Title
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Empirical evaluation of a new method for calculating signal to noise ratio (SNR) for microarray data analysis

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ABSTRACT

Signal-to-noise-ratio (SNR) thresholds for microarray data analysis were experimentally determined with an oligonucleotide array that contained perfect match (PM) and mismatch (MM) probes based upon four genes from *Shewanella oneidensis* MR-1. A new SNR calculation, called signal to both standard deviations ratio (SSDR) was developed, and evaluated along with other two methods, signal to standard deviation ratio (SSR), and signal to background ratio (SBR). At a low stringency, the thresholds of SSR, SBR, and SSDR were 2.5, 1.60 and 0.80 with oligonucleotide and PCR amplicon as target templates, and 2.0, 1.60 and 0.70 with genomic DNA as target templates. Slightly higher thresholds were obtained at the high stringency condition. The thresholds of SSR and SSDR decreased with an increase in the complexity of targets (e.g. target types), and the presence of background DNA, and a decrease in the composition of targets, while SBR remained unchanged under all situations. The lowest percentage of false positives (FP) and false negatives (FN) was observed with the SSDR calculation method, suggesting that it may be a better SNR calculation for more accurate determination of SNR thresholds. Positive spots identified by SNR thresholds were verified by the Student t-test, and consistent results were observed. This study provides general guidance for users to select appropriate SNR thresholds for different samples under different hybridization conditions.
INTRODUCTION

Microarrays have become a routine tool for studying gene functions, regulations and networks in a variety of biological systems. Currently, this technology has been also applied to drug discovery and validation (7), microbial diagnostics (4, 10, 16, 20, 22, 31), mutation and single polymorphism nucleotide (SNP) detection (9), strain comparison and genotyping (1, 8, 21), species identification (32), array sequencing (35), environmental detection and monitoring (5, 6, 13, 24, 27, 28, 33), and evolutionary processes (14). However, due to small spot sizes, different degrees of uniformity of printing pins, and uneven hybridization, microarray spots inherently have relatively high noise, which presents a variety of challenges for quantitative analysis of microarray data. For example, how to distinguish a real signal from its background is still an unsolved problem, and a subset of this question is what parameters and thresholds should be used to differentiate a signal from a noise.

The signal-to-noise ratio (SNR) has been used to define a positive spot, and two general methods are currently used to calculate SNR values. One is to use the ratio of the differences between signal mean and background noise divided by background standard deviation (2). This calculation method has been commonly used in many signal-processing disciplines, such as radio, electronics and imaging (2, 30), and the threshold is usually set to 3.0 (30). The other method is to use the ratio of signal median divided by background median, and the threshold was set to 1.50 (26), and it was modified to calculate the SNR for a probe with replicate spots and set the threshold of 2.0 (18, 19). However, the determination of these thresholds is arbitrary and has not been experimentally validated. Although the background standard deviation of pixel intensities for each spot is included in the first calculation method, the signal standard deviation is not considered in either of the two SNR calculation methods. In addition, an SNR threshold
may vary with different types of targets, target compositions, and hybridization conditions, and hence it could be difficult to set a universal SNR threshold. Therefore new SNR calculation methods to include both signal and background standard deviations and experimental evaluations of SNR thresholds are needed.

The objectives of this study were to: (i) evaluate a new SNR calculation method for SNR calculation, (ii) determine appropriate SNR thresholds for differentiating signals from noises based on different SNR calculation methods, and (iii) examine the effects of target types, background DNA, and target compositions on the threshold determination. Our results demonstrated that our new calculation performed better than two other existing calculations, and that SNR thresholds were affected by hybridization stringency, types of target templates, background DNAs, and compositions of the target templates. Those results provide general guidance for users to select appropriate SNR thresholds under different conditions.

METHODS AND MATERIALS

Oligonucleotide probe design and microarray construction

50mer and 70mer perfect match (PM) and mismatch (MM) oligonucleotide probes were prepared as previously described (12). Briefly, four genes (SO1679, SO1744, SO2680, and SO0848) were selected from the Shewanella oneidensis MR-1 genome. For each gene, one 50- or 70-mer PM probe, and 45 MM (with 1 to 37 mismatches) probes were generated with three random MM probes at each level. All 368 designed oligonucleotides were commercially synthesized without modification by MWG Biotech Inc. (High Point, NC). The concentration of oligonucleotide probes was adjusted to 100 pmol/µl. Oligonucleotide probes prepared in 50% DMSO (Sigma Chemical Co., MO) were spotted onto UltraGAPS glass slides (Corning Life Science, NY) using
a PixSys 5500 robotic printer (Cartesian Technologies Inc., CA). Each probe had four replicates on a single slide. In total, there were 1472 (368 x 4) spots on the array. After printing, the oligonucleotide probes were fixed onto the slides by UV cross-linking (600 mJ of energy) according to the protocol of the manufacturer (Corning Life Science, NY).

5 **Target template preparations**

Four 70mer artificial targets (T1-SO1679, T2-SO1744, T3-SO2680 and T4-SO0848) that were complementary to the 70mer PM probes were synthesized by the Molecular Structure Facility at Michigan State University (East lasing, MI). The artificial oligonucleotide targets were labeled at the 5’-end with Cy5 (T1-SO1679, T2-SO1744 and T3-SO2680) or Cy3 (T4-SO0848) fluorescent dye during synthesis. The 70mer oligonucleotide targets also contained sequences of 50mer oligonucleotide targets.

Gene-specific primers were selected for the four selected genes (Supplementary Table S1) with each PCR product about 500 bp covering both 50mer and 70mer probe sequences. Each gene was amplified with *S. oneidensis* MR-1 genomic DNA as template using the standard PCR amplification protocol. The amplified PCR products were purified using the QIAquick PCR Purification Kit (QIAGEN Inc., CA) according to the protocol of the manufacturer. The purified PCR fragments were visualized and checked the sizes via agarose gel electrophoresis, and then quantified using the PicoGreen dsDNA Assay Kit (Invitrogen, CA).

Genomic DNAs from four bacteria were also used as target DNAs. *Shewanella oneidensis* MR-1, *Escherichia coli* S17, and *Pseudomonas sp.* strain G179 were grown in the LB medium to stationary phases, and *Desulfovibrio vulgaris* Hildenborough were grown in the LS medium. The cells were collected by centrifugation at 4000 x g at room temperature for 10 min. Their genomic DNAs were isolated and purified as described previously (34). *Methanococcus maripludis*
gDNA was provided by Sergey Stolyar at the University of Washington (Settle, WA). The yeast *Saccharomyces cerevisiae* was grown in the YPD medium to the saturation, and its genomic DNA was extracted using the glass bead method as described by Hoffman and Winston (15).

To test how bacterial ratios affect the determination of SNRs, *S. oneidensis* MR-1 gDNA was mixed with other four bacterial gDNAs (*D. vulgaris* Hildenborough, *E. coli* S17, *Pseudomonas sp.* strain G179, and *M. maripludis*) with three different ratios: A = 10 (*S. oneidensis* MR-1):1:1:1:1; B = 1 (*S. oneidensis* MR-1):1:1:1:1, and C = 1 (*S. oneidensis* MR-1):10:10:10:10, respectively. Each sample had the same amount of total gDNA (2.5 µg).

**Probe labeling, microarray hybridization, and image quantification**

PCR amplicons, the purified genomic DNA from pure cultures (500 ng), and mixed genomic DNAs (2.5 µg) were fluorescently labeled by random priming using Klenow fragment of DNA polymerase (12). Mixture I (35 µl) containing certain amounts (as indicated in different experiments) of genomic DNA and 20 µl of random primers (Invitrogen, CA) was heated at 98°C for 3 to 5 min, cooled on ice and then centrifuged. Mixture II (15 µl) containing 1 µl of 5 mM dATP, dGTP and dTTP and 2.5 mM dCTP, 2 µl (80 U) of Klenow (Invitrogen, CA) and 0.5 µl of Cy3 dye (Amersham BioSciences, UK) was added to mixture I. A total of 50 µl labeling reaction solution was incubated for 3 hr at 42°C. The labeling reaction was terminated by heating at 98°C for 3 min. The tubes were removed and placed on ice. The labeled cDNA targets were purified immediately using a QIAquick PCR purification column and concentrated in a Savant Speedvac centrifuge (Savant Instruments Inc., Holbrock, NY).

Labeled PCR amplicons, or genomic DNAs were resuspended in a 25 µl of hybridization solution containing 50% formamide, 5 x saline-sodium citrate (SSC), 0.1% SDS, and 0.1 mg/ml of herring sperm DNA (Invitrogen, CA). The hybridization solution was incubated at 95-98°C.
for 5 min, centrifuged to collect condensation, and kept at 50°C. The solution was immediately
applied onto the microarray slide and hybridization was carried out in a waterproof Corning
Hybridization chamber (Corning Life Science, NY) submerged in a 45°C water bath in the dark
for 16 h (12). Washing was performed immediately in the following steps: (i) in a solution
containing 2 x SSC and 0.1% SDS at 40°C for 5 min and repeated once, (ii) in a solution
containing 0.1 x SSC and 0.1% SDS at room temperature for 10 min and repeated once, and (iii)
in 0.1 x SSC at room temperature for 2 min and repeated once. Slides were dried by compressed
air prior to scanning. The same batch slides and the same settings were used for all experiments.
The laser power was set to 95%, and photomultiplier tube (PMT) efficiency was set to 70%. Five
slides (4 replicated spots in each slide) were used for each condition, and hence each spot had up
to 20 data points. Hybridized microarray slides were scanned using ScanArray™ Express
microarray analysis system (Perkin Elmer®, MA). Spot signal, spot quality, and background
fluorescent intensities of scanned images were quantified with ImaGene version 6.0
(Biodiscovery Inc., Los Angeles, CA).

15 Data analysis

Data analysis included the following four major steps. (i) Defining positive and negative spot
pools: Microarray detection mainly depends on probe specificity and hybridization stringency
(e.g. temperature), and two levels of stringency were used in this study. A high-level stringency
is expected to eliminate cross-hybridization for the probes with a higher probe-target similarity, a
longer continuous stretch length, and a lower free energy. At both stringencies, positive and
negative pools were defined (Supplementary Table S2 & S3). At the high stringency, a positive
50mer probe has a sequence identity >90%, a stretch length >20, and free energy <-35 kcal/mol
with its non-targets, and a negative probe has a sequence identity ≤90%, a stretch length ≤20,
and free energy $\geq -35$ kcal/mol with its non-targets. Our previous experimental results showed that such high stringency hybridization can be achieved at 50°C plus 50% formamide (17). Similarly, a positive 70mer probe has a sequence identity $>90\%$, a stretch length $>25$, and free energy $<-50$ kcal/mol with its non-targets, and a negative probe has a sequence identity $\leq 90\%$, a stretch length $\leq 25$, and free energy $\geq -50$ kcal/mol with its non-targets. At a low stringency, a positive 50mer probe has a sequence identity $>85\%$, a stretch length $>15$, and free energy $<-30$ kcal/mol with its non-targets, and a negative probe has a sequence identity $\leq 85\%$, a stretch length $\leq 15$, and free energy $\geq -30$ kcal/mol with its non-targets (12). The low stringency generally corresponds to hybridization at 42°C plus 50% formamide. Similarly, a positive 70mer probe has a sequence identity $>85\%$, a stretch length $>20$, and free energy $<-40$ kcal/mol with its non-targets, and a negative probe has a sequence identity $\leq 85\%$, a stretch length $\leq 20$, and free energy $\geq -40$ kcal/mol with its non-targets (12). In addition, the probes that do not qualify for either positive pool or negative pool were ignored for further analysis. (ii) Microarray spot analysis: Spot intensity data were extracted from ImaGene output files. The values for gene ID, flag, signal mean ($\bar{S}$), background mean ($\bar{B}$), signal standard deviation ($\sigma$), and background standard deviation ($\sigma_B$) were extracted from ImaGene output files. After the removal of bad spots, the rest of spots (including potential empty spots and good spots) were kept for further analysis. All processes were conducted with Microsoft Excel software. (iii) Calculation of SNR values: For each spot, three methods were used to calculate SNR values:

\[
SSR = \frac{(\bar{S} - \bar{B})}{\sigma} \quad (I)
\]

\[
SBR = \frac{\bar{S}}{\bar{B}} \quad (II)
\]
\[
SSDR = \frac{(\bar{S} - \bar{B})}{(\sigma_s + \sigma_b)}
\] (III)

Where \(\bar{S}\) and \(\bar{B}\) are the signal mean and the background mean of pixel intensities, respectively, and \(\sigma_s\) and \(\sigma_b\) are the standard deviation of signal and background, respectively. Based on false positive and false negative spots at different values of SSR, SBR, and SSDR (in comparison with the defined positive and negative spot pools), their thresholds were determined by (a) minimizing false positives, (b) minimizing false negatives, and (c) optimizing the overall percentage of false positives and false negatives. (vi) The student-t test analysis of threshold-identified positive spots: The values of signal (\(S\)) and background (\(B\)) were extracted for a probe with replicate spots from ImaGene output files, and their mean (\(\bar{S}_m\) and \(\bar{B}_m\), respectively) and standard deviation (\(\sigma_{s.m}\) and \(\sigma_{b.m}\), respectively) values were calculated. Outliers were removed if |\(S - \bar{S}_m\)| ≥ 2.0 * \(\sigma_{s.m}\), or |\(B - \bar{B}_m\)| ≥ 2.0 * \(\sigma_{b.m}\), and this process continued until outliers were recursively removed. The final \(\bar{S}_m\), \(\bar{B}_m\), \(\sigma_{s.m}\) and \(\sigma_{b.m}\) were used for the Student t-test, and the significance between \(\bar{S}_m\) and \(\bar{B}_m\) was statistically evaluated for each probe at a given p value.

**Data analysis for Desulfovibrio vulgaris Hildenborough microarrays**

Both wild type and \(\Delta fur\) mutant of \(D. vulgaris\) cells were grown in the LS4D medium with 60 \(\mu\)M of iron, and microarray data were obtained as previously described (3). SSDR method was used to detect positive spots with the threshold of 0.80, and details of data analysis were conducted as previously described (3).

**Data analysis for GeoChip with a soil sample**

A soil sample was taken from a plot at BioCON (23), and 5 g of soil was used to extract DNA. GeoChip (13) was used to detect functional genes in such a microbial soil community. SSR, SBR and SSDR were used to detect positive spots with thresholds of 2.0, 1.6, and 0.8, respectively,
and details for labeling, hybridization, and scanning were performed as described previously (13).

RESULTS

A new SNR calculation method

To consider the signal intensity and background noise as well as their standard deviations for each spot, a new calculation method, termed SSDR (signal to both standard deviations ratio), was developed. SSDR differs from other two SNR calculation methods (SSR and SBR) in that it takes account into the signal standard deviation as a part of the denominator. The relationship between SSDR and signal or background intensity (together with their standard deviations) can be simply represented in Fig. 1, which shows both signal and background standard deviations are equally important for determination of SNR thresholds. When SSDR is ≥1.0, the difference between the signal intensity and background noise is equal or larger than the sum of the signal and background standard deviations. In this case, the pixel values of signal intensity are completely separated from those of background noises (Fig. 1). Intuitively, such a spot should represent positive signal. When SSDR < 1.0, overlaps of the pixel values between signals and background noises exist (Fig. 1). In this case, some spots could be positive while some are not, but the key question is what is the minimum SNR (e.g. SSDR) threshold for distinguishing the signal from its background noise. Thus in this study, we will experimentally determine the threshold of SSDR for differentiating signals from noises.

Experimental determination of SNR thresholds

To determine appropriate thresholds for distinguishing signal from noise for a single spot on the array, four synthesized targets were hybridized with the array at a final concentration of 10 pg
per oligonucleotide. Based on the predefined positive and negative pools at the low stringency, 60 (27 for 50mer, 33 for 70mer) probes are expected to be positives, 249 negative, and 59 ignored (Supplementary Table S2). The ignored probes fail to satisfy the definition of positive or negative spots. Based on the predicted pools of the positive and negative spots, the number of the false positive (FP) and false negative (FN) spots were calculated at different scenarios. First, false positive spots were minimized. To have no false positives, the thresholds of SSR, SBR, and SSDR should be 5.0, 5.0, and 1.0, respectively (Table 1, Fig. 2). If 1% FP spots were allowed, the thresholds were 4.0 for SSR, 3.5 for SBR, and 0.90 for SSDR (Table 1, Fig. 2). The thresholds would be 2.0, 1.8, and 0.70 for SSR, SBR and SSDR, respectively when 5% FP spots could be tolerated (Table 1, Fig. 2). Second, false negatives were minimized. The thresholds of SSR, SBR, and SSDR should be 0.5, 0.5, and 0.3, respectively if there were no FN spots (Table 1, Fig. 2). If 1% FN spots were allowed, the thresholds were 1.5 for SSR, 1.2 for SBR, and 0.70 for SSDR (Table 1, Fig. 2). The thresholds would be 2.5, 1.6, and 0.85 for SSR, SBR and SSDR, respectively when 5% FN spots were allowed (Table 1, Fig. 2). In addition, the thresholds of SSR, SBR and SSDR were determined by optimizing the total percentage of FP and FN spots. Generally speaking, higher percentages of FP were observed at a lower threshold of SSR, SBR, or SSDR. For example, the percentages of FP were 11.8%, 12.2%, and 7.9% at SSR = 1.5, SBR = 1.4, and SSDR = 0.5, respectively, which led to 13.0%, 14.9%, and 8.3% of total percentages of FP and FN spots, respectively (Fig. 2). On the contrary, higher percentages of FN were observed at a higher threshold of SSR, SBR, or SSDR. For example, the percentages of FN were 17.1%, 19.0%, and 12.8% at SSR = 4.0, SBR = 4.0, and SSDR = 1.2, respectively, resulting in 18.2%, 19.9%, and 13.1% of total percentages of FP and FN spots, respectively (Fig. 2). However, relatively low and stable percentages of FP and FN spots were shown when the values
of SSR, SBR or SSDR were in a certain range. For example, when SSR were between 2.0 and 3.0, the percentages of FP and FN spots were 8.0-9.7%; those percentages were 10.0-14.9% when SBR were 1.4-3.0; SSDR were 0.6-1.0 when those percentages were 5.0-8.0% (Fig. 2). Therefore, the above results indicate that the thresholds of SSR, SBR and SSDR can be in a certain range with a relatively low percentage of FP and FN spots although optimal thresholds were determined to be SSR = 2.5, SBR =1.6, and SSDR = 0.80.

Under a high stringency, 33 (13 for 50mer and 20 for 70mer) probes were positives, 280 (147 for 50mer and 137 for 70mer) were negative, and 55 were ignored (Supplementary Table S3). The thresholds of SSR, SBR and SSDR were determined using the same strategies as described above. First, through the minimization of false positives, the thresholds of SSR, SBR, and SSDR were determined to be 5.0, 5.0 and 1.1, respectively when no FP spots were allowed; those thresholds were 4.5 for SSR, 4.0 for SBR and 1.0 for SSDR if 1% FP spots were allowed; if 5% FP spots were tolerated, those thresholds of SSR, SBR and SSDR were 2.5, 2.0 and 0.70, respectively (Table 1, Fig. 3). Second, through the minimization of false negatives, the thresholds of SSR, SBR and SSDR were determined to be approximately 1.0, 1.0, and 0.5, respectively when no FN spots were allowed; if 1% FN spots were allowed, those thresholds were 2.0 for SSR, 1.4 for SBR, and 0.75 for SSDR; they would be 3.0 for SSR, 1.8 for SBR, and 0.95 for SSDR if 5% FN spots were tolerated (Table 1, Fig. 3). Finally, by optimizing the total percentage of FP and FN spots on the array, the thresholds of SSR, SBR and SSDR were determined to be 3.0, 2.0 and 0.90, respectively (Fig. 3). The results demonstrated that the thresholds of SSR, SBR and SSDR increased with an increase in stringency of defined positive and negative probe pools. In addition, both Fig. 2 and Fig. 3 showed that the lowest percentages of FP and FN spots were observed with the SSDR calculation, and that an optimization of
percentage of FP and FN appeared to be the best method for SNR determination. Therefore, for further experiments, the defined positive and negative pools with the low stringency were used, and an optimization of false positives and false negatives was considered the best method for SNR determination.

Effects of target types on the SNR threshold determination

To determine the impacts of target types on the threshold selection, 100 pg of each PCR amplicon or 500 ng of *S. oneidensis* MR-1 gDNA was also labeled with Cy3 and hybridized with the array, and the thresholds of SNR, SBR and SSDR were determined by optimizing the percentage of FN and FP spots. The same thresholds were obtained for PCR amplicon targets as the synthesized oligonucleotides although the PCR amplicon targets caused slightly higher percentages of total FN and FP than synthesized oligonucleotides. For example, the thresholds of SSR were 2.5 for oligonucleotide and PCR amplicon targets when the percentages of FP and FN were 8.0% and 8.7%, respectively (Fig. 4A). However, the thresholds of SSR of 2.0 (Fig. 4A) and SSDR of 0.70 (Fig. 4C) for gDNA were lower than those for synthesized oligonucleotides, or PCR amplicons. The percentages of total FN and FP of gDNA were a bit higher than synthesized oligonucleotide, or PCR amplicon targets (Fig. 4). For example, the percentage of FN and FP was 7.1% for gDNA compared to 5.0% for oligonucleotide targets and 6.51% for PCR targets when the SSDR thresholds of 0.8, 0.8 and 0.7 were used for oligonucleotide, PCR amplicon, and gDNA targets, respectively (Fig. 4C). In contrast to SSR and SSDR, SBR remained unchanged with different types of targets. The results also confirmed that the lowest percentage of false positives and false negatives was observed with the SSDR calculation method.

Effects of background DNA on threshold determination
When microarrays are used for community analysis, significant amount of DNAs from non-target organisms as background exists, and it could affect SNR threshold determination. To examine the effect of such background DNA on the SSR, SBR and SSDR thresholds, 500 ng of *S. oneidensis* gDNA, or 10 pg per oligonucleotide target was mixed with 1.0 µg of the yeast gDNA, and their thresholds were determined as described in Fig. 2. With the yeast gDNA as background, the thresholds of SSR and SSDR for *S. oneidensis* gDNA were determined to be 1.75 and 0.65, respectively, which were slightly lower than those without the yeast gDNA as background (Fig. 5A). Similarly, the thresholds of SSR and SSDR changed from 2.5 and 0.80 to 2.0 and 0.70, respectively when synthesized oligonucleotide targets were spiked into the yeast gDNA (Fig. 5B). However, the thresholds of SBR did not change with the target type, or the background DNA (Fig. 5). The results indicate that the thresholds of SSR and SSDR decreased with the addition of yeast gDNA as background, but that the threshold of SBR stayed the same.

To further understand why background DNA caused a decrease in the thresholds of SSR and SSDR, the changes in signal mean, background mean, and their standard deviations for each spot with the yeast DNA as non-target DNAs were compared with those without the yeast DNA (Fig. 6). When the yeast gDNA was added into the *S. oneidensis* gDNA, the trends of the signal mean and the background mean did not change, but the average signal and background standard deviations increased to 124% and 134%, respectively compared to *S. oneidensis* gDNA only (Fig. 6A). Similarly, when the oligonucleotide targets was used as target templates with the background yeast gDNA, the average signal mean and the average background mean did not change significantly, but both average signal and background standard deviations increased to 129% and 148%, respectively in comparison with the oligonucleotide targets only (Fig. 6B). The
results indicated that an increase in both signal and background standard deviations might result in lower thresholds of SSR and SSDR when non-target DNAs are present.

### Determination of SNR thresholds for artificial bacterial mixtures

To examine how DNA mixtures with different compositions affect the SNR threshold determination, *S. oneidensis* gDNA was mixed with other four bacteria in the ratios of (A) 10:1:1:1:1, (B) 1:1:1:1:1, and (C) 1:10:10:10:10, and each mixture had 2.50 µg of gDNA in total. The optimal thresholds of SSR, SBR, and SSDR were determined to be 2.00, 1.60, and 0.70, respectively for Mixture (A), and 1.75, 1.60, and 0.60, respectively for Mixture (B) ([Table 2](#)).

There were only about 23.3% of the defined positive spots were detected on the array for Mixture (C), so no thresholds of SSR, SBR, or SSDR could be estimated ([Table 2](#)). The results showed that the thresholds of SSR and SSDR were decreased with a decrease in the percentage of the target (*S. oneidensis* gDNA) in the sample, but that the thresholds of SBR were not affected, which is also consistent with the results observed with different types of target or with the yeast DNA. It is possible that a decrease in target concentration in a mixed sample may lead to a higher rate for FN or/and FN + FP.

### Verification of identified positive spots

To further understand if the identified positive spots based on the above thresholds have significantly higher signals than their backgrounds, the Student t test was used to determine if a probe with replicate spots was positive at a given p value. Since genomic DNA is most commonly used target, this experiment was carried out with *S. oneidensis* MR-1 gDNA (500 ng).

The predefined positives (at a low stringency), the t-test identified positives (at p<0.01), and SNR threshold-identified (2.0 for SSR, 1.6 for SBR and 0.70 for SSDR) positives were compared, and relatively consistent results were observed ([Table 3](#)). Among 368 probes, 60, 249
and 59 were defined as positive, negative, and ignored, respectively under a low stringency. Based on t-test, a total of 76 probes were identified as positives with 58 from the defined positives, 3 from the defined negatives, and 15 from the ignored pool at p<0.01 (Table 3). Similar numbers of positives to the t-test analysis were identified based on the SNR thresholds determined above. For example, at the SSDR threshold of 0.70, 81, 79, and 75 positives were identified at positive rates of >50%, >70%, and >90%, respectively (Table 3). These results demonstrated that the positive spots or probes identified by SNR thresholds and by the Student t-test were very similar, which was also consistent with the predefined positives and negatives.

Determination of positive spots by SSDR threshold for pure culture and soil samples

To demonstrate the application of SSDR thresholds for determining positive spots, two sets of data were used. One was pure cultures of wild type (WT) and $\Delta$fur mutant (JW707) Desulfovibrio vulgaris Hildenborough (DvH) with the DvH oligonucleotide microarray (3), and the other was a BioCON soil sample with GeoChip (13). For the first data set, the SSDR threshold of 0.80 was used. The average SSDR for the fur probe was 0.25 for the $\Delta$fur mutant, and 2.16 for WT, confirming the absence of this gene in the mutant (Table 4). Fur is a transcriptional regulator, and negatively regulates several genes in the fur regulon when it binds to a promoter. The microarray data did show that genes such as feoA, feoB, fld, and gdp predicted in the fur regulon (25) were up-regulated in the mutant JW707 (Table 4). The Fur regulator has been showed to be involved in oxidative stress responses, which are mainly controlled by the PerR regulator (25). Indeed, our results also showed that ahpC, rbr and perR were over-expressed in the JW707 mutant (Table 4). In addition, it was observed that the expression of genes (cobI, COG-fepB, fepC, and COG-fepD) involved in iron uptake was repressed, and that the expression of genes (bfr and ftn) involved in iron storage was induced (Table 4). This is consistent with the fact that
more iron may accumulate in the mutant due to the absence of Fur protein. It is noted that
different cutoffs for up-regulation and down-regulation were used in this study (two-fold) and
the previous study (3).

Despite our successful demonstration of SSDR application for pure cultures, a similar
demonstration with environmental samples, such as soil, is much more difficult. Thus in this
study, we used one soil sample with three hybridizations to see the number of detected positive
spots, and their unique and overlap spots among replicates (Table 5). With the thresholds of 2.0
for SSR, 1.6 for SBR, and 0.80 for SSDR, the average detected spots were 3858, 4372, and 3828
for SSR, SBR, and SSDR, respectively (Table 5). Although the fewest positive spots (3903) were
detected by SSDR, it had the highest overlap spot number (3761) and rate (96.3%), but the
lowest unique spot number (97) and rate (2.5%), indicating that SSDR is a more accurate method
to discriminate true signals from background noise (Table 5). Therefore, the above results
demonstrated that the SSDR method with an appropriate threshold could be used to determine
positive spots for both pure culture and environmental (e.g. soil) samples.

DISCUSSION

How to distinguish a real signal from its background remains challenging in microarray data
analysis, and this study focuses on the experimental determination of SNR thresholds. The
determination of SNR thresholds is an important step for the generation of high quality
microarray data, and its accuracy is critical for the subsequent data processing and biological
interpretation of microarray results. Thus this study experimentally determined the thresholds of
SNR under different scenarios. The results of this study should provide guidance for users to
select appropriate SNR thresholds for their experiments.
Considering the standard deviations of pixel intensity of both signal and background, a new calculation method was developed. It has a couple of advantages. First, the signal standard deviation was considered as a parameter together with background standard deviation. Since the pixel intensities of a spot are not uniform, its standard deviation significantly affects the ability of distinguishing a true signal from its background. In this case, a consideration of signal standard deviation can more accurately reflect microarray hybridization behaviors, and more reliably identify a true spot and its threshold. Second, our experimental data demonstrated that fewer false positives and negatives were observed with this method compared to two other methods. SBR did not change with target types, or background DNA since this calculation does not consider signal standard deviation or background standard deviation, but it generally had a high percentage of FN and FP spots, and it may not be a good parameter to distinguish a true signal from its background noise. Therefore, this new method may be used for a general SNR calculation, and more accurate thresholds could be obtained with this calculation.

Three possible scenarios, minimizing false positives, minimizing false negatives and optimizing false positives and false negatives, were considered to determine the ranges of SNR thresholds for detecting real signals, but the threshold values for optimal false positives and negatives could be used more often. By optimizing the percentage of FP and FN spots, those thresholds of SSR and SBR determined in this experiment appeared to be lower compared to other commonly accepted thresholds. For example, the threshold of SSR was set to be 3.0 (30), and SBR to be 1.50 (26) or 2.0 (19). Considering all three methods for SNR determination, the ranges of SNR thresholds for gDNA targets were summarized in Table 6. For example, the thresholds of SSR were in the range of 0.5 (no FN), 2.0 (optimal), to 4.0 (no FP), and those of SSDR were in a range of 0.3 (no FN), 0.7 (optimal), to 0.9 (no FP) under a low stringent
condition. Those ranges provide a general guideline for users to select appropriate SNR thresholds based on their experiments. There are two points needed to be mentioned. One is that the error rate of 5% (FP + FN) was used in his study, which is considered reasonable since microarray data have relatively high variations due to various reasons, such as the small size, degrees of uniformity of printing pins, and uneven hybridization. The other is that those SNR threshold values determined here for DNA microarray studies under different stringencies and different target types or/and concentrations may only be applied to long (50-70mer) oligonucleotide microarrays. An application of such parameters to short (18-25mer) oligonucleotide microarrays remains unclear, which needs to be further evaluated.

It is known that probe specificity and the stringency of hybridization conditions affect the determination of SNR thresholds. Two stringency conditions were used in this study. As expected, a lower threshold (e.g. SSR = 2.0, SBR = 1.6, and SSDR = 0.80) can be used for detecting specific hybridizations under high stringent hybridization conditions (e.g. at a high temperature of 50°C), and a higher threshold (e.g. SSR = 3.0, SBR = 2.0, and SSDR = 0.90) may be required for detecting specific hybridizations under low stringent hybridization conditions (e.g. at a low temperature of 42°C).

Many factors, such as target type, background DNAs, target composition, and target amount in the tested sample affect the SNR threshold determination. Microarray hybridization signal intensity is determined by the number of probe molecules bound to microarray surface, the number of labeled targets present in the sample, and their ratios, which are closely related to target type and their concentrations. In this study, the synthesized oligonucleotides and PCR amplicons are the simplest target, and both are similar, and they had almost the same thresholds. *S. oneidensis* MR-1 gDNA is more complex, and its threshold was a bit lower. Similarly, the
complexity of target is expected to increase in the presence of background DNA, and hence a lower threshold was observed. Further analysis revealed that this might be due to an increase in background standard deviation. This is validated by the fact that the thresholds of SBR did not change with the target type or with background DNA. With the mixed templates, Mixture (A) contained > 70% of real target (S. oneidensis gDNA), the threshold did not change significantly. However, a slightly decrease in threshold was observed in Mixture (B) with 20% of real target, and it became undeterminable for Mixture C containing about 2.5% of real target. The decrease of the thresholds with a decrease of the target template composition can be explained by an increase in sample noise when the target concentration decreased. Sample noise is mostly from labeled molecules in a sample. For example, labeled target solutions can react in a non-specific manner on microarrays, which masks the interactions between a probe and its target and obscures the microarray signal. Therefore, an increase in non-target concentrations leads to an increase in noise, which may reduce SNR thresholds to compromise microarray detectivity. This is also consistent with our observations for different types of target or with background DNA since labeled non-targets such as background DNAs cause a significant amount of background noise.

As previous studies showed, the detection limits for 50mer oligonucleotide and 70mer oligonucleotide arrays were estimated to be 25 to 100 ng of gDNA (11) for a pure culture although a higher sensitivity (5~10 ng gDNA) was also observed (24, 29). In the presence of background DNA, the detection limit for 50mer oligonucleotide was estimated to be 50~100 ng of gDNA (24, 29). In the Mixture C, the real target was about 63 ng of gDNA, so it was not surprising that only 23.3% of defined positive probes had true signals. These results suggest that
a threshold might change with target compositions, which are closely related to the microarray sensitivity.

It is also noted that the amount of target may affect the threshold determination. For example, a higher threshold may be required when a relatively large amount of target is used. In this study, we used the optimal concentrations of 10 pg for each oligonucleotide, 100 pg for each PCR amplicon, and 500 ng for gDNA, which are considered equivalent amounts of the target in samples. This is a simulation for a pure culture, or a mixture of a few known microorganisms. For a sample with many unknown microorganisms, such as microbial communities in soil and the human intestinal tract, a determination of SNR thresholds may be even more challenging. Because of unequal abundance, low-abundant genes/microorganisms may not be detected even at a relatively low threshold.

In summary, three methods were used to calculate SNR values, and the newly developed calculation showed a better performance for distinguishing a true signal from its background than the other two methods. The positives identified based on SNR thresholds were verified by the Student t-test across many replicate data, and consistent results were obtained. This study provides guidance for the selection of SNR thresholds for different samples, such as PCR amplicons, and genomic DNA from pure cultures and simple mixed cultures.

ACKNOWLEDGEMENTS

The authors thank Meiying Xu for providing GeoChip data of the BioCON soil sample and Yuting Liang for providing Desulfovibrio vulgaris Hildenborough microarray data of both wild type and the Δfur mutant. This research was supported by The United States Department of
Energy under Genomics:GTL program through the Virtual Institute of Microbial Stress and Survival (VIMSS; http://vimss.lbl.gov), and the Environmental Remediation Science Program.
REFERENCES


FIGURE LEGENDS

Fig. 1. Schematic presentation of the SSDR calculation method. A, B and C represent SSDR<1.0, SSDR = 1.0 and SSDR >1.0, respectively. All the four parameters used in calculation extracted from the ImaGene output files (Manual of ImaGene).

Fig. 2. Determination of thresholds of SSR (A), SBR (B), and SSDR (C) at a low stringency by minimizing the percentage of false positive and false negative spots. 10 pg of each synthesized oligonucleotide was used to hybridize with the array, and 5 replicate slides were used. SSR, SBR and SSDR were determined to be 2.5, 1.6, and 0.80, respectively.

Fig. 3. Determination of thresholds of SSR (A), SBR (B), and SSDR (C) at a high stringency by minimizing the percentage of false positive and false negative spots. 10 pg of each synthesized oligonucleotide was used to hybridize with the array, and 5 replicate slides were used. SSR, SBR and SSDR were determined to be 3.0, 2.0, and 0.90, respectively.

Fig. 4. Effects of target types on the thresholds and the percentages of false positive (FP), false negatives (FN), and both (FP+FN) for SSR (A), SBR (B), and SSDR (C). The left y-axis presents the optimal threshold, and the right y-axis presents the percentage of FP, FN, or FP + FN under the optimal threshold. Targets used were synthesized oligonucleotides (10 pg each), PCR amplicons (100 pg each), and S. oneidensis MR1 gDNA (500 ng). The more significant p value is shown on the top of each column with the following notions: nd=no difference,
Fig. 5. Effects of background DNA on the determination of SSR, SBR and SSDR thresholds. 500 ng of *S. oneidensis* MR-1 gDNA (A) and 10 pg for each synthesized oligonucleotide (B) were spiked into 1.0 µg of yeast gDNA. For synthesized oligonucleotide targets, the yeast gDNA was first labeled and then mixed with the spiked oligonucleotides. *S. oneidensis* MR-1 gDNA was first mixed with the yeast gDNA, and then labeled together. The significance is shown on the top of each column with the following notions: nd=no difference, *=p<0.10, **=p<0.05, and ***=p<0.01 (the Student t test) when thresholds with background DNA were compared to those without background DNA.

Fig. 6. Comparison of changes in signal mean, background mean, signal standard deviation and background standard deviation for each spot on the array when the yeast gDNA was added into the *S. oneidensis* gDNA (A), or the synthesized oligonucleotide targets (B).
Table 1. The thresholds of SSR, SBR and SSDR determined by minimizing the percentage of false positive (FP) spots or false negative (FN) spots on the array using synthesized oligonucleotide targets under low and high stringencies.

<table>
<thead>
<tr>
<th></th>
<th>A. The thresholds of SSR, SBR and SSDR at the defined low stringency</th>
<th>B. The thresholds of SSR, SBR and SSDR at the defined high stringency</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SSR</td>
<td>SBR</td>
</tr>
<tr>
<td>No FP</td>
<td>5.0</td>
<td>5.0</td>
</tr>
<tr>
<td>1% FP</td>
<td>4.0</td>
<td>3.5</td>
</tr>
<tr>
<td>5% FP</td>
<td>2.0</td>
<td>1.8</td>
</tr>
<tr>
<td>5% FN</td>
<td>2.5</td>
<td>1.6</td>
</tr>
<tr>
<td>1% FN</td>
<td>1.5</td>
<td>1.2</td>
</tr>
<tr>
<td>No FN</td>
<td>0.5</td>
<td>0.5</td>
</tr>
</tbody>
</table>
Table 2. The thresholds of SSR, SBR and SSDR and the percentages of false positives, false negatives, or both for artificial bacterial mixtures. Genomic DNAs from Mixture A, B and C containing *S. oneidensis* MR-1 (bold)) and other four bacteria with different ratios were used as targets. SSR, SBR, SSDR, and percentages of false positives and false negatives were determined as described in Fig. 2. Five slides were used.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of defined positive spots</td>
<td>300</td>
<td>300</td>
<td>300</td>
</tr>
<tr>
<td>% of detected positive spots</td>
<td>318</td>
<td>311</td>
<td>70</td>
</tr>
<tr>
<td><strong>SSR</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Threshold</td>
<td>2.0</td>
<td>1.75</td>
<td>ND</td>
</tr>
<tr>
<td>% of false positives (FP)</td>
<td>4.3</td>
<td>3.5</td>
<td>0</td>
</tr>
<tr>
<td>% of false negatives (FN)</td>
<td>3.3</td>
<td>3.4</td>
<td>76.7</td>
</tr>
<tr>
<td>% of total FP and FN</td>
<td>7.6</td>
<td>6.9</td>
<td>76.7</td>
</tr>
<tr>
<td><strong>SBR</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Threshold</td>
<td>1.60</td>
<td>1.60</td>
<td>ND</td>
</tr>
<tr>
<td>% of false positives (FP)</td>
<td>4.7</td>
<td>3.6</td>
<td>0</td>
</tr>
<tr>
<td>% of false negatives (FN)</td>
<td>3.3</td>
<td>4.7</td>
<td>76.7</td>
</tr>
<tr>
<td>% of total FP and FN</td>
<td>8.0</td>
<td>8.3</td>
<td>76.7</td>
</tr>
<tr>
<td><strong>SSDR</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Threshold</td>
<td>0.70</td>
<td>0.60</td>
<td>ND</td>
</tr>
<tr>
<td>% of false positives (FP)</td>
<td>2.7</td>
<td>2.2</td>
<td>0</td>
</tr>
<tr>
<td>% of false negatives (FN)</td>
<td>2.8</td>
<td>3.7</td>
<td>76.7</td>
</tr>
<tr>
<td>% of total FP and FN</td>
<td>5.5</td>
<td>5.9</td>
<td>76.7</td>
</tr>
</tbody>
</table>
Table 3. Comparison of positive probes identified by probe design criteria, by the Student t-test, and by SNR thresholds. 368 probes were valid for analysis when 500 ng of labeled S. oneidensis MR1 gDNA hybridized with the array. Five slides were used with four replicates in each slide, so each probe had up to 20 spots.

A. Defined and t-test identified positive probes at the low stringency (p<0.01 for the Student t test)

<table>
<thead>
<tr>
<th></th>
<th>No. of probes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Defined positives</td>
<td>60&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Defined negatives</td>
<td>249&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Ignored</td>
<td>59&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>No. of t-test positives</td>
<td>57&lt;sup&gt;a&lt;/sup&gt; + 4&lt;sup&gt;b&lt;/sup&gt; + 15&lt;sup&gt;c&lt;/sup&gt; = 76</td>
</tr>
<tr>
<td>No. of t-test negatives</td>
<td>3&lt;sup&gt;a&lt;/sup&gt; + 245&lt;sup&gt;b&lt;/sup&gt; + 44&lt;sup&gt;c&lt;/sup&gt; = 292</td>
</tr>
</tbody>
</table>

B. SNR-threshold-identified positive probes at different positive rates (PR = the number of positive spots identified by SNR thresholds *100/total number of spots for each probe).

<table>
<thead>
<tr>
<th>Threshold</th>
<th>PR &gt; 50%</th>
<th>PR &gt; 70%</th>
<th>PR &gt; 90%</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of SSR-identified positives (% of t-test positives)</td>
<td>2.0</td>
<td>58&lt;sup&gt;a&lt;/sup&gt; + 7&lt;sup&gt;b&lt;/sup&gt; + 21&lt;sup&gt;c&lt;/sup&gt; = 86 (113%)</td>
<td>58&lt;sup&gt;a&lt;/sup&gt; + 5&lt;sup&gt;b&lt;/sup&gt; + 19&lt;sup&gt;c&lt;/sup&gt; = 82 (108%)</td>
</tr>
<tr>
<td>No. of SBR-identified positives (% of t-test positives)</td>
<td>1.6</td>
<td>58&lt;sup&gt;a&lt;/sup&gt; + 8&lt;sup&gt;b&lt;/sup&gt; + 25&lt;sup&gt;c&lt;/sup&gt; = 91 (120%)</td>
<td>57&lt;sup&gt;a&lt;/sup&gt; + 6&lt;sup&gt;b&lt;/sup&gt; + 23&lt;sup&gt;c&lt;/sup&gt; = 86 (113%)</td>
</tr>
<tr>
<td>No. of SSDR-identified positives (% of t-test positives)</td>
<td>0.70</td>
<td>59&lt;sup&gt;a&lt;/sup&gt; + 4&lt;sup&gt;b&lt;/sup&gt; + 18&lt;sup&gt;c&lt;/sup&gt; = 81 (107%)</td>
<td>59&lt;sup&gt;a&lt;/sup&gt; + 3&lt;sup&gt;b&lt;/sup&gt; + 17&lt;sup&gt;c&lt;/sup&gt; = 79 (104%)</td>
</tr>
</tbody>
</table>

<sup>a</sup> the size of defined positive probe pool;  
<sup>b</sup> the size of defined negative probe pool;  
<sup>c</sup> the number of ignored probes based on defined positive, negative, and ignored probe pools.
Table 4. Examples of transcriptional changes of function-known genes in \textit{Afur} mutant (JW707) and wild type (WT) of \textit{D. vulgaris Hildenborough}.

<table>
<thead>
<tr>
<th>Category/DVU</th>
<th>Gene</th>
<th>Annotated function</th>
<th>SSDR(^a)</th>
<th>Expression ratio (JW707/WT)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>JW707</td>
<td>WT</td>
</tr>
<tr>
<td><strong>Genes in the predicted Fur regulon(^b)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DVU0303</td>
<td>\textit{genZ}</td>
<td>GenZ, hypothetical protein</td>
<td>2.16±0.285(^c)</td>
<td>1.78±0.172(^c)</td>
</tr>
<tr>
<td>DVU0304</td>
<td>\textit{genY}</td>
<td>GenY, hypothetical protein</td>
<td>2.47±0.277</td>
<td>2.15±0.122</td>
</tr>
<tr>
<td>DVU0763</td>
<td>\textit{gdp}</td>
<td>GGDEF domain protein</td>
<td>2.28±0.321</td>
<td>1.96±0.231</td>
</tr>
<tr>
<td>DVU0942</td>
<td>\textit{fur}</td>
<td>Fur, transcriptional regulator</td>
<td>0.25±0.036</td>
<td>2.16±0.116</td>
</tr>
<tr>
<td>DVU2571</td>
<td>\textit{feoB}</td>
<td>Ferrous iron transport protein B</td>
<td>1.97±0.166</td>
<td>1.95±0.142</td>
</tr>
<tr>
<td>DVU2572</td>
<td>\textit{feoA}</td>
<td>Ferrous iron transport protein A</td>
<td>1.72±0.321</td>
<td>1.83±0.211</td>
</tr>
<tr>
<td>DVU2574</td>
<td>\textit{feoA}</td>
<td>Ferrous ion transport protein</td>
<td>2.17±0.277</td>
<td>1.93±0.102</td>
</tr>
<tr>
<td>DVU2680</td>
<td>\textit{flb}</td>
<td>Flavodoxin</td>
<td>1.88±0.130</td>
<td>2.06±0.133</td>
</tr>
<tr>
<td><strong>Genes in the predicted PerR regulon(^b)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DVU2247</td>
<td>\textit{ahpC}</td>
<td>Antioxidant, AhpC/Tsa family</td>
<td>2.02±0.220</td>
<td>2.03±0.186</td>
</tr>
<tr>
<td>DVU2318</td>
<td>\textit{rbr}</td>
<td>Rubrerythrin, putative</td>
<td>2.47±0.277</td>
<td>1.76±0.122</td>
</tr>
<tr>
<td>DVU3095</td>
<td>\textit{perR}</td>
<td>PerR, transcriptional regulator</td>
<td>1.88±0.213</td>
<td>1.90±0.133</td>
</tr>
<tr>
<td><strong>Other iron-related genes</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DVU0646</td>
<td>\textit{cobI}</td>
<td>Precorrin-2 C20-methyltransferase</td>
<td>1.28±0.096</td>
<td>2.35±0.182</td>
</tr>
<tr>
<td>DVU0647</td>
<td>\textit{COG-}feoB</td>
<td>Iron compound ABC transporter, iron-binding protein</td>
<td>0.93±0.071</td>
<td>2.11±0.171</td>
</tr>
<tr>
<td>DVU0648</td>
<td>\textit{feoC}</td>
<td>Iron compound ABC transporter, ATP-binding protein</td>
<td>1.17±0.277</td>
<td>1.84±0.132</td>
</tr>
<tr>
<td>DVU0649</td>
<td>\textit{COG-}feoD</td>
<td>Iron compound ABC transporter, permease protein</td>
<td>1.20±0.076</td>
<td>2.26±0.119</td>
</tr>
<tr>
<td>DVU1397</td>
<td>\textit{bfr}</td>
<td>Bacterioferritin</td>
<td>2.27±0.217</td>
<td>2.16±0.212</td>
</tr>
<tr>
<td>DVU1568</td>
<td>\textit{ftn}</td>
<td>Ferritin</td>
<td>1.89±0.173</td>
<td>2.33±0.222</td>
</tr>
</tbody>
</table>

\(^a\) SSDR was calculated from Cy5-labeled cDNA signal while Cy3-labeled gDNA was used for both JW707 and WT; \(^b\) Those are predicted by Rodionov et al. (25); \(^c\) Mean±SD (n=6); \(^d\) Not determined due to the lack of Cy5 signal of the \textit{Afur} mutant.
Table 5. The number of detected, unique and overlap spots among replicate A, B, and C. Three different methods, SSR, SBR and SSDR and their pre-determined thresholds were used for the detection of positive spots.

<table>
<thead>
<tr>
<th></th>
<th>SSR</th>
<th>SBR</th>
<th>SSDR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold</td>
<td>2.0</td>
<td>1.6</td>
<td>0.80</td>
</tr>
<tr>
<td>Detected positive spots (mean ± sd)</td>
<td>3858±157</td>
<td>4372±322</td>
<td>3828±60</td>
</tr>
<tr>
<td>Total positive spots (A U B U C)</td>
<td>4132</td>
<td>4743</td>
<td>3903</td>
</tr>
<tr>
<td>a. Unique positive spots among three replicates</td>
<td>232 (5.6%)</td>
<td>566 (12%)</td>
<td>97 (2.5%)</td>
</tr>
<tr>
<td>b. Overlapped positive spots among two replicates [(A ∩ B) U (A ∩ C) U (B ∩ C)]</td>
<td>263 (6.4%)</td>
<td>521 (11%)</td>
<td>45 (1.2%)</td>
</tr>
<tr>
<td>c. Overlapped positive spots among three replicates (A ∩ B ∩ C)</td>
<td>3637 (88%)</td>
<td>3656 (77%)</td>
<td>3761 (96.3%)</td>
</tr>
</tbody>
</table>
**Table 6.** A summary of the ranges of experimentally determined SNR threshold under low and high stringent conditions using the *S. oneidensis* MR1 gDNA target.

<table>
<thead>
<tr>
<th></th>
<th>Low stringency</th>
<th></th>
<th>High stringency</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No FN</td>
<td><strong>Optimal</strong></td>
<td>No FP</td>
<td>No FN</td>
<td><strong>Optimal</strong></td>
</tr>
<tr>
<td>SSR</td>
<td>0.5</td>
<td>2.0</td>
<td>4.0</td>
<td>1.0</td>
<td><strong>2.5</strong></td>
</tr>
<tr>
<td>SBR</td>
<td>0.5</td>
<td>1.6</td>
<td>4.0</td>
<td>1.1</td>
<td><strong>1.8</strong></td>
</tr>
<tr>
<td>SSDR</td>
<td>0.3</td>
<td>0.7</td>
<td>0.9</td>
<td>0.4</td>
<td><strong>0.8</strong></td>
</tr>
</tbody>
</table>