Assessing unmodified 70-mer oligonucleotide probe performance on glass-slide microarrays

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Abstract

Background: Long oligonucleotide microarrays are potentially more cost- and management-efficient than cDNA microarrays, but there is little information on the relative performance of these two probe types. The feasibility of using unmodified oligonucleotides to accurately measure changes in gene expression is also unclear.

Results: Unmodified sense and antisense 70-mer oligonucleotides representing 75 known rat genes and 10 Arabidopsis control genes were synthesized, printed and UV cross-linked onto glass slides. Printed alongside were PCR-amplified cDNA clones corresponding to the same genes, enabling us to compare the two probe types simultaneously. Our study was designed to evaluate the mRNA profiles of heart and brain, along with Arabidopsis cRNA spiked into the labeling reaction at different relative copy number. Hybridization signal intensity did not correlate with probe type but depended on the extent of UV irradiation. To determine the effect of oligonucleotide concentration on hybridization signal, 70-mers were serially diluted. No significant change in gene-expression ratio or loss in hybridization signal was detected, even at the lowest concentration tested (6.25 nM). In many instances, signal intensity actually increased with decreasing concentration. The correlation coefficient between oligonucleotide and cDNA probes for identifying differentially expressed genes was 0.80, with an average coefficient of variation of 13.4%. Approximately 8% of the genes showed discordant results with the two probe types, and in each case the cDNA results were more accurate, as determined by real-time PCR.

Conclusions: Microarrays of UV cross-linked unmodified oligonucleotides provided sensitive and specific measurements for most of the genes studied.

Background

The advent of microarray technology has enabled scientists to investigate biological questions in a more global fashion. Instead of studying genes individually, the expression of thousands of genes can be analyzed simultaneously using probes attached to the surface of a microscope slide [1-6].
The cDNA microarray represents a popular array type in which double-stranded PCR products amplified from expressed sequence tag (EST) clones are spotted onto glass slides [7,8], allowing gene-expression profiles to be determined with high reproducibility and efficiency. However, construction of cDNA microarrays presents a number of challenges, largely related to costs associated with clone validation, tracking and maintenance. The laborious and problematic tracking of cDNA clones and PCR amplicons may lead to 10-30% misidentification of clones [6]. For all practical purposes, sequence verification of array elements is an ongoing necessity. Other limitations of cDNA microarrays are their difficulty, because of cross-hybridization, in discriminating expression patterns of homologous genes, alternative splice variants and antisense RNAs.

Alternatively, microarrays can be composed of short oligonucleotides (25 bases) synthesized directly onto a solid matrix using photolithographic technology (Affymetrix) [2,9] or constructed from long oligonucleotides (55-70 bases) spotted onto glass slides [10-12]. To mimic the Affymetrix design of freely moving probes tethered at one end onto a solid support, in-house manufactured or commercially available long oligonucleotides are modified by the addition of a 5′ amino group for covalent attachment onto pre-activated glass slides [5,10]. This oligonucleotide design strategy has been widely viewed as a prerequisite for accurate gene-expression measurements. However, there is no clear evidence that other covalent attachments do not form. With oligonucleotide arrays, problems related to clone tracking, handling of glycerol stocks and failed PCR amplifications are avoided. The completion of numerous microbial, plant and eukaryotic genomes, as well as extensive EST data, provides sufficient sequence information to design unique oligonucleotides capable of distinguishing homologous genes and alternative splice variants. As such, oligonucleotide probes have an added flexibility over PCR amplicons.

Comprehensive studies comparing the Affymetrix approach with cDNA arrays have only recently appeared in the literature [13,14]. Studies comparing long oligonucleotides to cDNA arrays have not been as forthcoming. In the only two study [13], a second study found a correlation of r = 0.328 between matched results from the same two platforms [14]. Discordant results were not resolved in the latter study. Here we test the performance of unmodified 70-mers printed alongside PCR amplicons. Using this unique study design, both probe types can be simultaneously interrogated with a complex target composed of both cellular and synthetic RNA.

Results
Optimal attachment parameters for 70-mers and PCR amplicons on the same slide
The success of microarray assays requires stable binding and retention of probes throughout the entire printing/blocking/hybridization/washing process. Oligonucleotides were spotted alongside PCR amplicons onto TeleChem SuperAmine aminated slides, and immobilized by ultraviolet (UV) cross-linking. To determine the optimal UV cross-linking energy required for efficient oligonucleotide immobilization, a series of spotted arrays from the same printing session were subjected to increasing UV energy (70, 150, 250, and 450 mJ/cm²). Deposition and retention of the probe onto aminated slides were assessed by: staining with Vistra Green solution and subsequent fluorescence scanning at 532 nm; and hybridization with Cy-labeled targets derived from rat brain and heart RNA. Using these two methods, optimal retention of both oligonucleotide and PCR amplicon probes was determined to occur between 250 and 450 mJ/cm² as described below.

Directly following probe deposition, UV cross-linking at 250 mJ/cm² and Vistra Green staining, the measured fluorescence intensity from oligonucleotide probes was typically higher than PCR amplicons (Figure 1a). The efficiency of the oligonucleotide immobilization strategy was tested by stringently washing the same slide overnight in 0.2% SDS (at 42°C, SSC absent in wash solution) to remove Vistra Green and any loosely bound nucleic acids, restained with Vistra Green and scanned (Figure 1b). There was essentially no change in the average median intensity values for either the oligonucleotide or PCR amplicon probes after incubation with detergent (Figure 1c).

These results show that immobilization of unmodified 70-mer oligonucleotides to SuperAmine aminated slides by high-UV cross-linking energy is sufficient and comparable to PCR amplicons. Clearly, our oligonucleotide immobilization protocol should be sufficient to sustain routine microarray hybridization and wash procedures, which are much less
The importance of titrating UV cross-linking energy for oligonucleotide immobilization is exemplified by a series of spotted arrays hybridized with Cy-labeled targets derived from rat brain and heart RNA. As seen in Figure 2, there was an increase in the appearance of hybridized spots and signal intensity as the energy of cross-linking was increased. At lower cross-linking energies (for example, 70 and 150 mJ/cm²) it is apparent that oligonucleotide probes were not sufficiently attached to the surface of aminated slides. In contrast, PCR amplicons are sufficiently immobilized onto the aminated surface of glass slides at these intensities [15]. On the basis of the hybridization experiments and in agreement with Vistra Green staining results, optimal attachment of 70-mer oligonucleotides occurred between 250 and 450 mJ/cm². An improvement in oligonucleotide retention onto aminated slides was not seen at intensities higher than 450 mJ/cm² (data not shown). Of interest was the finding that different slide chemistries (that is, poly-L-lysine, aldehyde, aminosilane, epoxide) had different UV cross-linking titration curves for optimal attachment of 70-mer oligonucleotides, and increasing the cross-linking intensity in some slide types actually decreased apparent probe deposition (data not shown). Even more surprising was the observation that slides with the same or similar slide chemistry from different vendors exhibited marked differences in the optimal UV cross-linking energy for probe attachment.

**Sensitivity of unmodified 70-mers on aminated slides**

Defining assay sensitivity is important as the ability to measure gene-expression changes is desired not only for moderate and abundant mRNAs but also for rare transcripts. One method for assessing the sensitivity of microarray assays is to use exogenous spiking controls. These controls also help to identify systematic problems associated with target labeling, slide hybridization and scanning. For this purpose, we developed a set of 10 Arabidopsis control cDNA plasmids. Each of the 10 plasmids was used to synthesize cRNA in vitro. cRNAs were quantitated and differentially spiked into heart and brain RNA samples at specific copy numbers based on the following assumptions: 360,000 mRNA transcripts per cell, 20 pg total RNA per cell and 1 pg mRNA transcript per cell; hence, 100 ‘spike’ copies/cell...
would be equivalent to about 0.14 ng ‘spike’ transcript in 10 μg total RNA [12]. To normalize our array data, we spiked five of the ten control cRNAs into the Cy3 and Cy5 labeling reactions at equal copy number, ranging from 40 to 100 copies per cell. The remaining five control cRNAs were spiked into the labeling reactions at concentrations ranging from 1 to 300 copies per cell. We printed Arabidopsis 70-mers and PCR amplicons across six different sectors in order to measure intra-slide variability. Inter-slide variability was evaluated across independent hybridizations and target labeling. Figures 3a and b compare the intra- and inter-slide variation of oligonucleotides to discern twofold changes in Arabidopsis targets. Similar results were obtained for the detection of targets spiked at threefold ratios (data not shown). These data show that our oligonucleotide array platform was able to detect two- and threefold changes in transcript number at a sensitivity of one to two copies per cell. Comparing the sensitivity of oligonucleotide probes to PCR amplicons indicates that the two probe types perform equally well at consistently detecting twofold changes in rare transcript levels (Figure 3c).

Arabidopsis probe elements serve as excellent negative controls when exogenous cRNA is not added to the labeling reaction. In the absence of cRNA spiking, cross-hybridization of Cy-labeled rat targets to Arabidopsis probe elements was negligible (data not shown).

Concordance between probe types when measuring differential gene expression in biological samples

To compare the accuracy of unmodified oligonucleotide arrays to conventional cDNA arrays, we hybridized slides containing both probe types with equal amounts of Cy-labeled target derived from rat heart and brain RNA. Hybridization to 36 out of 38 antisense-strand oligonucleotide probes was negligible to nonexistent under our incubation and wash protocols (data not shown). A positive hybridization signal (threefold above background) was obtained for 65 of the 75 sense-strand oligonucleotide probes interrogated with labeled target. Accordingly, each of the corresponding 65 PCR amplicon probes had a positive hybridization signal (threefold above background). As the intra-slide variation was low (average standard deviation (SD) was around 0.07 and around 0.11 for log₂-transformed ratio values derived from PCR amplicon and 70-mer spots, respectively), ratio values from replicate spots within a slide were averaged. Averaged ratios derived from independent experiments were plotted and regression analysis was performed to assess reproducibility of arrays containing oligonucleotide targets (Figure 4a). A similar comparison was performed on PCR amplicon probes (Figure 4b). Both oligonucleotide and PCR amplicon probes showed high reproducibility between replicate experiments (that is, different slides hybridized with targets generated from different batches of RNA samples) with correlation coefficients (r) of 0.95 and 0.96, respectively. Next, we compared oligonucleotide-derived ratios with those obtained from PCR amplicon probes (Figure 4c). A correlation coefficient of r = 0.80 (p < 0.05) and a slope close to unity were obtained, indicating that unmodified oligonucleotide and PCR amplicon probes gave comparable expression ratios. Moreover, there was agreement in the calculated average coefficient of variation (13.4%) for the expression ratios computed from the two probe types.

Validation of microarray results with real-time PCR

Of the 65 represented genes that had a positive hybridization signal with both the oligonucleotide and PCR amplicon probe types, 60 were in agreement of each other (Figure 4c). Real-time PCR was used to test the accuracy of our microarray results (Figure 5a). Six genes exhibiting a range of expression differences in heart and brain were selected for validation. These included histone H4, which did not exhibit differential expression in the two tissues (log₂ ratio around 0); cytochrome oxidase IV, kynurenine 3-hydroxylase, serine/threonine protein kinase and 14-3-3 protein gamma which exhibited two- to threefold differences (log₂ ratio = 1 to 1.6); and desmin which was around 32-fold differentially expressed between the two tissues (log₂ ratio approximately 5). In each case, the expression ratios derived from oligonucleotide and PCR amplicon probes were in accord with real-time PCR results.

There were five notable discrepancies between the two probe types, as compared to the 60 that were in agreement. A discrepancy was defined as a change equal to or greater than twofold measured with one probe type and no change (or a change in the opposite direction) measured with the other probe type. The resolution of the discordant results for inositol-1,4,5-trisphosphate receptor, H+-ATPase, branched aminotransferase and epoxide hydrolase is presented in Figure 5b. In each case, real-time PCR results were in agreement with the PCR amplicon-derived expression ratios. Interestingly, each of the oligonucleotide-derived expression ratios erroneously suggested that these genes were not differentially expressed in heart and brain tissues.

Effect of oligonucleotide probe concentration on signal intensity

In our initial experiments, unmodified 70-mer oligonucleotides were printed onto TeleChem slides at a relatively high concentration of 50 μM. By comparison, the concentration for printing 5’-amino linker modified 50-mer oligonucleotides was 20 μM [12]. To test the performance of unmodified oligonucleotides at lower printing concentrations, seven rat 70-mer oligonucleotides were serially diluted from 50, 25, 12.5 to 6.25 μM. Each oligonucleotide was chosen on the basis of earlier microarray results showing that both oligonucleotide and cDNA probes could hybridize to heart and brain targets, and that hybridization intensities associated with the seven different gene elements varied by at least an order in magnitude. The selected probes included...
SCG10 and desmin, which were highly differentially expressed in brain and heart, respectively; 14-3-3-gamma and profilin which were expressed at around twofold higher levels in brain and heart, respectively; and histone H4, 14-3-3-theta and thymosin beta-4, which showed no difference in expression between brain and heart. All diluted oligonucleotides along with their corresponding undiluted PCR amplicons (approximately 100-200 nM) were spotted onto the array at least four times. A stained representative section of an array printed with different starting concentrations of oligonucleotides and a single concentration of the corresponding PCR amplicons is depicted in Figure 6. As the starting oligonucleotide concentration was decreased from 50 μM to 6.25 μM (eightfold dilution), DNA fluorescence decreased on average twofold for the oligonucleotides (data not shown). This suggests that the capacity of the slides to
 retain 70-mer oligonucleotides in a typical 100 μm diameter spot approached saturation at the higher concentrations.

Arrays containing diluted oligonucleotides were hybridized with labeled targets derived from brain and heart RNA as before. The median Cy3 and Cy5 hybridization intensities (minus background) were summed for each oligonucleotide concentration along with their corresponding cDNA probe (Figure 7a). A comparison of oligonucleotide and cDNA probes clearly demonstrates that the longer probe length of the latter does not necessarily translate to greater hybridization signal intensities. While the PCR amplicon probes for 14-3-3 protein theta, desmin and thymosin beta-4 generated higher signals than the corresponding oligonucleotide elements, the converse was observed for 14-3-3 protein gamma, histone H4 and profilin. Moreover, there was no correlation (r = 0.06, p > 0.05) between the hybridization signal intensities acquired from PCR amplicon probes and the corresponding oligonucleotide probes. Of interest was the apparent inverse correlation between oligonucleotide concentration and hybridization intensity. Hybridization intensities actually increased with decreasing

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**Figure 4**

Scatter plots of ratios from replicate experiments for both 70-mer and PCR amplicon probes. (a) Plot of log₂ ratios detected by oligonucleotide probes from two independent experiments. The plot shows a high correlation between experiments with a correlation coefficient (r) of 0.95 and a slope of 0.89. Results are representative of five similar comparisons. (b) Plot of log₂ ratios detected by PCR amplicons from two independent experiments. The plot shows a high correlation between experiments, with r = 0.96 and a slope of 0.84. Results are representative of five similar comparisons. (c) Correlation of gene-expression ratios detected by 70-mer and PCR amplicon probes. A high r of 0.80 and a slope of 0.73 were obtained when log₂ ratio values from oligonucleotide probes were plotted against PCR amplicons. Results are representative of five similar comparisons.
oligonucleotide concentration for 14-3-3 protein gamma, desmin, SCG10, and to a lesser extent thymosin beta-4 (Figure 7a).

For the oligonucleotides corresponding to differentially expressed genes (for example, SCG10, desmin, 14-3-3-gamma, profilin), the log₂ ratios from four independent hybridizations (including flip dye experiments) were averaged and plotted in Figure 7b. The calculated ratios were highly reproducible and similar across the entire concentration range tested. This suggests that an oligonucleotide concentration as low as 6.25 µM is sufficient for accurate determination of relative expression differences. As the absolute levels of these four transcripts in rat heart and brain are not known with certainty, we repeated these experiments with known concentrations of synthetic Arabidopsis cRNA that were differentially spiked into rat heart and brain RNA. Six Arabidopsis oligonucleotides were accordingly diluted and printed onto aminated slides to test their ability to discriminate twofold differences in synthetic cRNA concentrations ranging from 10 to 300 copies per cell. Our data clearly show that an Arabidopsis oligonucleotide probe concentration as low as 6.25 µM was sufficient to accurately determine twofold differences in cRNA species at a ratio of 20/10 copies per cell (Figure 8).

Discussion

In the study reported here, we systematically compared the performance of unmodified 70-mer oligonucleotides to traditional PCR amplicons, both probe types printed and UV cross-linked onto glass slides coated with primary amine groups. Direct comparisons are best accomplished when both probes are printed alongside each other, allowing for simultaneous interrogation with a complex target. Hence, analysis is not confounded by uneven aminosilane coating in different batches of slides, inconsistencies in the array resulting from different print sessions, differences in day-to-day label incorporation, or variations in day-to-day
hybridization and wash procedures. A correlation coefficient ($r$) of 0.80 was obtained from our analysis, indicating that the two probe types gave comparable expression ratios. One variable that was not controlled for in our study was the number of cross-links per DNA molecule. Given a constant UV exposure, many more cross-links per molecule of cDNA probe are presumably formed compared to the shorter oligonucleotide probe. It is possible that the correlation coefficient was not higher as a result of the differential reaction of the two probe types to UV irradiation.

We designed our arrays to contain 75 different probes corresponding to mammalian signal transduction genes with a wide range of expression levels. In heart versus brain comparisons, oligonucleotide probes, like their cDNA probe counterparts, could reproducibly discern differences in mRNA populations as low as twofold (namely, 14-3-3 protein gamma) and as high as around 90-fold (namely, creatine kinase). Hence, the dynamic range of unmodified oligonucleotides is at least two orders of magnitude in fold-change measurements.

Figure 7
The role of oligonucleotide concentration on microarray hybridization measurements. (a) Comparison of hybridization intensities of seven genes detected by oligonucleotides printed at different concentrations. Corresponding PCR amplicons at 100-200 ng/µl are included for comparison. The medium background subtracted fluorescent intensities were summed for the Cy3 and Cy5 channels for each gene and plotted. (b) Comparison of the expression ratios of differentially expressed genes detected by oligonucleotides printed at different concentrations. Data are the mean ± SD of four independent experiments.
In the course of our work, we generated a resource of 10 Arabidopsis spiking control cRNAs along with their corresponding 70-mer oligonucleotide and PCR amplicon probes. As part of our quality-control procedures, all microarray assays routinely incorporate the spiking controls. These reagents will allow the microarray user to add specific concentrations of known transcripts into a complex mix of mammalian target RNA in order to assess, for example, hybridization kinetics, intra-slide variability, inter-slide variability, sensitivity and effectiveness of normalization algorithms. On the basis of experiments with the spiking controls, unmodified oligonucleotides can be used to detect twofold changes in transcript number at a level of 2-20 mRNA copies per cell. It is important to note that our protocol for generating first-strand cDNA target involves the use of random primers. At the outset, the Arabidopsis cRNAs were engineered to contain a 3’ poly(A) tail. Hence, alternative protocols using oligo(dT) to prime mRNA for the synthesis of labeled target [15,16] can still take advantage of our spiking control set.

In our initial assessment of cDNA and 70-mer oligonucleotide probe types, the latter was printed at a concentration of 50 μM. Even at a printing concentration as low as 6 μM, oligonucleotide probes were capable of discerning twofold expression differences in complex cellular RNA mixtures and in synthetic spiked cRNAs. In fact, decreasing the oligonucleotide printing concentration from 50 to 6 μM had the effect of increasing the hybridization signal around two- to sixfold for a number of the probes (Figure 7a). The reason is unclear, but it is possible that high-density packing of an oligonucleotide probe within the confines of a small spot interferes with fluorescence emission of the target or hybridization efficiency. Alternatively, the higher spotting concentrations may favor cross-linking of the oligonucleotide probes to each other following UV irradiation. In either case, this phenomenon appears to be sequence dependent as not all probes exhibited this behavior. The present study also demonstrates that longer probes are not necessarily associated with higher hybridization signals, as the hybridization signals from half of the 70-mer oligonucleotide probes were actually higher than or equivalent to their corresponding PCR amplicons, which have an average length of 1 kilobase (kb). Taken together, the combination of unmodified oligonucleotides and low printing concentrations has resulted in an approximately 16-fold reduction in reagent costs. An issue not evaluated in the present study, but one that has significant cost-saving potential, is the effect of reducing the length of unmodified oligonucleotides on microarray sensitivity. Clearly, this is an area for future investigation.

Of the five discordant results found between oligonucleotide and cDNA arrays, real-time PCR data validated the accuracy of the cDNA probe type in every case (Figure 5b). It seems likely that a failure in oligonucleotide probe design was responsible for the discordant data. Analysis of the discordant oligonucleotide sequences (that is, inositol-1,4,5-trisphosphate receptor, H+-ATPase, branched aminotransferase and epoxide hydrolase) did not reveal any obvious secondary structure that might interfere with hybridization. Treatment of spotted arrays with UV light is thought to induce free-radical-based coupling between thymidine residues on the oligonucleotide and carbon atoms on the alkyl amine groups of coated glass slides (Todd Martinsky, TeleChem International, personal communication). The T content of discordant and discordant oligonucleotides was similar, with average values of 26% and 29%, respectively, suggesting that UV cross-linking was not preferentially disrupting hybridization specificity of discordant 70-mers. Moreover, there was a lack of correlation (r = 0.03, p > 0.05) between T content of the oligonucleotides and corresponding hybridization signal intensities. Of interest, however, was the finding that the discordant oligonucleotides had an average GC content of 57% compared to the concordant oligonucleotide average of 50%. Accordingly, the hybridization signal associated with the discordant oligonucleotides was around two- to threefold higher than the concordant oligonucleotides, suggesting that ‘non-specific’ Cy-labeled targets were cross-hybridizing with the discordant oligonucleotides. This possibility is clearly illustrated for the H+-ATPase gene (Figure 9). Within the H+-ATPase 70-mer sequence is a stretch of 20 contiguous nucleotides perfectly matching a
region in the tumor endothelial marker 8 mRNA. It has been shown previously that 15 contiguous nucleotides are sufficient for cross-hybridization of non-target species [12]. There are two important points to note. First, the PCR amplicon for H+-ATPase also contains the same 20 contiguous nucleotides (Figure 9). Regardless of this, this particular probe was still able to distinguish differential expression of the H+-ATPase gene in heart and brain tissue. We postulate that a large fraction of H+-ATPase-specific Cy-labeled targets (which on average should be 100-200 nucleotides long) were available for hybridization to complementary sequences found on the longer PCR amplicon probe but absent on the shorter 70-mer probe (for example, sequences downstream of the 70-mer). Second, tumor endothelial marker 8 mRNA was identified in mouse. The orthologous rat mRNA has not been cloned yet, which is reflected by the more than 2,5 million mouse EST sequences present in dbEST, compared to only 351,827 ESTs for the rat. On the basis of BLAST searches of human and mouse sequences, contiguous non-target sequences could also be identified in the discordant oligonucleotides for inositol-1,4,5-trisphosphate receptor, branched aminotransferase and epoxide hydrolase. Hence, future oligonucleotide design considerations should include an analysis of mouse and human sequences, because of the relatively small number of available rat sequence for expressed transcripts. In addition, the synthesis of redundant probe sets (for example, two 70-mers per gene) might be warranted to help decrease false negatives by an order of magnitude.

The mechanism of the adherence of unmodified oligonucleotides to glass slides has been addressed [11]. Attachment involves noncovalent interactions such as electrostatic interactions, where the negatively charged phosphate backbone of the oligonucleotide is attracted to the positively charged surface of the glass slide (for example, a surface containing protonated alkyl amines). Whereas noncovalent interactions appear to be the predominant mechanism for oligonucleotide attachment, covalent linkage is likely to have an important supplementary role in UV-irradiated microarrays. This seems plausible as our stringent overnight washes in strong detergent did not appreciably detach unmodified oligonucleotides from the slide surface. The importance of UV cross-linking cannot be overemphasized. Under-irradiation of cDNA arrays is known to cause insufficient binding of DNA and over-irradiation results in over-nicking of DNA samples [17]. A further complicating factor is our finding that oligonucleotides printed onto different slide chemistries (or slides with similar chemistries from different vendors) will have very different optimal UV titration curves. In our hands, optimal UV cross-linking occurred at 450 and 70 mJ/cm² for oligonucleotides printed onto TeleChem SuperAmine™ and Corning GAP I™ slides, respectively. For TeleChem slides, under-irradiation (70-150 mJ/cm²) causes insufficient oligonucleotide attachment. For Corning slides, over-irradiation (150-450 mJ/cm²) results in a decrease in the hybridization signal that may reflect excessive covalent attachment of oligonucleotides. As UV cross-linking may adversely affect oligonucleotide accessibility to
labeled target during hybridization, we cannot discount the possibility that alternative attachment strategies (for example, 5'-amino-modified oligonucleotides) may provide greater sensitivity and specificity. This issue needs to be explored in the future.

In the present study, we provide evidence that the performance of unmodified 70-mer oligonucleotides is comparable to cDNAs printed on glass slides. Optimal conditions were identified for oligonucleotide attachment and hybridization/wash conditions, resulting in high assay sensitivity and reproducibility. Our results show that unmodified oligonucleotides can provide an accurate, reproducible and cost-effective means to measure gene-expression profiles. Of interest is the fact that our hybridizations were successfully carried out on slides that simultaneously contained both PCR amplicons and oligonucleotides. Hence, future microarrays can be constructed in a modular fashion, with oligonucleotide-based elements being added to existing PCR amplicons as more genomic sequence information is gathered, in the absence of readily available cDNA clones. Lastly, our findings have broader implications, suggesting that the combination of expression measurements across different platforms (for example, Affymetrix and cDNA arrays, unmodified long oligonucleotides and cDNA arrays) within a single analysis maybe feasible [18].

Materials and methods

Constructing exogenous spiking cRNA controls and a PCR ampliton printing set to assess oligonucleotide sensitivity

Ten Arabidopsis thaliana genes corresponding to chlorophyll a/b-binding protein (Cab), lipid transfer protein 4 (Ltp4), lipid transfer protein 6 (Ltp6), NAC1, ribulose-5-phosphate kinase (PRKase), ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (rbcL), rubisco activase (Ra), root cap 1 (RCP1), triosphosphate isomerase (TIM), and papain-type cysteine endopeptidase (XCP2) were designed to detect tissue-specific transcripts from either rice (Oryza sativa) or potato (Solanum tuberosum). The corresponding 500 bp of each gene in the Arabidopsis cDNA library were amplified by PCR with the SP6 (5'-ATTAGTGACAC-TATAG-3') and M13R primers. To generate cRNAs containing a 3' poly(A) tail, pSP64poly(A) constructs were linearized with EcoRI and in vitro transcribed from the SP6 promoter using the MEGAscript™ High Yield Transcription Kit (Ambion, Austin, TX). The Arabidopsis cRNA set was designed to serve as spiking controls to assess a broad range of copy numbers (rare, moderate, abundant) and varying expression ratios (1:3, 1:2, 1:1, 2:1, 3:1). For printing control 70-mer, oligonucleotides were synthesized on the basis of the corresponding 500 bp of each gene in the Arabidopsis control spiking cRNA set (Table 1). Development of our control set of 10 Arabidopsis oligonucleotides and corresponding set of PCR amplicons subcloned into pSP64poly(A) serve as a valuable quality-control resource for cDNA/oligo microarrays. The Arabidopsis control spiking cRNA vector set and protocols will be made freely available to academic investigators upon request.

Rat 70-mer oligonucleotide design

To minimize cross-hybridization, oligonucleotides of 70 bases (unmodified) were designed using the computer program, Pick70 [19]. Oligonucleotide design considerations included uniqueness, avoidance of internal self-annealing structures, narrow T_m range (75-80°C) over the entire oligonucleotide set and masking of low-complexity regions. The TIGR Rat Gene Index containing a non-redundant set of expressed mRNA sequences [20] was used as the ‘complete genome source’ for selecting 70-mer oligonucleotide sequences with Pick70. Oligonucleotides were chosen to represent ‘housekeeping’ and signal transduction genes, while other 70-mers were designed to detect tissue-specific transcripts from either brain or heart (for example, those for SCG10, creatine kinase, and desmin). The sequences of the 70-mers and corresponding GenBank accession numbers of the genes are available as an additional data file. Oligonucleotides were synthesized at a 50 nmol scale by Invitrogen/Life Technologies (Cloud, CA) or Operon Technologies (Alameda, CA), and resuspended in sterile millQ water to a final concentration of 100 μM. We selected individual cDNA clones from the TIGR Rat Gene Index whose EST sequences corresponded to the same gene from which the 70-mers were designed. Rat cDNA clone inserts were amplified by PCR with M13F (5'-GGTTTCCAGTCAGAGTGTTG-3') and M13R (5'-TGAGCGATAACAGTTCACAC-3') primers [15,16]. Insert size ranged from 0.5 to 1.5 kb.
**Microarray fabrication**

Oligonucleotides (50 μM, except where indicated otherwise) and PCR amplicons (100-200 nM) in 50% DMSO were printed onto SuperAmine slides (TeleChem International, Sunnyvale, CA) using an Intelligent Automation Systems (IAS) arrayer (Cambridge, MA) with a 12-pen print head [16]. The rat 70-mers and PCR amplicons were printed in quadruplicate while the 10 Arabidopsis 70-mers and PCR amplicons were spotted into six different sectors on the slide. After printing, DNA was cross-linked to the slides by UV irradiation with a Stratalinker UV Crosslinker (Stratagene, La Jolla, CA) and stored in a vacuum chamber until use. To assess oligonucleotide retention, slides were UV cross-linked and stained for 10 min in Vistra Green Nucleic Acid staining solution (Amersham Pharmacia, Piscataway, NJ) at a 1:10,000 dilution. Afterwards, slides were washed at least five times, 1 min each, in milliQ water at room temperature, centrifuged to dryness (500 rpm x 5 min), and scanned at 535 nm using a dual laser GenePix 4000B scanner (Axon Instruments, Foster City, CA). Subsequently, slides were gently agitated in 0.2% SDS (no salt) overnight at 42°C, washed extensively with water, scanned to ensure that the dye was completely removed, retained with Vistra Green and rescanned.

**Target labeling and array hybridization**

To generate labeled single-stranded cDNA target, 10 μg total RNA from rat heart or brain (Clontech, Palo Alto, CA) was reverse transcribed for 2-3 h at 42°C in the presence of 6 μg random primers (Invitrogen/Life Technologies), 1x first-strand synthesis buffer (Invitrogen/Life Technologies), 10 mM DTT, dNTP mix (25 mM dATP, 25 mM dCTP, 25 mM dGTP, 15 mM dTTP, 10 mM amino allyl-dUTP), and 200 units SuperScript II reverse transcriptase (Invitrogen/Life Technologies). RNA was hydrolyzed with 200 mM NaOH and 100 mM EDTA for 15 min at 65°C, then neutralized with 200 mM HCl. First-strand cDNA was purified from unincorporated amino allyl-dUTPs on QIAquick PCR purification columns (Qiagen, Valencia, CA) according to manufacturer’s instructions, except that QIAquick wash buffer was replaced with 5 mM K+ phosphate buffer (pH 8.5) containing 80% ethanol, and cDNA was eluted with 4 mM K+ phosphate buffer (pH 8.5). Eluted cDNA was lyophilized, resuspended in 4.5 μl 0.1 M Na2CO3 buffer (pH 9), mixed with either Cy3 or Cy5 NHS-ester (Amersham Pharmacia), and incubated for 1 h in the dark at room temperature. Cy3- and Cy5-labeled cDNA targets were then purified on QIAquick PCR purification columns, combined and concentrated by lyophilization, and hybridized to the microarray at 42°C for

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**Table 1**

**Arabidopsis 70-mer oligonucleotide probes**

| Abbreviation | Complete gene name | GenBank accession number | 70-mer sequence |
|--------------|--------------------|--------------------------|-----------------|
| RCP1         | Root cap 1 protein | AF168390                 | 5’-AACAGATCTTTGCTGGGACAACCTTCTTCTGATTGCAGCTATAATCATGGCTCTGAACTGGGA-3’ |
| Cab          | Photosystem I chlorophyll | X56062 | 5’-TGCTCGCTTCTCTGATCTCGAGAAAGCATTAGGATATGGAAACTGGGTTTGACAGGATCGCCATCGGAG-3’ |
| TIM          | Triosephosphate isomerase | AF247559 | 5’-GGACCACTCAAGAACCTTGGCATCATTCTTCTGGCCACAAAGGACCTAGCTGCTGAC-3’ |
| Ltp4         | Lipid transfer protein 4 | AF159801 | 5’-AGTCCCATCTTCTAAGCCAGAATGACCTTCTGCAGAAGCATTAGGATATGGAAACTGGGTTTGACAGGATCGCCATCGGAG-3’ |
| PRKase       | Ribulose-5-phosphate kinase | X58149 | 5’-AAAAACAAGGAGATTTCAACACACTCATCTTCTGCAACAAAGGACCTAGCTGCTGAC-3’ |
| Ltp6         | Lipid transfer protein 6 | AF159803 | 5’-GCCGTTGCTTGTCTCGCTCTTGCTCTACACCGGAGCAGCAGCTGCTGAC-3’ |
| rbcL         | Ribulose-1,5-bisphosphate carboxylase/oxygenase, large subunit | ATU91966 | 5’-AGTCCGTTCTTCTACTCTGACTCATGAGAACACCTCTCCGAGATTTGGAATCTCATGACAAACCTCATAG-3’ |
| Ra           | Rubisco activase | X14212 | 5’-AGCCTGCTGCAGGCTGTTATGTTGATACTCAGAATCAGTCAACACAGATGTTAACCCACACAG-3’ |
| XCP2         | Papain-type cysteine endopeptidase | AF191028 | 5’-GCTTTCTTCCAGCCTTACCTCATTCCGTGATATCCTCCCGAGGATTTGGAATCTCATGACAAACCTCATAG-3’ |
| NAC1         | NAM-like protein | AF198054 | 5’-GGCTGCTGACATGCGTCTATCCACGTGATGAGATCCTTACATCAACTTTCCGAGGATTTGGAATCTCATGACAAACCTCATAG-3’ |
16 h in hybridization solution containing 50% formamide, 5x SSC, 0.1% SDS, 20 μg mouse Cot-1 DNA and 10 μg poly(dA). Reverse dye labeling of samples was employed in separate experiments to account for any bias in dye coupling or emission efficiency of Cy dyes. After hybridization, microarray slides were washed by immersion into 2x SSC, 0.2% SDS for 5 min at 42°C, 0.2x SSC, 0.1% SDS for 1 min at room temperature, 0.2x SSC for 1 min at room temperature, and 0.05x SSC twice for 1 min at room temperature, dried by centrifugation, and immediately scanned. Different hybridization and wash conditions were tested (data not shown). The procedures described above have been optimized for both PCR amplicon and oligonucleotide probes regardless of whether the two probe types are printed together or separately. We chose a random primer labeling scheme so that oligonucleotide probe design would not be restricted to any particular region of the mRNA molecule. In contrast, oligo(dT),12-18 priming protocols [16] limit design considerations to the 3’ end of the mRNA molecule.

Array image processing and data analysis
Cy3 and Cy5 fluorescence on microarray slides were scanned at 10 μm resolution using a GenePix 4000B scanner and saved as two single TIFF images. The intensities of spots on the two images were subsequently analyzed with GenePix Pro 3.0 software and a dataset was output. We used the following criteria to flag bad or extremely weak spots from the array dataset: spot area < 70 pixels, % saturated pixels > 50%, and sum of the median signal intensity < 1,000. Normalization of the array dataset was based on total median normalization, expression ratios were calculated for each non-flagged spot and log2 transformed.

Additional data files
The sequences of the 70-mers and corresponding GenBank accession numbers of the genes are available as an additional data file with the online version of this paper.

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