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Evaluating the performance of new approaches to spot quantification and differential expression in 2-dimensional gel electrophoresis studies

Jeffrey S. Morris^{*},¹, Brittan N. Clark², Wei Wei¹, and Howard B. Gutstein²

¹Department of Biostatistics, The University of Texas M D Anderson Cancer Center, Houston, Texas USA

²Departments of Anesthesiology and Biochemistry, The University of Texas M D Anderson Cancer Center, Houston, Texas USA

Abstract

Spot detection and quantification for 2-DE are challenging and important tasks to fully extract the proteomic information from these data. Traditional analytical methods have significant weaknesses, including spot mismatching and missing data, which require time-consuming manual editing to correct, dramatically decreasing throughput and compromising the objectivity and reproducibility of the analysis. To address this issue, we developed *Pinnacle*, a novel, quick, automatic non-commercial method that borrows strength across gels in spot detection and has been shown to yield more precise spot quantifications than traditional methods. New commercial software, notably *SameSpots*, has also recently been developed as an improvement over traditional workflows. In this paper, we briefly describe *Pinnacle* and compare its performance to *SameSpots* in spot detection, spot quantification precision, and differential expression. Our analysis is performed in a rigorous fashion that, unlike other comparisons in the literature, summarizes performance across all spots detected on the gels, and we manually optimize *SameSpots* results while simply running *Pinnacle* with standard settings and no manual editing. While both methods showed marked improvement over a commercially available traditional method *PG240*, *Pinnacle* consistently yielded spot quantifications with greater validity and reliability, avoided spot delineation problems, and detected more differentially expressed proteins than *SameSpots*, and represents a significant non-commercial alternative for 2-DE processing.

Keywords

Two-dimensional polyacrylamide gel electrophoresis; Differential In-Gel Electrophoresis; False discovery rate; Differential expression; Bioinformatics; Spot detection; Image processing

Introduction

2-DE is an important method for proteomic analysis, allowing one to detect and measure thousands of proteins present within a biological sample. In 2-DE, proteins are physically separated within a polyacrylamide gel in two dimensions based on their *pI* and molecular

^{*}Corresponding Author: Department of Biostatistics, Division of Quantitative Sciences, The University of Texas M D Anderson Cancer Center, P O BOX 301402, Houston, TX 77230-1402, Phone: (713) 563-4284, Fax: (713) 563-4242, jefmorris@mdanderson.org.

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weight. Subsequently, the gel is scanned to produce a digital image, in which the proteins appear as spots with high staining intensities. Each gel image must be processed to detect and quantify the intensities of the protein spots. Given a study with N gels and p protein spots detected across all of the gels, after this processing we are left with an $N \times p$ matrix Y of spot intensities. For comparative proteomics studies, this matrix can then be surveyed using standard statistical methods to determine which protein spots are differentially expressed across treatment groups, or are otherwise related to whatever factor is of interest in that study.

It is of crucial importance to effectively detect, match, and quantify the spots across gels. It is also important to have adequate sensitivity for spot detection, so subsequent analyses survey all of the detectable proteins in the sample and do not miss proteins that are differentially expressed. Accurate spot matching is critical, as mismatched spots across gels result in different proteins being matched together in the analysis, compromising the ability to find significant differences. Precise spot quantifications are important to reduce variability and thus maximize power for detecting significant differences.

Most traditional methods for gel processing involve first detecting spots, estimating spot boundaries, and computing spot volumes for each individual gel, then matching the detected spots across gels. This approach is problematic because it results in missing data in the $N \times p$ matrix Y for any spots not detected on all of the gels. It is also prone to matching errors because cognate spots may be in slightly different positions on each gel, and matching is only done after detection. In addition, spot boundaries are difficult to accurately estimate, especially when spots overlap, and so the measurement error in the spot quantification is increased by methods that estimate spot boundaries separately for each individual gel. Most currently available software packages permit manual editing of these errors which can be very time consuming (1-4 h per gel [1][2]) and compromises the objectivity and reproducibility of the analysis. Most problematically, all of these problems worsen as the number of gels N in the study increases. This encourages researchers to conduct studies with fewer samples, leading to insufficient statistical power to detect significant differences when multiple testing is appropriately taken into account. These issues compromise the ability to conduct automatic, objective, high-throughput, 2-DE analyses using traditional approaches [2].

Some recently developed 2-DE processing workflows mitigate some of these problems. We have introduced a novel non-commercial method *Pinnacle* [3] for spot detection and quantification. *Pinnacle* operates on pre-aligned gel images, and starts by computing an average gel. Unlike the composite gels used in many commercial packages, this average gel is an arithmetic average, which allows one to borrow strength across gels in performing spot detection. Wavelet denoising is then applied, and spots are detected on the average gel by identifying “pinnacles,” or staining intensities that are local maxima in both the horizontal and vertical dimensions. Quantifications for each individual gel are taken as the maximum pixel intensity in the neighborhood of the (x, y) coordinate for each pinnacle detected in the average gel. This method is quick and automatic, results in no missing data, and the spot detection accuracy should improve as the number of gels in the study increase. The use of pinnacles instead of spot volumes for quantification eliminates the need to estimate spot boundaries, resulting in improved reliability and validity [3].

After we developed *Pinnacle* and performed the studies in that paper, Nonlinear Dynamics (Newcastle upon Tyne, UK) released a new commercial 2-DE analysis product, *Progenesis SameSpots*, which provides another alternative processing workflow to the traditional approaches. The basic workflow has some similarities to *Pinnacle*: align the images before processing to avoid the problems resulting from spot matching after detection, and perform spot quantification in a more uniform way across gels to increase quantification precision. Also working on aligned gels, *SameSpots* starts by performing spot detection simultaneously across

all of the images, obtaining a master list of detected spots and corresponding spot boundaries. Unlike *Pinnacle*, it does not compute an average gel or otherwise analytically borrow strength across gels in performing spot detection. Spot boundary masks are then overlaid on the aligned gels, and used to compute individual spot volumes, producing the $N \times p$ matrix Y . This method also results in no missing data, and its use of common spot boundaries for all of the gels should in principle increase the precision of spot volume estimates. Karp and colleagues [4] recently confirmed that the use of *SameSpots* reduced variability and in principle should increase power to detect differential expression when compared with traditional methods, and, in theory, should diminish the need for subjective editing. Note that unlike *Pinnacle*, which quantifies spots using pinnacle intensities, *SameSpots* quantifies spots using volumes after estimating spot boundaries.

Our previous study [3] compared the performance of *Pinnacle* with several available commercial packages, but all used the traditional workflow, which was all that was available at that time. In this study, we assess the performance of *Pinnacle* and *SameSpots*, representing a widely used commercial method with an alternative workflow. Our comparisons are done on four 2-DE data sets: two dilution series and two differential expression studies. Spot detection was evaluated by summarizing the number of spots automatically detected by each method, and by visually assessing the accuracy of detection and spot splitting. We separately investigated the two components of precision: *validity* and *reliability*. Validity was assessed by computing the linearity of the relationship between the spot quantification for detected spots and the protein loads in the two dilution series. Reliability was assessed by computing the % CV within each protein load group in the dilution series and within treatment groups in the comparative proteomics studies. Differential expression was evaluated by summarizing the number of spots in the comparative proteomics studies with p-values less than 0.05, 0.01, 0.005, and 0.001, and with local false discovery rates less than 0.15 and 0.20.

Our primary comparison is based on an automatic application of these methods to aligned gels with no manual editing. We also attempted to optimize *SameSpots* by evaluating alternative spot detection strategies and performing extensive manual editing of the spot detection and boundaries in *SameSpots* and compared these results. Using a graphical interface we are developing, it is quick and easy to edit *Pinnacle*'s spot detection results while viewing the average gel if desired, but in this paper we only assess the automatic *Pinnacle* results with standard values for tuning parameters and no manual editing. To assess both of these methods relative to more traditional workflows, we also compared *Pinnacle* and *SameSpots* with Progenesis *PG240* in terms of validity and reliability of spot quantification using the dilution series. We did not include *PG240* in the differential expression comparisons, as it is unclear how to properly handle the missing data in that context, and using standard ad hoc approaches to deal with the missing data, we have already compared *Pinnacle* and *PG240* in terms of differential expression in our previous study [3]. Unlike existing studies done in 2-DE, each analysis was done on all of the spots detected on the gels, not just a subset, to ensure that we provided a thorough assessment of the algorithms' performance.

Methods

Data Sets

Nishahara and Champion Study (NC)—These data were provided to us courtesy of Dr. Kathleen Champion-Francissen. The generation of this data set was described previously [5]. Briefly, these data consist of 28 gels from a dilution series using a sample of *E. coli*, with four replicate gels from each of seven different protein loads spanning a 100-fold range (0.5 μ g, 7.5 μ g, 10 μ g, 15 μ g, 30 μ g, 40 μ g, and 50 μ g).

SH-SY5Y Neuroblastoma Cell Dilution Series (SH)—These data were obtained from a dilution series performed by our lab. Details of the 2-DE, staining, and image capture were described previously [3]. These data consist of 18 gels from a dilution series created from SH-SY5Y neuroblastoma cells, with three replicate gels from each of 6 protein loads (5 μ g, 10 μ g, 25 μ g, 50 μ g, 100 μ g, 150 μ g).

D1 Differential expression study—These data were obtained by treating C6 glioma cells stably expressing the mu opioid receptor (mansour) with 1 μ M of the agonist fentanyl or an equivalent volume of water for 12 hours. Four independent samples of cells in each treatment group were then collected and extracted and a sequential extraction was performed using the ProteoPrep (Sigma) kit as previously described (mouledous), except that after hypotonic lysis the remaining pellet was extracted in 2% CHAPS, 7M urea, 2M thiourea, and 20 mM Tris buffer. Protein loads were equalized and IEF was performed in a running buffer consisting of the above plus 30% (v/v) trifluoroethanol (TFE). Second-dimension SDS-PAGE, staining and image capture were performed as previously described [3].

D2 Differential expression study—C6 glioma cells as described above were also used to generate this data set. However, after drug treatment, cells were pelleted and 50 mM ammonium bicarbonate added at a 1:5 (w/v) ratio, and the pellet gently resuspended. TFE/Chloroform at a ratio of 2:1 (v/v) was then added to the sample in a ratio of 7:1 (v/v). The sample was then vortexed vigorously, sonicated on ice, and incubated on ice for one hour with gentle vortexing. The TFE fraction was collected and lyophilized. Proteins were then resuspended in 1% C7BzO, 7 M urea, 2M thiourea, and 30% (v/v) TFE for IEF. Second-dimension SDS-PAGE, staining and image capture were performed as previously described [3]. In this data set, there were 5 fentanyl gels and 4 control gels.

Processing Methods

Alignment and Cropping of Gels—All analyses were performed on gel images that had been aligned so that cognate spots are at corresponding locations across different gels, and cropped to remove the boundary regions without proteins. For the two dilution series, a reference gel was chosen, and all of the gel images in the set were aligned to the reference gel using the commercial software package TT900 (Nonlinear Dynamics, Newcastle Upon Tyne, UK), which provides an interface for the user to manually align cognate spots on each gel to the reference gel. This alignment process took approximately 30-60 minutes per gel. TT900 was originally purchased as a stand-alone package, but is now integrated with *SameSpots*. After alignment, we read the gel images into Matlab (Mathworks, Inc., Natick, MA USA) for cropping and resaved the cropped, aligned images as .tif files. These files were used in all subsequent analyses. For the differential expression studies, a preliminary alignment was done using the automatic 2d gel alignment package RAIN [6], which was then manually refined and improved as necessary using TT900, which took far less time (5-10 minutes/gel) than using TT900 alone. These gels were also read into Matlab, cropped, and saved as .tif files.

Pinnacle—Here we briefly summarize the *Pinnacle* method and describe its key features. Working on aligned images, *Pinnacle* first computes an “average gel,” which involves averaging the intensities pixel by pixel across all of the gels in the experiment. Next, *Pinnacle* applies wavelet denoising methods to denoise the average gel, followed by spot detection, which involves identifying “pinnacles” on the wavelet-denoised average gel. A pixel is said to be a pinnacle if it is a local maximum in both the horizontal and vertical directions, and has intensity greater than a specified threshold. By default, the threshold is the 75th percentile on the gel, although the user can change this parameter, if desired. Unlike most other methods, spot quantification in *Pinnacle* is not done by estimating spot boundaries and computing spot volumes, but instead the quantification is taken to be the pixel intensity at the

pinnacle location. Specifically, the spot quantification for each individual gel is obtained by taking the maximum intensity within a rectangular window surrounding the pinnacle locations in the average gel. This window size is specified by the user, and by default, is 5 pixels by 5 pixels, accommodating slight remaining misalignment of spots between gels, but being small enough to avoid including pinnacles from any other cognate spots. Background correction can be done either globally or locally. We have found local background correction to work better for the data we have analyzed. Local background correction was done for each pinnacle by subtracting the minimum pixel intensity in a rectangular window surrounding the pinnacle. We find that large windows work well for background correction, so as to ensure that the window includes regions of the gel that are truly background, i.e. without protein content. By default, the window size for local background correction is 200×200 , which is what we use here (for 1024×1024 images). *Pinnacle* can be used for DIGE data by treating the image from each channel as a separate image in the detection process, and then taking the simple ratio (or difference on the log scale) of corresponding pinnacle intensities across the two channels for quantification. No other background correction is necessary for DIGE.

Four key aspects of *Pinnacle* are that it (1) uses an average gel for spot detection, (2) uses wavelet denoising to remove noise from the average gel image before detection, (3) detects spots by their pinnacles, and (4) quantifies spots by pinnacle intensities rather than spot volumes. The use of the average gel for spot detection has several favorable characteristics. It provides unambiguous spot detection since a pinnacle is a well-defined quantity. Second, it allows borrowing of strength across gels in determining what is a spot, since spot signals are reinforced across gels, while noise and artifacts are down-weighted. This, in principal, should lead to improved sensitivity and specificity for spot detection, and spot detection accuracy that increases with sample size. A spot need not be present in all of the gels to be identified in the average gel: given N gels, by the central limit theorem of statistics, we expect that the background noise should be reduced by a factor of \sqrt{N} , leading to a higher signal-to-noise ratio on the average gel than individual gels (and thus greater sensitivity for spot detection) as long as the spot is present on at least $1/\sqrt{N}$ of the gels. In the context of MALDI-MS peak detection, this principle has been clearly demonstrated in a simulation study [7], and in the future we plan to validate this principle for 2-DE using simulation studies based on a 2-DE simulation tool we are developing. Another alternative that might lead to even more sensitivity (but also less robustness to artifacts) would be to use a weighted average gel [8].

Wavelet denoising has become a standard tool in denoising images in all areas of signal processing, including 2-DE [9][10][3], although *Pinnacle* is the first method to explicitly use the wavelet denoising to aid spot detection. This process smoothes out spurious bumps in the image that correspond to high-frequency noise without attenuating the dominant features of the gel too strongly, which allows sensitive pinnacle-based spot detection while reducing the number of spurious detected spots that are in fact noise. This can provide additional power in differential expression studies, since spurious spots increase the false discovery rate estimates for other spots, and thus reduce power to detect them.

The detection of spots by their pinnacles rather than their boundaries has the potential to improve the resolving of overlapping spots. Færgestad, et al. [11] outline the many shortcomings spot-based methods have in resolving overlapping spots. They point out that most spots on a gel overlap with another spot, and that frequently an “envelope” of overlapping spots is called a single spot by many spot detection methods, and in that case it can become nearly impossible to detect differential expression in one of the spots in the envelope. Further, they mention that in overlapping spots, the determination of boundaries between the spots is crucial, and the fact that these boundaries are estimated separately across gels introduces serious quantification errors into the data. By basing spot detection on the pinnacles, *Pinnacle* avoids many of these problems and permits the resolving of overlapping spots since

even spots with a great degree of overlap typically have their own unique pinnacles. This can be seen in Figure 1 of Færgestad, et al.'s paper [11] illustrating an example with an “envelope of proteins” detected as a common spot by convention boundary-based spot detection methods. It is clear from that image that if an average were taken over the gels, two pinnacles would stand out distinctly from one another and be detected and quantified as separate spots.

The quantification of spots using pinnacle intensities rather than spot volumes precludes the estimation of spot boundaries. This has the potential to increase the reliability of spot quantifications, as spot boundary estimation is a difficult and error-prone process, especially when spots overlap. Even though spot volumes are traditionally used for quantification, if a given spot has a common shape across all of the gels, pinnacle intensities and true spot volumes should be strongly correlated with each other [3][12]. For quantification, the problem of overlapping spots is that the intensity at each pixel consists of a mixture of all spots spanning that pixel. By quantifying using only the pinnacle intensity, *Pinnacle* focuses on the part of the spot with maximum contribution from the spot of interest, the pinnacle, which potentially reduces the impact of the neighboring spots relative to volume-based methods whose quantification sums all pixel intensities within the estimated spot boundaries.

SameSpots—Working with aligned gels, Progenesis *SameSpots* starts by performing spot detection simultaneously across all of the images, obtaining a master list of detected spots and corresponding spot boundaries. These spot boundary masks are then overlaid on the individual gels, and are used to compute spot volumes, producing the $N \times p$ matrix Y .

The images for the four datasets were processed as single-stain/gel experiments. For the dilution series, the reference gel was selected from the highest protein load group. The reference gel remained constant throughout all analyses. As described previously, the images were aligned using TT900, therefore we proceeded to “analyze with *SameSpots*” without any further modifications. We did not employ the “mask of disinterest,” a tool used to exclude areas of the gel from analysis, because we had already cropped those regions from the gel images prior to analysis. Gels were grouped by protein load for the dilution series, and by treatment group for the comparative proteomics studies.

For the primary analysis, we evaluated *SameSpots* as an automatic method, with no manual editing of spots or spot boundaries. All detected spots were selected for analysis using the “spot review tool.” For background correction, we used the default “lowest on boundary,” which computes the spot background as the minimum pixel intensity adjacent to the spot boundary. Previous versions of Progenesis recommended the “Progenesis background subtraction” algorithm. We also evaluated this background correction method, but found that it resulted in considerably larger %CVs than “lowest on boundary.” In order to preserve the relationship between protein loads and spot volumes used to assess validity, normalization was not applied to the dilution series, but was applied to the differential expression data. The spot volumes for p spots for each of the N gels were exported as an $N \times p$ spreadsheet for further analysis.

We found that the spot detection algorithm within *SameSpots* at times resulted in a smaller spot count than expected, with the merging of numerous visibly distinct spots. In an attempt to improve spot count, we reapplied spot detection on the selected reference gel using the conventional (previous generation) Progenesis platform. The new spot boundaries were applied using the “SameSpots” option available within Progenesis (PG) to quantify the remaining gels in the experiment. This approach mimics the step-wise approach in *SameSpots* by detecting proteins on the reference gel and then applying the spot boundary “mask” to all of the gels in the experiment. This increased spot detection by an average of 27.6%, and improved spot splitting. No manual editing was performed for the primary comparisons.

As a secondary analysis, we edited the spots (split, merged, added, selected for deletion) found by *SameSpots* using the “spot review tool.” All of the edits were done by one author (BNC) to minimize user-induced variation. Spot volumes were then exported into an Excel spreadsheet. We found that the user interface in the conventional portion of the Progenesis platform (PG) had enhanced contrast and zooming capabilities, which permitted a better visual assessment of the spots than the interface in *SameSpots*. When viewing the *SameSpots*-edited spot boundaries in PG, we noticed that some of the spot boundaries could be further improved by additional manual editing in PG. Necessary edits were applied, and the *SameSpots* option was used to transfer the new spot boundaries to quantify cognate spots in the remaining gels in the experiment. This editing process removed the default settings applied by *SameSpots* (i.e., clearing all normalization from the experiment and background values from each edited gel). Therefore, the “lowest on boundary” background subtraction was re-applied, and normalization was reapplied for the differential expression studies, but not the dilution series. Spot volumes were exported into an Excel spreadsheet. The time required for manual editing was recorded.

In summary, we obtained spot volumes from *SameSpots* using four different methods, *SameSpots* alone (SameSpots), *SameSpots* aided by conventional Progenesis spot detection (SameSpots+PG), *SameSpots* with manual editing of spots (SameSpots+edits), and *SameSpots* with further manual editing in the conventional Progenesis platform (SameSpots+edits (PG)).

Progenesis PG240—We also applied a previous version of Progenesis (*PG240*, version 2006, Build 2405.1), which is based on the traditional approach, to the NC and SH dilution series so that we could determine whether newer methods (*SameSpots* and *Pinnacle*) showed improvements over traditional algorithms. We applied *PG240* (without *SameSpots*) to the aligned, cropped gel images previously described. The analysis wizard was used to set up the experiments. Gels were grouped by dilution series. Spot criteria for the reference gel included all of the spots present on at least one gel. No manual editing of the spots was performed, and the spot volumes were exported into an Excel spreadsheet. We excluded from the quantitative analysis spots that were not present in 3 of 4 replicates for at least 1 protein load group in the NC study, or in 3 of 3 replicates for at least 1 protein load group in the SH study. Spot volumes of zero were substituted for spots present on other gels that did not have a match on the current gel.

Statistical Comparisons

For the dilution series, we assessed the two different components of precision, *reliability* and *validity*, separately. To assess reliability of spot quantifications, we computed the %CV across replicates for each detected spot and each protein load group for each method. We considered %CV a better measure than standard deviation for reliability, as the standard deviation is typically correlated to the mean, while the coefficient of variation is not. For the comparative proteomics study, we computed %CV for each treatment group. As different numbers of spots were detected by the different methods, we compared methods by summarizing the distribution of %CV across treatment groups, computing the mean and 5-number summary (5th percentile, 25th percentile, 50th percentile, 75th percentile, 95th percentile) for each protein load group. We also combined them over the protein load groups to get a single set of summary statistics for each study.

To assess validity, for each method we computed the coefficient of variation (R^2) for the linear regression of the spot quantification vs. the protein load for each detected spot in the respective dilution series. Without normalization, more reliable spot quantification methods should yield stronger linear relationships, and thus greater R^2 , between spot quantifications and protein

loads. This measure can also be interpreted as the square of the Pearson correlation coefficient between the spot quantifications and protein loads. We summarized again the distribution of R^2 across spots within an experiment using the mean and 5-number summary.

The measures R^2 and %CV provide a reasonable summary of the precision of a spot quantification method, and increased precision of spot quantifications should correspond to increased power in differential expression studies. However, they are not in themselves sufficient measures for summarizing performance. For example, neither measure penalizes a method for merging several distinct spots. That type of error will not decrease the R^2 because each distinct spot should have the same ratio across gels in a dilution series; we may even expect the R^2 to be higher and the %CV to be lower due to the reduced variability of any inherent technical noise from summing the volumes across spots. However, a spot quantification method that merges spots will tend to mask differential expression in comparative proteomics studies.

Thus, we also compared the methods in terms of differentially expressed proteins using two studies, both of which were expected to show evidence of some differentially expressed proteins.

For the differential expression studies, we first log-transformed spot quantifications to stabilize the variance [13] and then performed two-sample t-tests for each spot to detect differences in protein expression between treatment groups. We summarized the number of spots with p-values <0.05, 0.01, 0.005, and 0.001. When searching for differentially expressed proteins over several hundred or thousand spots, it is essential to do some sort of multiple comparison adjustment to ensure that the differences observed are not simply due to random chance alone [14]. For example, with 1000 spots, even if there is in fact no true difference between the protein expression in the treatment groups, by basic statistical principles it is expected that roughly 50, 10, 5, and 1 spots should have p-values less than 0.05, 0.01, 0.005, and 0.001, respectively. One way to adjust for multiple comparisons is to consider the false discovery rate (FDR), which is the probability that a spot determined to be significantly different is, in fact, a false discovery [14][15]. We applied the method *fdrtool* in R [16] to the p-values for each spot, and computed the corresponding q-values, which indicate the probability of a given spot being a false positive assuming it is called significant [17]. We summarized the number of spots with q-values <0.20 and 0.15, spots for which the probability of being a false discovery is <0.20 and 0.15, respectively, if flagged as differentially expressed.

Results

Spot Detection

NC Study—Using the traditional Progenesis *PG240* method, a total of 1422 spots were detected in at least one gel in the NC study. Of these, only 312 spots (22%) were detected on all of the gels. There were 996 spots detected on at least 3 of 4 replicates in at least one protein load group. These spots were used for validity and reliability analyses.

Progenesis *SameSpots* detected 688 spots, and *Pinnacle* found 937 spots (see Figure 1d). The low number of spots detected by *SameSpots* appeared to be due to a merging of distinct spots in dense regions of the gel (Figure 1a). This problem has been previously observed [4]. *Pinnacle* did not have trouble resolving overlapping spots or defining spots in regions of high density because even overlapping spots have distinct pinnacles.

When PG was used for spot detection after gel alignment by *SameSpots*, 1034 spots were detected (see Figure 1b). This improvement appeared to be due to improved splitting of spots. After 240 minutes of manual spot editing of the reference gel using *SameSpots*, the number of

spots detected increased from 688 to 900 spots (see Figure 1c). After 60 additional minutes of manual spot editing in PG after *SameSpots* alignment, the spot count increased from 900 to 1082 spots.

SH Study—For *PG240*, a total of 1344 spots were detected on at least one gel. Of these, just 99 were detected in all of the gels, and 1094 were detected in all 3 replicates for at least one protein load group. *SameSpots* detected a total of 1037 spots, and again spot merging appeared to be an issue. The number increased to 1305 spots when PG was used to perform spot detection on the back end. After 310 minutes of manual editing, *SameSpots* detected 1244 spots, and with an additional 180 minutes of manual editing in PG, detected 1383 spots. *Pinnacle* found a total of 1322 spots.

D1 Study—*SameSpots* detected 1094 spots, with 1319 detected when PG was used on the back end of *SameSpots* to enhance detection. After 180 minutes of manual editing in *SameSpots*, 1153 spots were detected, and after an additional 25 minutes of manual editing in PG, 1255 spots were detected. *Pinnacle* found a total of 719 spots with automatic settings and no manual editing.

D2 Study—*SameSpots* detected 1410 spots, with 1928 detected when PG was used on the back end. After 180 minutes of manual editing in *SameSpots*, 1390 spots were detected, and after 25 additional minutes editing in PG, 1458 spots were detected. *Pinnacle* found a total of 1136 spots with automatic settings and no manual editing.

Discussion—We found *SameSpots* at times experienced problems with merging distinct spots in dense regions of the gel. This problem was largely mitigated when detection was done in PG after alignment using *SameSpots*. Extensive manual editing was required to correct errors in spot splitting and detection. *Pinnacle* appeared to do a good job of detecting spots without any manual editing, although if desired, the user can edit spots after detection in *Pinnacle* or try different tuning parameters to optimize results. We did not apply this option, so all of the results include only the spot lists automatically obtained by the *Pinnacle* algorithm using standard tuning parameter settings.

Validity

Validity of spot quantifications was summarized by the R^2 of a linear regression of the spot quantifications on the protein loads. Not all of the methods identified the same spots; therefore, we summarized the results by their distributions across all of the detected spots for each method, as shown in Table 1.

NC Study—Boxplots of R^2 for detected spots are contained in Figure 2(a). For the 937 spots detected by *Pinnacle*, the mean R^2 was 0.965, which corresponds to a correlation of 0.982 between the spot quantifications and the true protein loads. Of the 688 spots detected by *SameSpots*, the mean R^2 was 0.956. With PG spot detection and no manual editing, the mean R^2 for the 1034 detected spots was 0.948. After manual editing, the mean R^2 across the 900 detected spots was 0.958. After spot editing using PG, the mean R^2 was 0.956. Comparing the distributions of R^2 by looking at the 5-number summaries and boxplots, we see that the differences between *Pinnacle* and *SameSpots* was strongest at the lower and middle part of the distribution, i.e. for the less reliable spots. *Pinnacle* and each of the *SameSpots* methods greatly outperformed the traditional *PG240*, which resulted in spot quantifications with a mean R^2 of 0.900 across the 996 detected spots, and with dramatic differences at the low end of the R^2 distribution.

SH Study—Boxplots of R^2 for detected spots are contained in Figure 2(b). For the 1322 spots detected by *Pinnacle*, the mean R^2 was 0.879. This was much higher than the mean R^2 for the 1037 spots detected by *SameSpots*, which was 0.804. When *SameSpots* alignment was followed by spot detection using PG, the mean R^2 for 1305 spots improved to 0.867. After manual editing, the R^2 across 1244 detected spots was 0.876. With additional manual editing in PG, the mean R^2 was 0.870. Comparing the distributions of R^2 between the methods, we see that the lower quantiles of R^2 are considerably larger for *Pinnacle* than the *SameSpots* methods, and the upper quantiles were similar, with the *SameSpots* methods slightly higher. Again, *PG240* performed much worse than either *Pinnacle* or any variation of *SameSpots*, with a mean R^2 of 0.698 across 1094 detected spots, and many spots with low R^2 values.

Discussion—The gels in the NC dilution series were exceptionally clean and well run. This was reflected by the excellent validity measures found for this data set for all of the methods. The SH dilution series may be more representative of gels commonly seen in practice. That dilution series resulted in smaller, but still reasonable, R^2 values. However, the relative performance of the different processing methods was similar for both data sets. Both *Pinnacle* and *SameSpots* provided spot quantifications with much higher validity than *PG240*. This may be due to the missing data problem inherent in the traditional approaches, and the extra measurement variability induced by estimating separate spot boundaries for cognate spots in each gel in *PG240*. The results produced by *SameSpots* were improved when manual editing was performed to correct a number of spots that were inaccurately detected or split. In both dilution series, *Pinnacle* appeared to compare favorably to the *SameSpots* methods, with higher mean R^2 , improved lower quantiles, and similar higher quantiles in the R^2 distribution.

Reliability

Reliability of spot quantifications was summarized by the %CV for each spot within each group. The different methods did not detect the same sets of spots; therefore, we summarized the reliability of spot quantifications by their distributions across all of the detected spots for each method. Summary values for each protein load group are found in the supplementary material, including the mean %CV, 5-number summary, plus the number and proportion of reliable spots, i.e. with %CV>20. The summary values combining the results for all of the protein load groups are found in Table 2.

NC Study—Boxplots of %CV for the detected spots are contained in Figure 2(c). The mean %CV for *Pinnacle* was 18.8; for *SameSpots* it was 20.2, and 22.7 when PG was used for spot detection. After manual editing, the mean %CV was 20.2 after *SameSpots* editing, and 20.9 after also editing with PG. Looking at the distribution of %CV, we see that *Pinnacle* tended to have slightly smaller %CV quantiles throughout the distribution. For *PG240* alone, the mean %CV across the detected spots was 49.7.

SH Study—Boxplots of %CV for detected spots are contained in Figure 2(d). The mean %CV for *Pinnacle* was 27.5; for *SameSpots* it was 29.9, and 30.4 when PG was used for spot detection. After manual editing, the mean %CV was 29.4 after *SameSpots* editing, and 30.0 after also editing with PG. Again, the differences between *Pinnacle* and *SameSpots* were consistent throughout the distribution of spot %CV. The mean %CV for *PG240* alone was 66.9.

D1 Study—The mean %CV for *Pinnacle* was 25.6; for *SameSpots* it was 43.8, and 44.0 when PG was used for spot detection. After manual editing, the mean %CV was 40.5 after *SameSpots* editing, and 40.3 after also editing with PG. The differences in %CV distribution were seen across the entire distribution of spot %CV.

D2 Study—The mean %CV for *Pinnacle* was 28.8; for *SameSpots* it was 36.8, and 36.7 when PG was used for spot detection. After manual editing, the mean %CV was 30.2 after *SameSpots* editing, and 30.3 after also editing with PG. Again, we see consistent differences across the entire range of %CV.

Discussion—For both dilution series, *SameSpots* and *Pinnacle* had much smaller %CV than *PG240*, which is likely due to the missing data and extra technical variability from estimating separate spot boundaries for cognate spots on each gel. The %CV for *SameSpots* improved after manual editing, but *Pinnacle* still had considerably smaller %CV than any of the *SameSpots* methods across all of the studies. In the supplemental tables split out by protein load, we see that use of *Pinnacle* tended to find a higher number and higher proportion of spots that were reliable, defined as %CV<20. We propose that the use of pinnacles rather than spot volumes in quantification is a key factor in the improved reliability. Increased reliability is important in differential expression studies, as lower %CV should lead to increased power to detect differentially expressed proteins.

Differential Expression

The differential expression results for the comparative proteomics studies are in Table 3.

D1 Study—Although *Pinnacle* with default tuning parameters found fewer overall spots for this data set (719 for *Pinnacle* vs. 1094-1319 for the various *SameSpots* methods), it found more spots with p-values <0.05 (76 vs. 11-24), <0.01 (17 vs. 1-3), <0.005 (6 vs. 0-1), and <0.001 (3 vs. 0). Flagging spots with local false discovery rates <0.15 or <0.20 as differentially expressed, *Pinnacle* found 2 or 3 differentially expressed proteins, respectively, while none of the *SameSpots* methods found any.

D2 Study—Of the 1136 spots detected, *Pinnacle* flagged 5 or 7 spots as significantly differentially expressed with a local FDR of <0.15 or 0.20, respectively, while none were found using any of the *SameSpots* methods. *Pinnacle* found more spots with p-values <0.001 (5 vs. 1-2) than the *SameSpots* methods. In this study, *SameSpots* methods found more spots with p-values <0.005 (11-16 vs. 10), <0.01 (24-35 vs. 20), and <0.05 (123-159 vs. 107), with the difference roughly proportional to the total number of spots detected (1410-1928 vs. 1136).

Discussion—Even with smaller spot counts in these two studies, compared to *SameSpots*, *Pinnacle* found more spots with very small p-values. After adjusting for multiple testing using FDR methods, *Pinnacle* found significant spots, whereas the *SameSpots* methods found none. These results were not surprising given that spot quantifications using *Pinnacle* had higher validity and reliability than any of the *SameSpots* methods.

Conclusions

The newer preprocessing methods, *SameSpots* and *Pinnacle*, both show tremendous improvement over *PG240*, a method based on a traditional analytical approach, in terms of validity and reliability of spot quantifications for 2-DE. Both *SameSpots* and *Pinnacle* avoid missing data, and reduce technical variability because they do not try to estimate different spot boundaries for corresponding spots on each gel, which is a difficult and error-prone step that increases variability. *SameSpots* still estimates spot boundaries, but estimates a single boundary for each spot that is applied to quantify that spot for all of the aligned gels. This eliminates missing data, and reduces technical variability in the spot quantifications.

Pinnacle bases the entire spot detection and quantification process on the pinnacles of the spots, and so avoids estimating spot boundaries altogether. This may seem controversial to some, as

spot volumes are often viewed as the “correct” quantifications for proteins in a gel. However, given the characteristics of pinnacles—they highly correlate with volumes [3], are much easier to estimate, introduce less technical variability, and are less influenced by overlap from neighboring spots—an analytic plan using pinnacles seems to be a reasonable approach.

As far as we know, there have been no definitive studies demonstrating that estimated spot volumes yield better quantifications than pinnacle intensities and, in fact, our results here and in our previous study [3] suggest the opposite. We have demonstrated that *Pinnacle* on average yields spot quantifications with higher validity than volume-based *SameSpots*, even after *SameSpots* results are improved by hours of manual editing. *Pinnacle*-based spot quantifications were also consistently more reliable than *SameSpots*, yielding lower %CV in four different data sets. The significance of these improvements in spot quantification precision was strikingly demonstrated by the differential expression analysis. Using *Pinnacle* resulted in spots flagged as differentially expressed using FDR-based methods, while even extensive manual editing of the *SameSpots* analysis did not yield significant differential expression in our two examples.

The spot detection algorithm utilized by the *SameSpots* interface in the Progenesis package appears, at times, to have difficulty splitting overlapping spots in dense regions of the gel. The negative effects of the merging of distinct spots were not reflected in their validity and reliability results. This is because the merged “superspot” would still have reasonable R^2 and %CV in the dilution series due to the spots having the same ratio of volumes between gels of differing protein loads. In the differential expression setting, in which we expect different expression ratios between distinct spots on different gels, such merging of spots would in principle mask differential expression if one but not all of the proteins in the “superspot” were in fact differentially expressed [11]. This might also partially explain *Pinnacle*'s better performance in the differential expression studies.

Since the completion of the studies reported in this paper, other companies have updated their 2-DE processing software to use a similar workflow as *SameSpots*, aligning the gels and using common spot outlines across gels for quantification, including REDFIN Solo (Ludesi; Malmo Sweden) and Delta2D (Decodon; Greifswald, Germany). These commercial packages were not yet available at the time this study was done, so were not compared in this study.

One situation in which we do not expect *Pinnacle*, or any other 2-DE analysis package, to work well is when pixel intensities on the gel image reach saturation, disrupting the linear relationship between pinnacle intensities and spot volumes, and potentially obscuring differential expression [18]. We avoid saturating any spots by adjusting the contrast settings when imaging gels, and we recommend that approach for all methods.

In conclusion, we have demonstrated that new analysis algorithms based on image alignment and standardized spot boundaries (or pinnacles) between gels markedly improve the robustness of analysis when compared with previous methods involving conventional spot detection and individual matching techniques. Thus, these new methods more accurately extract the proteomic information from the gel images, overcoming some of technical problems that in the past have prevented 2DGE and DIGE from realizing their full potential for biomedical research.

Pinnacle is a non-commercial method that differs from the available commercial packages in its use of pinnacle detection on wavelet-denoised average gels for spot detection, and its use of pinnacle intensities in lieu of spot volumes for quantification. In the studies presented here, *Pinnacle* demonstrated greater measurement precision, permitting the identification of many more differentially expressed spots, than prominent commercial software *SameSpots*, even after extensive hand editing was done to optimize the *SameSpots* results. Further, *Pinnacle* is

automatic, eliminating the potential for inadvertent bias, and did not experience the problems with spot-merging of overlapping spots that was sometimes seen with *SameSpots*. It represents a significant non-commercial alternative for preprocessing 2-DE images.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations Used

FDR	false discovery rate
SH	SH-SY5Y neuroblastoma cell dilution series
MALDI-MS	Matrix assisted laser desorption ionization time-of-flight mass spectrometry
NC	Nishihara and Champion dilution series
pI	isoelectric point
R ²	Coefficient of variation (squared correlation) for validity
2d	two-dimensional
2-DE	two-dimensional polyacrylamide gel electrophoresis
D1	First differential expression study
D2	Second differential expression study
%CV	Coefficient of variation (percentage) for reliability
DIGE	Differential In-Gel Electrophoresis

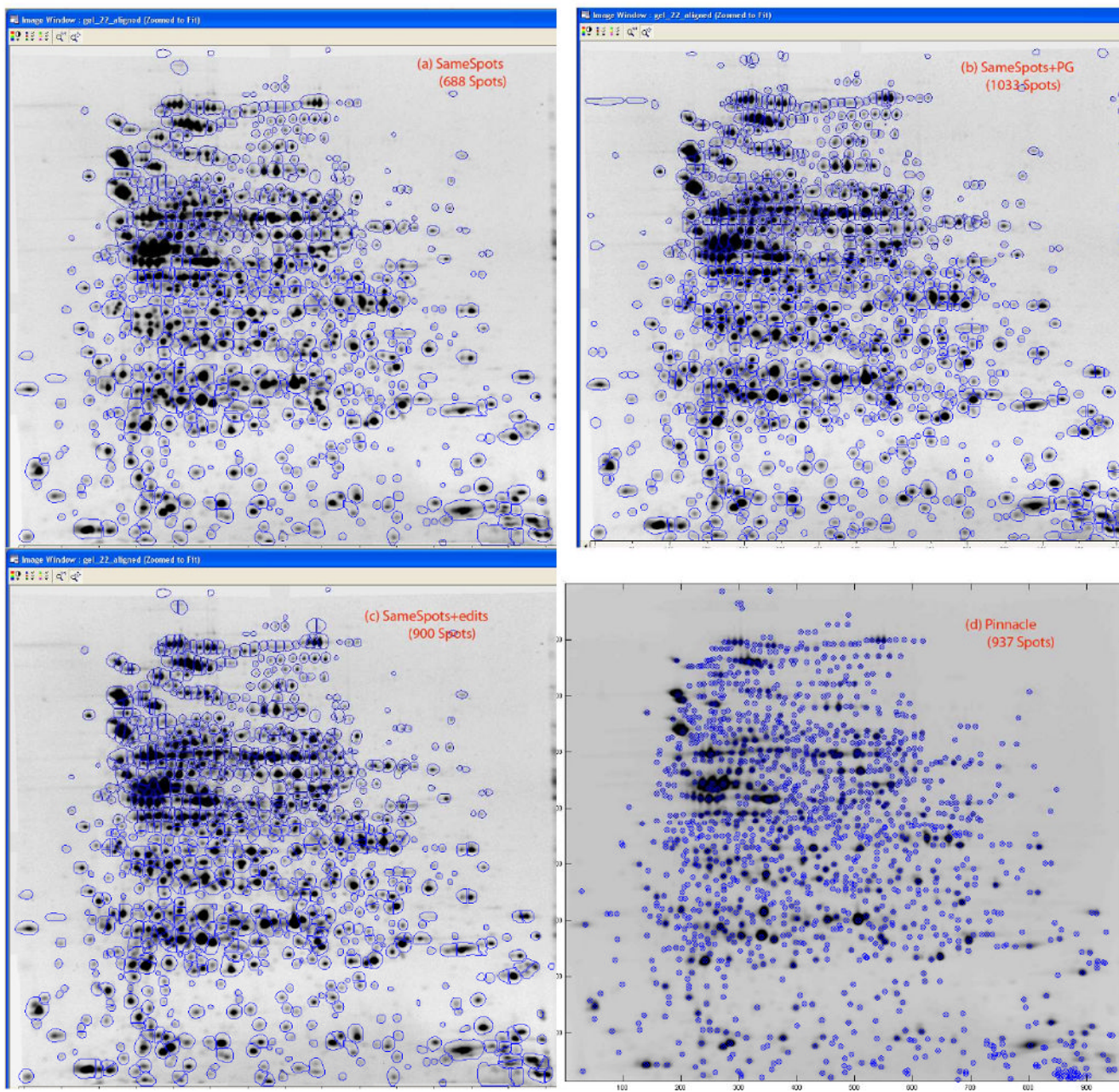


Figure 1. Spot Detection, NC Study

Results of spot detection on Nishahara and Champion study for (a) SameSpots, (b) SameSpots with PG spot detection, (c) SameSpots with manual editing, and (d) Pinnacle. For (a)-(c), the image is from gel 22; spots are marked by their estimated boundaries. For (d), the image is the average gel used for spot detection; pinnacles for detected spots are marked by a circled x.

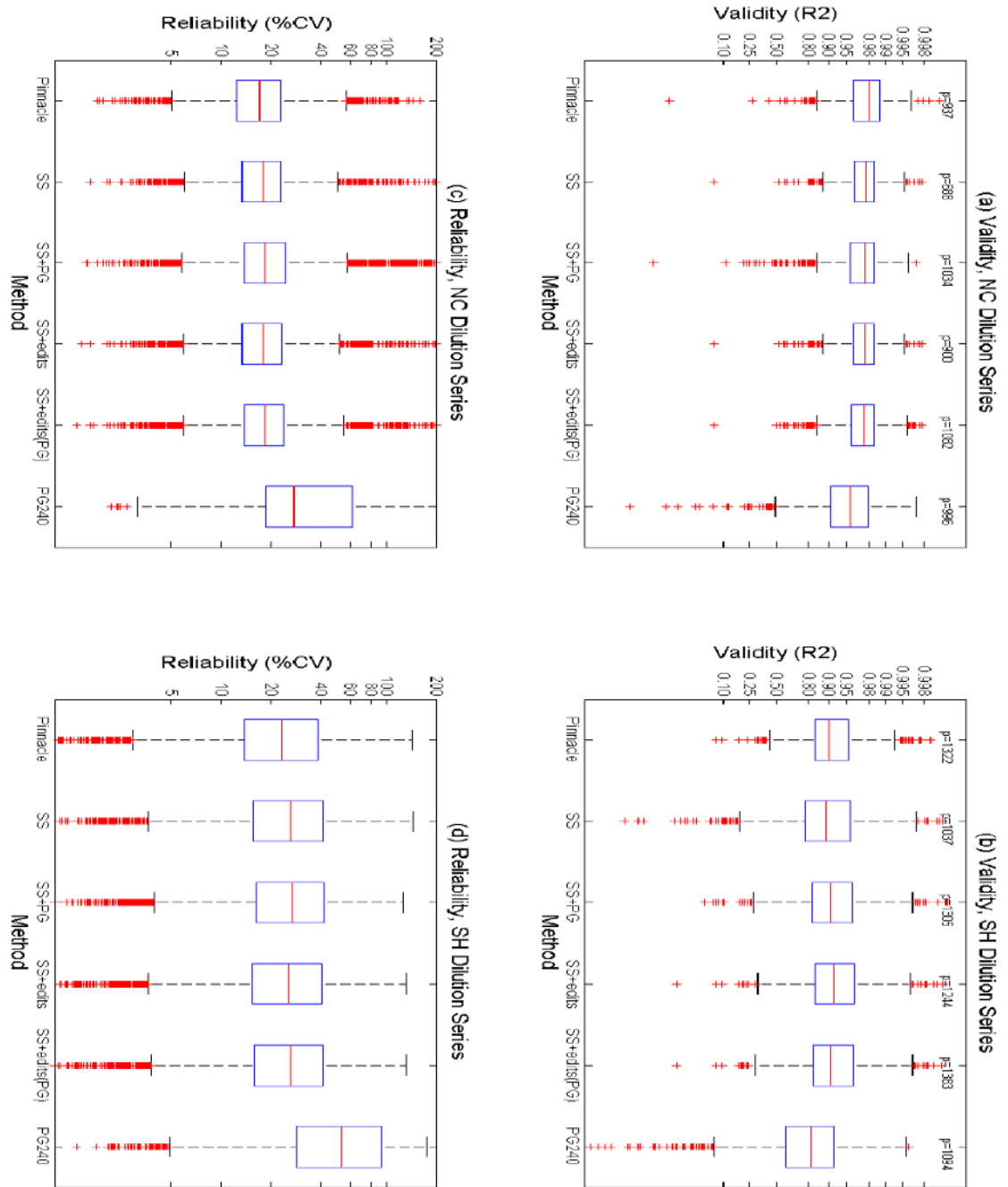


Figure 2. Boxplots of Validity and Reliability of Detected Spots, NC and SH Dilution Series
 Boxplot of (a) validity (R^2), Nishahara and Champion Study (NC), (b) validity (R^2), SH-SY5Y Neuroblastoma Cells Dilution Series (SH), (c) reliability (%CV), NC study, (d) reliability (%CV), SH study, for all spots detected by various methods, including Pinnacle, SameSpots (SS), SameSpots+PG (SS+PG), SameSpots+edits (SS+edits), SameSpots+edits-PG (SS+edits-PG), and PG240, with the numerical results plotted on a log axis for clarity of presentation.

Table 1

Validity of Quantifications for Detected Spots

Summary of R^2 measuring linearity of quantification method across protein loads within dilution series for all spots automatically detected by the various preprocessing methods, including (1) Pinnacle, (2) SameSpots, (3) SameSpots with PG spot detection (SameSpots +PG), (4) SameSpots with manual editing (SameSpots+edits), (5) SameSpots with manual editing in PG (SameSpots+edits (PG)), and (6) Progenesis PG240 (PG240).

Study	Method	Number of Spots	R^2 (Validity)					
			Mean	Q05	Q25	Median	Q75	Q95
NC	Pinnacle	937	0.965	0.899	0.966	0.980	0.987	0.992
	SameSpots	688	0.956	0.889	0.934	0.977	0.984	0.992
	SameSpots +PG	1034	0.948	0.818	0.959	0.976	0.984	0.992
	SameSpots+edits	900	0.958	0.892	0.963	0.977	0.984	0.992
	SameSpots+edits (PG)	1082	0.956	0.878	0.960	0.976	0.984	0.992
	PG240	996	0.900	0.571	0.909	0.957	0.980	0.990
SH	Pinnacle	1322	0.879	0.734	0.840	0.900	0.954	0.986
	SameSpots	1037	0.804	0.087	0.780	0.895	0.959	0.990
	SameSpots +PG	1305	0.867	0.598	0.821	0.910	0.962	0.991
	SameSpots+edits	1244	0.876	0.622	0.836	0.917	0.964	0.990
	SameSpots+edits (PG)	1383	0.870	0.610	0.827	0.909	0.962	0.990
	PG240	1094	0.698	0.039	0.601	0.815	0.920	0.979

Table 2

Reliability of Quantifications for Detected Spots

Summary of %CV for spots detected by various preprocessing methods, across all dilution series and treatment groups for differential expression studies. Methods include (1) Pinnacle, (2) SameSpots, (3) SameSpots with PG spot detection (SameSpots+PG), (4) SameSpots with manual editing with SameSpots (SameSpots+edits), (5) SameSpots with manual editing with PG (SameSpots+edits (PG)), (6) Progenesis PG240 (PG240).

Study	Method	Number of Spots	%CV (Reliability)						
			Mean	Q05	Q25	Median	Q75	Q95	
NC	Pinnacle	937	18.8	7.4	12.5	17.1	22.8	33.9	
	SameSpots	688	20.2	8.2	13.4	17.9	22.8	36.2	
	SameSpots+PG	1033	22.7	8.0	13.7	18.5	24.4	44.4	
	SameSpots+edits	900	20.2	7.9	13.4	17.9	23.0	35.7	
	SameSpots+edits (PG)	1082	20.9	7.8	13.7	18.3	23.9	38.1	
	PG240	566	49.7	10.5	18.6	27.4	61.5	200.0	
SH	Pinnacle	1322	27.5	5.2	13.7	23.2	38.3	61.0	
	SameSpots	1037	29.9	5.9	15.5	26.4	41.1	64.4	
	SameSpots+PG	1305	30.4	6.3	16.2	27.1	41.8	64.9	
	SameSpots+edits	1244	29.4	5.7	15.4	25.6	40.5	64.0	
	SameSpots+edits (PG)	1383	30.0	6.1	15.9	26.3	41.2	65.1	
	PG240	391	66.9	11.1	28.4	53.0	92.3	173.2	
D1	Pinnacle	719	25.6	11.0	16.5	22.4	30.1	53.0	
	SameSpots	1094	43.8	22.5	30.7	37.8	51.6	85.6	
	SameSpots+PG	1319	44.0	23.6	31.5	38.3	51.1	79.2	
	SameSpots+edits	1153	40.5	22.5	31.0	36.8	46.6	69.3	
	SameSpots+edits (PG)	1255	40.3	22.5	31.0	36.8	46.5	68.5	
	PG240	1136	28.8	10.4	16.2	21.9	31.4	75.0	
D2	Pinnacle	1410	36.8	14.7	21.6	29.1	44.3	86.0	
	SameSpots	1928	36.7	15.3	23.4	32.2	44.7	72.0	
	SameSpots+PG	1390	30.2	14.3	20.7	27.2	35.7	56.0	
	SameSpots+edits	1458	30.3	14.2	20.8	27.5	36.1	55.6	
	SameSpots+edits (PG)								
	PG240								

Table 3
Differential Expression Comparison, Comparative Proteomics Studies

Summary of performance of preprocessing methods in detecting differential expression in comparative proteomics studies. The p-values are from two-sample t-tests performed on the log-transformed spot quantifications for the various methods. The q-values are local FDR estimates, and measure the probability that a given spot would be a false positive if it is declared differentially expressed.

Method	Number of Spots	q-values		p-values			
		<0.15	<0.20	<0.001	<0.005	<0.01	<0.05
D1	Pinnacle	2	3	3	6	17	76
	SameSpots	0	0	0	0	1	17
	SameSpots +PG	0	0	0	1	3	24
	SameSpots +edits	0	0	0	0	1	11
D2	SameSpots +edits (PG)	0	0	0	0	1	13
	Pinnacle	5	7	5	10	20	107
	SameSpots	0	0	2	11	24	123
	SameSpots +PG	0	0	2	16	35	151
SameSpots +edits		0	0	1	14	32	157
	SameSpots +edits (PG)	0	0	0	12	27	159