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Translational Database Selection and Multiplexed Sequence Capture for Up Front Filtering of Reliable Breast Cancer Biomarker Candidates

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Abstract

Biomarker identification is of utmost importance for the development of novel diagnostics and therapeutics. Here we make use of a translational database selection strategy, utilizing data from the Human Protein Atlas (HPA) on differentially expressed protein patterns in healthy and breast cancer tissues as a means to filter out potential biomarkers for underlying genetic causatives of the disease. DNA was isolated from ten breast cancer biopsies, and the protein coding and flanking non-coding genomic regions corresponding to the selected proteins were extracted in a multiplexed format from the samples using a single DNA sequence capture array. Deep sequencing revealed an even enrichment of the multiplexed samples and a great variation of genetic alterations in the tumors of the sampled individuals. Benefiting from the upstream filtering method, the final set of biomarker candidates could be completely verified through bidirectional Sanger sequencing, revealing a 40 percent false positive rate despite high read coverage. Of the variants encountered in translated regions, nine novel non-synonymous variations were identified and verified, two of which were present in more than one of the ten tumor samples.

Introduction

Discovery of biomarkers has traditionally been mediated by interpretation of transcriptome data generated using array-based expression profiling platforms [1,2]. This has for instance resulted in better understanding of prostate cancers, where the initial screening for serum PSA has led to earlier detection of the disease, but also to inaccurate diagnosis, necessitating discovery of new and better biomarkers [3,4]. It has earlier been shown that biomarker discovery benefits from integration of genomic and proteomic technologies [5]. Recent data have also demonstrated a stronger correlation between transcripts and proteins than previously anticipated [6,7]. Thus, translating aberrant protein expression patterns to transcript differences is of high interest, and the possibility that some of these changes are encoded in the genome offers new openings to identify causative mutations. Today’s novel technologies and large-scale efforts for proteomic screening are providing the grounds for a great increase of pace in such biomarker discovery. The Human Protein Atlas is one example where the proteome is being screened for differences in expression patterns in a large collection of cancers and corresponding healthy tissues [5,9]. Antibodies are created for each human protein in a mono-specific polyclonal fashion and are used to stain tissue microarrays in order to determine the expression patterns and levels.

In addition to more efficient proteomic screening, the advent of massively parallel sequencing methods [10,11,12] has greatly increased the throughput of genomic data [13] and has also provided new platforms for transcriptomic screening past the traditional array based technologies [14,15]. Whereas genomes of healthy and diseased tissues can be sequenced in full today [16,17], the sequencing throughput is still not able to provide us with enough data to screen vast numbers of genomes in parallel in a cost-efficient manner. The response to this has traditionally been the selection of genomic regions of interest by PCR [18], but has lately been replaced by methods for extraction of regions of interest by sequence enrichment strategies [19,20,21].

Here we demonstrate a model for further cost reduction and increased efficiency in biomarker discovery by employing up-front database selection in combination with sample barcoding [22] and multiplexed sequence capture enrichment, to rationally filter out potential biomarkers at an early stage.

Materials and Methods

Ethics statement

Ethical permission was obtained from the Ethics Committee at Lund University whereby informed consent was deemed not to be required other than by the opt-out method. The study was
conducted according to Declaration of Helsinki Principles. The data were analyzed anonymously.

Translational database selection
The Human Protein Atlas database [8] was searched for proteins with a clearly differential staining pattern in healthy breast tissue and breast cancer tissue requiring: i) breast glandular cells with no staining and ii) breast cancer tumor cells with at least five patients with strong staining. A complementary search for breast glandular cells with strong staining and breast cancer tumor cells with at least ten patients with weak or no staining was also carried out.

Further, the proteins found through the search results were screened to match a number of criteria of interest. These included overall differential staining of healthy and cancerous tissues for the particular protein and high assay validation scores. Proteins were scored as particularly interesting if they were present in a transmembrane region, and if they contained a signaling peptide. In total, 41 proteins were selected in this way and an additional 10 proteins known to be associated with cancer from the literature were added to the list. (Table S1)

Selection and design of regions for genomic enrichment
The coding exons for the proteins selected through the HPA database were extracted from the UCSC human reference genome (hg18) database. Additionally the 5'UTR and 3'UTR regions were included, as well as 1000 basepairs upstream from the 5'UTR. To facilitate efficient capture of the targeted genomic regions, the selected regions were expanded to a minimum of 250 basepairs and regions with a resulting overlap were fused together. 479 regions totaling 303,788 basepairs, 89,705 of which were protein coding, were selected in this way.

The selected regions were submitted to the array manufacturer (Nimblegen, Madison, WI, USA) for manufacturing of 385k-feature enrichment arrays. The final design after internal processing and filtering of repetitive regions contained 581 tiled regions spanning a total of 303,986 target bases.

Sampling of tumors and DNA extraction
Tumors were surgically removed from the patients, trimmed for healthy tissue and instantly put into a freezer at -20°C. >99% of the cells were judged to be of tumor origin. For extraction of DNA ten pieces approximately 1 mm³ each were cut out from each tumor and put into a fastprep tube (164102930, Lysing matrix D, Fisher Scientific, Gothenburg, Sweden). 360 µl of ATL buffer from the DNasey (Qiagen, Valencia, CA, USA) kit was added to the tube that was then processed 2 times 60 seconds on a Fastprep FP210 system (Qiogene, Carlsbad, CA, USA). The homogenized liquid phase was pipetted into a Qiashredder column (Qiagen, Valencia, CA, USA) that was centrifuged for 1 minute at 13k rpm. 40 µl of protease K (Qiagen, DNasey kit) was added to the shredded material followed by a 15-minute incubation of the sample at 56°C. 300 µl of buffer AL (DNasey, Qiagen) and 300 µl 96% ethanol was added to the sample and the resulting mixture was split in half and transferred into 2 DNasey Mini spin columns. The columns were processed according to manufacturers instructions (DNasey, Qiagen) after which each sample was eluted twice with 200 µl of EB buffer (DNasey, Qiagen) in separate tubes, totaling 4 tubes with 200 µl eluate each for each initial tumor sample.

Following elution each sample was ethanol precipitated by adding 20 µl 3 M NaAc and 300 µl -20°C 96% ethanol, and then incubated at -80°C for 15 minutes. After freezing the sample was centrifuged at 13k rpm for 25 minutes in room temperature. The sample was washed with 500 µl -20°C 70% ethanol and centrifuged at 13k rpm for another 15 minutes in room temperature. The liquid was removed and the sample tubes were dried at room temperature over night with open lids under a protective cover. The following day 20 µl 1xTE buffer was added to each sample tube to dissolve the extracted DNA after which the tubes for each sample were pooled and analyzed for concentration and purity on a NanoDrop N-1000 spectrophotometer (NanoDrop, Wilmington, DE, USA).

Healthy reference tissue from each patient sample was processed in the same way to obtain DNA for validation sequencing of variations indicative of potential biomarkers.

DNA enrichment and sequencing
A total of ten DNA samples from breast cancer tumors were processed into sequencing libraries using an in house developed automated protocol [23] based on the GS FLX titanium Library preparation method (Roche/454, Branford, CT, USA) using multiplex identifier handles (MID 1–10, Roche/454). The samples were then pooled in equimolar ratios into a single tube and enriched for target sequences by hybridizing to the custom Nimblegen 385k array (Roche/Nimblegen, Madison, WI, USA) previously designed and manufactured for the project. Following enrichment the pooled library was titrated and sequenced according to manufacturers instructions (Roche/454) on a GS FLX using long-read titanium chemistry.

Data analysis
The data corresponding to each sample was mapped to the human reference genome (hg19) using the Roche/454 GS Mapper software (Newbler version 2.3), and the file containing the resulting high confidence variants (HCdiffs.txt) was used for further analysis. The single nucleotide variants were annotated using custom perl scripts and the knownGene transcript database (UCSC). The variants were sorted by the number of samples they were present in. Information from dbSNP 130 was used to extract previous knowledge about specific SNPs.

Validation of variations
Following Roche/454 sequencing and data analysis as described above, 15 novel non-synonymous variations were encountered, spread across the different samples (Table S2). These were verified through amplification by PCR followed by bidirectional Sanger sequencing of the corresponding genomic regions. Nine out of these variations were confirmed as heterozygous in both tumor tissue and healthy reference tissue, whereas six of the variants were not visible in either tissue in the validation experiments, hence deemed false positives.

Results
DNA from breast cancer samples and surrounding healthy breast tissue (Figure 1) was isolated from ten surgically removed tumors and selected genes were investigated using array-based enrichment and DNA sequencing. In total, 581 genomic regions corresponding to the 51 selected differentially expressed or literature derived proteins (Table S1) were surveyed for mutations. Out of the 1,109,321 sequencing reads generated, 22.63% mapped uniquely to the human genome and overlapped with the 303,936 bases associated with the 581 target regions. Of the generated bases, 69.3 million mapped to the target, corresponding to an average 228 times sequence coverage of the target. >99.9% of the target was covered by at least one read. The distribution of reads between the different samples was good in general (Table 1)
however multiplex identifier 3 (MID 3) proved to have been very inefficiently amplified in the emPCR, as verified later by qPCR of the MIDs [23], and the sample labeled with MID 9 generated very low concentrations at the library preparation, which resulted in a lower molar amount of DNA from the MID 9 tagged sample in the pool.

In total 1,982 single nucleotide variations (SNVs) at 579 unique positions in the target regions were found in the tumor samples when compared to the human reference sequence (hg19). A higher rate of variation, per base, was seen in 5’UTR, 3’UTR, promotor and intron sequence of the target (0.07%), than in protein coding exon sequence (0.05%). Of the 149 unique positions with SNVs in protein coding sequence (488 SNVs total), 66 were subjected to an amino acid change. 15 of these non-synonymous alterations had not previously been reported in dbSNP build 130. All novel SNVs were detected in frequencies above 20% of the reads. Six of these non-previously reported variations were encountered in more than one tumor sample, and were confined to three genes, one in SATB1, four in MUC5AC and one in DDX26B (Table 2).

Confirmatory Sanger sequencing was carried out for all 15 novel SNVs in the tumor samples and normal reference tissue. This resulted in confirmation of nine of the 15 SNVs as heterozygous variations present in both tumor and normal tissue. The six remaining variations turned out to be false positives.

**Discussion**

The percentage of reads mapping to the targeted regions for the used enrichment platform has previously been reported around 60–80% [19]. There are several possible reasons for the modest fraction of sequencing reads that mapped to the targeted regions in

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mapped reads</th>
<th>Mapped bases</th>
<th>Ave coverage</th>
<th>SNVs</th>
<th>Coding SNVs</th>
</tr>
</thead>
<tbody>
<tr>
<td>MID1</td>
<td>35243</td>
<td>9.9mil</td>
<td>33</td>
<td>247</td>
<td>66</td>
</tr>
<tr>
<td>MID2</td>
<td>30264</td>
<td>8.6mil</td>
<td>28</td>
<td>215</td>
<td>50</td>
</tr>
<tr>
<td>MID3</td>
<td>2739</td>
<td>0.8mil</td>
<td>3</td>
<td>27</td>
<td>8</td>
</tr>
<tr>
<td>MID4</td>
<td>22952</td>
<td>6.4mil</td>
<td>21</td>
<td>217</td>
<td>55</td>
</tr>
<tr>
<td>MID5</td>
<td>30869</td>
<td>8.5mil</td>
<td>28</td>
<td>230</td>
<td>46</td>
</tr>
<tr>
<td>MID6</td>
<td>18608</td>
<td>5.2mil</td>
<td>17</td>
<td>221</td>
<td>63</td>
</tr>
<tr>
<td>MID7</td>
<td>20239</td>
<td>5.6mil</td>
<td>18</td>
<td>185</td>
<td>48</td>
</tr>
<tr>
<td>MID8</td>
<td>38539</td>
<td>10.7mil</td>
<td>35</td>
<td>252</td>
<td>53</td>
</tr>
<tr>
<td>MID9</td>
<td>7815</td>
<td>2.2mil</td>
<td>7</td>
<td>129</td>
<td>34</td>
</tr>
<tr>
<td>MID10</td>
<td>40046</td>
<td>11.4mil</td>
<td>38</td>
<td>259</td>
<td>65</td>
</tr>
<tr>
<td>Total</td>
<td>247314</td>
<td>69.3mil</td>
<td>228</td>
<td>1982</td>
<td>488</td>
</tr>
</tbody>
</table>

Distribution of mapped reads, bases, coverage and single nucleotide variants (SNVs) across the ten multiplex enriched breast cancer tumor samples. An even distribution across all samples except those tagged with multiplex identifier tag 3 and 9. Poor amplification characteristics of MID 3 was later verified by qPCR [21] to be the reason for poor representation of the corresponding sample among the sequence reads. The sample tagged with MID 9 was added in a lower molar amount upon pooling due to a low DNA concentration after library preparation.

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Translational Filtering of Biomarkers

Table 2. Encountered single nucleotide variants (SNVs) overview.

<table>
<thead>
<tr>
<th>Total unique positions with single nucleotide variants (SNVs)</th>
<th>579</th>
</tr>
</thead>
<tbody>
<tr>
<td>In target genes</td>
<td>467</td>
</tr>
<tr>
<td>In exons (incl UTR)</td>
<td>266</td>
</tr>
<tr>
<td>In introns</td>
<td>201</td>
</tr>
<tr>
<td>In coding sequence (CDS)</td>
<td>149</td>
</tr>
<tr>
<td>In non-coding sequence</td>
<td>430</td>
</tr>
<tr>
<td>Non-synonymous in CDS</td>
<td>66</td>
</tr>
<tr>
<td>Novel non-synonymous not in dbSNP</td>
<td>15</td>
</tr>
<tr>
<td>Verified novel non-synonymous not in dbSNP</td>
<td>9</td>
</tr>
<tr>
<td>Novel non-synonymous present in more than one patient</td>
<td>6</td>
</tr>
</tbody>
</table>

An overview of the single nucleotide variations encountered across the samples. 149 unique positions with single nucleotide variations (SNVs) were found in protein coding sequence, 15 of which gave rise to a different amino acid and were previously unreported in dbSNP (version 130). In total six of these turned out to be false positives when verified by bidirectional Sanger sequencing. Two of the remaining mutations were present in more than one individual and were located in the SATB1 and DDX26 genes. doi:10.1371/journal.pone.0020794.t002

Among the novel non-synonymous SNVs encountered 40% turned out to be false positives. Two of the remaining previously unreported non-synonymous SNVs were found in the genomes of more than one individual. Each was verified as a heterozygous SNV present in both normal and cancerous tissue. Their presence in SATB1, previously linked to breast cancer promotion [26], and DDX26B, to our knowledge previously unreferenced in relation to breast cancer, provides further support for continued examination of the role of the genes in conjunction to breast cancer.

Lastly, the higher frequency of variations seen in non-coding sequence as compared to coding sequence has previously been reported in healthy [18] and cancerous [17] tissue, strengthening the scientific grounds for the logical reasoning that non-protein-coding regions of the genome are less subjected to evolutionary constraints.

In summary, the employment of a translational database selection strategy in combination with multiplexed enrichment by sequence capture provides a tool for careful biomarker discovery. Given the abundance of false positives generated by massive sequencing approaches, employing a rational selection strategy prior to sequencing can provide an efficient means to limit the number of variants at the end of the pipeline, enabling a complete variant verification process and a more reliable final list of biomarker candidates.

Supporting Information

Table S1 Selected genes. 41 proteins and their corresponding genes were selected through the HPA database and 10 more proteins and their corresponding genes known to be associated with cancer from literature were added to the list. (DOC)

Table S2 Novel non-synonymous SNVs. Among the novel non-synonymous SNVs encountered 40% turned out to be false positives. Two of the remaining SNVs were present in more than one individual. (DOC)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: PLS SH JL. Performed the experiments: PLS HM. Analyzed the data: PLS MKB. Contributed reagents/materials/analysis tools: MKB KJ SH JL. Wrote the paper: PLS MKB JL.

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