IIB), III, and IV, respectively. Tumor staging and histology was used as given in the patient’s records. Blood was drawn preoperatively from treatment-naïve patients. Thirty-two of the patient samples were collected at the Cleveland Clinic satellite facility in Florida, USA. A detailed patient description is given in Supplementary Table S1. Sequence data from 28 apparently healthy control samples were collected. Six breast cancer samples were collected at the Cleveland Clinic satellite facility in Florida, USA. A detailed patient description is given in Supplementary Table S1. Sequence data from 28 apparently healthy female controls were previously generated and stored in a CNA database that was described recently (21). Additional 87 collected samples of patients and controls were analyzed as an independent noncancerous control set and 31 samples collected from a patient with multiple myeloma and liver sarcoma served as a malignant control (Table 1). All donors and patients provided their informed consent.

Sampling

Serum samples were collected as described (22) and stored at −80°C prior to further processing. Briefly, frozen serum was centrifuged at 4,000 × g for 20 min and 200 μL of the supernatant was used in the High Pure Nucleic Acids Extraction Kit (Roche) according to the instructions of the manufacturer.

Generation of Random DNA Libraries, Sequencing, and Sequence Analysis

The serum DNA was processed, sequenced, and analyzed as described previously (21). Briefly, extracted DNA was amplified using the GenomePlex Single Cell Whole Genome Amplification Kit (WGA4, Sigma-Aldrich) in duplicates, which were pooled for subsequent sequencing. Although probably not necessary for CNA, the shearing procedure as included in the WGA4 protocol was done for comparability with, e.g., genomic results. Previously, we have tested the WGA4 procedure using either unsheared or sheared genomic DNA and could not detect essential differences in sequencing results (21). Using the adapters that are attached during the whole genome amplification procedure, a sample-specific molecular barcode was attached to the amplicons by PCR. The resulting DNA preparations were pooled and sequenced using a Roche/454 GS-FLX high-throughput sequencer. After sequencing, the sequences were assigned to the individual patients and control samples according to the 10-bp tag sequence. An overview of the further sequence analysis pipeline is depicted in Supplementary Fig. S1.

Statistical Procedure

Normalization for each sample and each repetitive element (per assigned nucleotide) was done either on the basis of the total assigned nucleotides of human genomic origin or of human repetitive element origin (Supplementary Fig. S1). The normalized values were used to construct receiver operator characteristics (ROC) curves for each repetitive element present in at least 95% of the serum samples. The C value was calculated as area under the receiver operating characteristic (ROC) curve by applying the linear trapezoidal rule (23). Repetitive elements with a C value of at least 0.62 were considered as independent variables for the subsequent calculation of binominal multivariate regression models. The general linear model was fitted using a training set comprising normal female controls (n = 67) and patients with tumor stages II to IV (n = 28). An information-theoretical model selection with subsequent multimodel inference approach was taken (24). First, the corrected Akaike’s information criterion values were calculated for all possible models of two to five predictors (e.g., 174,500 when 30 independent variables were considered). Akaike’s information criterion values were calculated, according to Burnham and Anderson (25), as transformation of the residual sum of squares of the linear multivariate regressions. These values were used to calculate normalized Akaike weights (ω) in such a way that they sum up to 1 for all models as described (25).
Those models accounting for the lower 97.5% of the 5% trimmed $\omega$ sum were discarded. For each predictor (independent variable) Akaike $\omega$ of all remaining models containing that predictor were summed ($\sum \omega$). The six variables with the highest Akaike $\sum \omega$ were used to constitute the final model. For these six variables, the respective estimates were averaged (25). In an additional approach, to further reduce the chance of random group effects on modeling, a random resampling strategy was applied. On average, each sample was resampled into 172 bootstrap rounds (SD, 9), in which 46 normal controls and 19 stage II to IV patient samples were used in each round. In total, 255 rounds were done that yielded $3.7 \times 10^6$ models, of which for each calculation the best (highest $\omega$) 1%, but not less than 40 models, were selected and the predictor estimates were recorded as given above. This resulted in $\sim 1.1 \times 10^5$ models, from which the sum of $\sum \omega$ per independent variable was calculated. Again, the six predictors with the highest weight sums were used for model definition, in which average estimates of independent variables were calculated as above.

For each study sample, the calculated intercept and the predictor estimates (slope) were used to calculate the model value of the samples. To assess the validity of the achieved models, samples originating from patients with tumor stage I ($n = 10$) were used as a validation set, together with samples from individuals with other and nonmalignant medical conditions. The performance of the obtained models was assessed by the generation of ROC curves from those calculated model values. Sensitivity and specificity were derived at the point of best separation, which was defined as the maximized Youden’s index value (26).

**Results**

**Sequencing Results**

A total of 423,150 sequences for cancer patients and 953,545 sequences for apparently healthy female controls were obtained, totaling $7.5 \times 10^7$ and $1.61 \times 10^8$ nucleotides, respectively. Together with the additional controls, a total of $2.8 \times 10^6$ sequences were analyzed herein. Supplementary Table S2 gives a summary of the average fragment and nucleotide counts in each group and the average number of nucleotides of human genomic origin. Nucleotides of human origin are defined as nucleotides that were either identified as human repetitive elements by RepeatMasker (27, 28) or produced significant hits after BLAST (29) searching of the human genome.

**Representation of Repetitive Elements in the CNA**

Generally, the serum CNA pool of patients with breast cancer contains significantly more repetitive elements (52%; SD, 0.8%) than the serum CNA of normal control subjects (51%; SD, 1%). Although small, the difference is significant ($P = 0.0013$; median test) when all patients with invasive carcinomas in stage II to IV were compared with the healthy female control cohort.

Two different normalization approaches were taken to compare repetitive element representation. The normalization on total repetitive elements instead of total genomic

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### Table 1. Sample sets used for model development and validation

<table>
<thead>
<tr>
<th>Training/model development</th>
<th>Name of sample set</th>
<th>$n$</th>
<th>Gender (female/male)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal female training set</td>
<td>67</td>
<td>67/0</td>
<td>Samples were obtained from apparently healthy women</td>
<td></td>
</tr>
<tr>
<td>Stage II to IV training set</td>
<td>28</td>
<td>28/0</td>
<td>Contains 20 samples of tumor stage II, 2 samples of tumor stage III, and 6 samples of tumor stage IV</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Validation</th>
<th>Name of sample set</th>
<th>$n$</th>
<th>Gender (female/male)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal female training set</td>
<td>67</td>
<td>67/0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast cancer validation set</td>
<td>10</td>
<td>10/0</td>
<td>Contains only patients with tumors in stage I, which were not included in the training set</td>
<td></td>
</tr>
<tr>
<td>Noncancerous validation set</td>
<td>87</td>
<td>34/53</td>
<td>Combination of independent healthy controls and patients with other noncancerous medical conditions. Other medical conditions include multiple sclerosis ($n = 34$), diabetes mellitus ($n = 1$), and autism ($n = 1$)</td>
<td></td>
</tr>
<tr>
<td>Multiple myeloma</td>
<td>31</td>
<td>0/31</td>
<td>Samples were obtained from one male patient at different time points</td>
<td></td>
</tr>
</tbody>
</table>
hit length identifies which repetitive elements are overrepresented or underrepresented within the group of repetitive elements, in which AT-rich sequences show the largest deviation in both normalizations. A ROC curve was calculated for each repetitive element. A ROC curve plots the true positive rate against the false positive rate for the different diagnostic cutpoints and the area under the curve (C value) is a measure of test accuracy. Supplementary Table S3 summarizes the ROC curve data of those repetitive elements that yielded C values of >0.62 for the two different normalizations. The data shows that there is no single repetitive element that can be used alone to obtain good separation between patients with breast cancer and controls.

### Binomial Multivariate Regression Modeling

The repetitive elements listed in Supplementary Table S3 were subsequently used for developing binomial multivariate models. Models were calculated for normalization by the number of repetitive nucleotides (modelnormRE) and the total hit length of genomic origin (modelnormGO).

The models were fitted using data from the stages II to IV training set versus the normal female training set (Table 1). Table 2 gives the predictors, weight sums, and respective averaged estimators for both models, together with the C statistic from ROC curves.

The stage I patient data was subsequently used as an independent data set (breast cancer validation set) to validate the model performance. Table 3 provides the diagnostic performance characteristics of modelnormRE and modelnormGO in the different patient groups. The modelnormRE yielded a higher C value than modelnormGO when applied to the breast cancer validation set (Table 3).

Further validation of the model predictors employed a random resampling approach (modelRRnormRE), generated from the initial training data set. As illustrated in Table 2, the predictors of modelnormRE with the exception of FLAM-C were validated by the modelRRnormRE. FLAM-C was replaced by MLT2A2, which had a higher \( \Omega \omega \) in the random resampling modeling. Following the same approach as conducted for the first two models, the modelRRnormRE was applied to the breast cancer validation set (stage I) and to the combined data set (stage I-IV) using ROC curve analysis. The performance characteristics for the three different groupings of the cancer samples compared with the normal female training set, are given in Table 3. Supplementary Fig. S2 shows the calculated model values and the ROC curves for this model. Although the C value of modelRRnormRE (0.9; confidence interval, 0.76-0.97) in the stage II to IV training set was lower than that of modelnormGO (0.94; confidence interval, 0.8-0.99), the former yielded a slightly higher C value when the breast cancer validation set was compared with the normal female training set.

### Model Validation Using an Independent Control Group

The modelRRnormRE was analyzed additionally in a cohort of 87 individuals that had noncancerous diseases (1.26 \( \times 10^8 \) sequences with 2.1 \( \times 10^8 \) nucleotides) and in a multievent time course of a patient suffering from multiple myeloma (\( n = 31; 320,342 \) sequences with 5.8 \( \times 10^7 \) nucleotides). A detailed description of the samples is given in Table 1. The distribution of the obtained scores are shown in Fig. 1. The two different breast cancer patient groups are displayed for comparison. It is obvious that neither the multiple myeloma nor the independent nonmalignant control set is associated with serum DNA sequences.
similar to invasive ductal carcinoma. From ROC curve analysis using the noncancerous validation set versus the breast cancer validation set, a C value of 0.986 (0.927-0.998), with a specificity of 95% and a sensitivity of 90%, was calculated. At 100% specificity, a corresponding sensitivity of 70% was obtained.

Discussion

High-throughput sequencing of total serum DNA shows differential representation of certain repetitive elements in the CNA of patients with breast cancer compared with healthy controls. The comprehensive database generated allowed the simultaneous comparison of all known repetitive elements of patients with breast cancer versus normal and nonmalignant controls. The normalized nucleotide counts of repetitive elements were evaluated and those giving the best separation were included in modeling. Here, we chose model selection using an information-theoretical approach with multimodel inference, which is a relatively new concept in biological sciences providing a more robust alternative compared with traditional approaches of hypotheses testing (24, 25). As expected, due to the large number of possible predictive variables obtained with shotgun sequencing of the CNA pool, the data supported more than one possible model. In such cases, model averaging provides robust inference that is not conditional on a single “best” model. In addition, a random resampling strategy of 255 rounds was applied, which confirmed five of the six initial predictors for the final model\textsubscript{normRE}. From the initial model\textsubscript{normRE}, only the free left Alu monomer (FLAM-C; ref. 30) was replaced by MLT2A2, an endogenous retrovirus-related element. This suggests that the Alu monomer FLAM-C is a weaker predictor for breast ductal carcinoma than the LINE elements and AT-rich sequences. This conclusion is strengthened by the fact that MLT2A2 is included in both the model\textsubscript{RRnormRE} and the model\textsubscript{normGO}.

The final models were tested for their performance with independent data obtained from patients with stage I invasive ductal carcinomas. Interestingly, both model\textsubscript{normRE} and model\textsubscript{RRnormRE} yielded higher area under the ROC curve values in the stage I patient cohort with the latter providing the highest C value when applied to the data obtained from stage I cancers. We used the C values obtained for the validation data set as a general measure for the goodness of a model, although the obtained differences were not statistically significant, as shown by the overlapping confidence intervals. Further studies are needed to verify this trend. Nevertheless,

Table 3. Diagnostic performance of model\textsubscript{normRE}, model\textsubscript{normGO}, and model\textsubscript{RRnormRE} in different patient groups versus the normal female training set

<table>
<thead>
<tr>
<th>Model</th>
<th>Patient group</th>
<th>C value*</th>
<th>Specificity†</th>
<th>Sensitivity</th>
<th>Sensitivity‡ (Spec. 97.5)</th>
<th>Maximum accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model\textsubscript{normGO}</td>
<td>Stage II to IV</td>
<td>0.94 (0.80-0.99)</td>
<td>0.90</td>
<td>0.93</td>
<td>0.61</td>
<td>0.91</td>
</tr>
<tr>
<td></td>
<td>Stage I</td>
<td>0.90 (0.62-0.97)</td>
<td>0.94</td>
<td>0.80</td>
<td>0.50</td>
<td>0.92</td>
</tr>
<tr>
<td></td>
<td>Stage I to IV</td>
<td>0.93 (0.80-0.98)</td>
<td>0.88</td>
<td>0.89</td>
<td>0.58</td>
<td>0.89</td>
</tr>
<tr>
<td>Model\textsubscript{normRE}</td>
<td>Stage II to IV</td>
<td>0.89 (0.74-0.96)</td>
<td>0.92</td>
<td>0.79</td>
<td>0.57</td>
<td>0.88</td>
</tr>
<tr>
<td></td>
<td>Stage I</td>
<td>0.94 (0.76-0.99)</td>
<td>0.92</td>
<td>0.90</td>
<td>0.70</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>Stage I to IV</td>
<td>0.91 (0.78-0.97)</td>
<td>0.92</td>
<td>0.82</td>
<td>0.50</td>
<td>0.89</td>
</tr>
<tr>
<td>Model\textsubscript{RRnormRE}</td>
<td>Stage II to IV</td>
<td>0.90 (0.76-0.97)</td>
<td>0.82</td>
<td>0.82</td>
<td>0.57</td>
<td>0.84</td>
</tr>
<tr>
<td></td>
<td>Stage I</td>
<td>0.97 (0.89-1.00)</td>
<td>0.87</td>
<td>1.00</td>
<td>0.70</td>
<td>0.95</td>
</tr>
<tr>
<td></td>
<td>Stage I to IV</td>
<td>0.92 (0.81-0.97)</td>
<td>0.82</td>
<td>0.87</td>
<td>0.50</td>
<td>0.85</td>
</tr>
</tbody>
</table>

NOTE: Model performance was assessed by calculating the area under the ROC curve.
*C value (area under the ROC curve with 95% confidence intervals in brackets).
†Specificity and sensitivity were calculated at the point of best separation.
‡Sensitivity at a specificity of 97.5%.
changes in the chromatin order are known to be relevant and acetylation or methylation of histones leading to epigenetic modifications such as methylation of DNA some positioning in the genome of the cancerous cells. Repetitive elements that might reflect a differential nucleosome revealed differences in the relative amounts of different digestion by nucleases (19). Our sequencing approach revealed that the protein-bound DNA is protected from bodies, and that the protein-bound DNA is protected from bodies, and that the protein-bound DNA is protected from bodies, and that the protein-bound DNA is protected from.

Generally, cell-free DNA in the blood is associated with histones and it is assumed that these circulating nucleosomes originate from apoptotic or necrotic cells in the body, and that the protein-bound DNA is protected from digestion by nucleases (19). Our sequencing approach revealed differences in the relative amounts of different repetitive elements that might reflect a differential nucleosome positioning in the genome of the cancerous cells. Epigenetic modifications such as methylation of DNA and acetylation or methylation of histones leading to changes in the chromatin order are known to be relevant to cancer (33). Recently, sequence-specific histone methylation was detected in the plasma of patients with multiple myeloma (34). All models presented herein contain several LINE1 elements that were lower in patients with breast cancer than in normal controls. Alterations in other LINE1 elements in serum have already been associated with breast cancer (35). Interestingly, extensive hypomethylation of LINE elements and endogenous retroviruses have been found in several cancers (36), and hypomethylation does have an effect on nucleosome positioning (33). Furthermore, LINE elements in humans are predominantly found in chromosomal G bands also known as facultative heterochromatin (37, 38). Interestingly, there is evidence of nonrandom degradation of DNA in leukemic cells in which apoptotic DNA preferentially hybridizes to heterochromatin (39). We found AT-rich sequences being a strong positive predictor in all of the calculated models. Recently, it was reported that SATB1, the special AT-rich binding protein, is highly expressed in aggressive breast cancer cells (40). SATB1 belongs to a family of nuclear matrix binding proteins, which constitute and maintain the DNA nuclear architecture, thereby regulating cell-specific gene expression. SATB1 is normally cleaved by caspases during the early stages of apoptosis (41). An overexpression of SATB1 in breast cancer cells could, therefore, be one possible reason for the relative overrepresentation of AT-rich sequences in the breast cancer-specific circulating DNA pool. Although speculative, it seems possible that apoptotic DNA degradation in cancer cells differs from nonmalignant cells and that these differences could be detected by sequencing of apoptotic DNA remnants in the serum.

Alternatively, differential repetitive element representation in the DNA pool may originate from different processes by which DNA is released from cancer and normal cells. Several forms of cell death other than apoptosis (e.g., necrosis, autophagy, or mitotic catastrophe) as well as an active release of newly synthesized DNA have been discussed as possible sources of cancer-specific CNA (31, 42). The detection of MLT2A2 as a strong predictor in two of our models also suggests that the CNA pool might also contain retrovirus-like particles that reflect distinct retroviral activity of endogenous retroviruses in cancer cells. Retroviral particles have been detected in various malignant tumors (43) and in the T47-D human breast cancer cell line (44).

Based on recent reports, a bias introduced by random amplification methods including the WGA4 cannot be ruled out (21, 45), in which the latter was shown to introduce the least bias (45). To avoid a significant influence on the comparative data, samples were prepared strictly the same way, two independent WGA4 reactions were pooled for each sample and breast cancer samples and normal controls were run together in mixed batches to exclude bias introduced by individual sequencing runs.

In conclusion, we report specific DNA-transposable element sequences that could discriminate all stages of invasive ductal breast carcinoma with significant specificity and sensitivity. Although not practical as such for standard

FIGURE 1. Distribution of scores calculated using modelRRnormRE on data obtained from different patient groups. The 75th and 25th percentile are plotted as boxes, whiskers display the 97.5th and 2.5th percentiles. BrCa (all), all patients with breast cancer regardless of tumor staging; BrCa (validation set), 10 patients with tumors in stage I; NORM, normal group as used in training sets; MS, multiple sclerosis patients; NC, multiple sclerosis and normal data combined as noncancerous group. For a detailed description of samples, see Table 1.

modelRRnormRE is especially promising for detecting stage I invasive ductal carcinoma with high sensitivity. A previously published CNA-based breast cancer analysis, which detected differences in the integrity of circulating Alu elements, showed no clear discrimination of breast cancer in stage I disease (31). Screening tests for the early detection of cancer is universally accepted to reduce cancer mortality. Nevertheless, careful optimization is required because an inordinately high false-positive rate will cause unnecessary diagnostic work-ups (32). With a sensitivity of 70% at a specificity of 100% in independent control and patient data sets, the modelRRnormRE seems promising as a basis for the development of future screening tools. Adjustment of the specificity cutoff to 95% provides a high predicted sensitivity of 90%. The results obtained at different time points from a multiple myeloma patient that developed a second malignant neoplasm suggest that this model may not be affected by cancers in general.

The origin of serum DNA sequences is controversial. Generally, cell-free DNA in the blood is associated with histones and it is assumed that these circulating nucleosomes originate from apoptotic or necrotic cells in the body, and that the protein-bound DNA is protected from digestion by nucleases (19). Our sequencing approach revealed differences in the relative amounts of different repetitive elements that might reflect a differential nucleosome positioning in the genome of the cancerous cells. Epigenetic modifications such as methylation of DNA and acetylation or methylation of histones leading to changes in the chromatin order are known to be relevant
screening or prognostic purposes, the detected patterns provide a knowledge base for the development of a serum-based assay. Automated methods such as microarrays or quantitative multiplex PCR, once validated against the sequence data reported here, will further investigation in a prospective clinical setting. The data is promising for the development of a cost-effective, serum-based screening assay.

Disclosure of Potential Conflicts of Interest
All authors, ownership interest, Chronix Biomedical.

References


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