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Quantitative Western Blot Analysis with Replicate Samples

LI-COR

Abstract

Replicate measurements are critical for quantitative Western blot (QWB) analysis. The purpose of QWB is to monitor changes in the relative abundance or modification of a target protein within a group of samples. Does the experimental treatment cause an increase or decrease in target abundance, compared to the untreated or control condition?

Replicate samples confirm the validity of observed changes in protein levels. Without replication, it is impossible to know if an effect is real or simply an artifact of experimental noise or variation. Biological and technical replicates are both important, but each type of replicate addresses different questions [1,2,3].

Types of Replicate Measurements

Technical replicates are repeated measurements used to establish the variability of a protocol orassay, and determine if an experimental effect is large enough to be reliably distinguished from the assay noise [1].

- Examples may include loading of multiple lanes with each sample on the same blot, running multiple blots in parallel, or repeating the blot with the same samples on different days.
- Technical replicates evaluate the precision and reproducibility of an assay, to determine if the observed effect can be reliably measured. When technical replicates are highly variable, it is more difficult to separate the observed effect from the assay variation. You may need to identify and reduce sources of error in your protocol to increase the precision of your assay. Technical replicates do not address the biological relevance of the results.

Biological replicates are parallel measurements of biologically distinct and independently generated samples, used to control for biological variation and determine if the experimental

effect is biologically relevant. The effect should be reproducibly observed in independent biological samples. Demonstration of a similar effect in another biological context or system can provide further confirmation.

- Examples include analysis of samples from multiple mice rather than a single mouse, or from multiple batches of independently cultured and treated cells.
- To demonstrate the same effect in a different experimental context, the experiment might be repeated in multiple cell lines, in related cell types or tissues, or with other biological systems.

An appropriate replication strategy should be developed for each experimental context. Several recent papers discuss considerations for choosing technical and biological replicates [1,2,3].

Reporting Changes in Relative Abundance

QWB methods typically use ratiometric analysis to determine relative abundance of the target protein and compare relative protein levels across a group of samples.

Ratiometric analysis is a form of 'self-calibration' that expresses the relative abundance of target protein as a ratio between the experimental sample and the control. It compares the intensity of the target band in each experimental sample to the intensity of the target band in the control sample [4,5,6].

- Normalize your QWB data using an appropriate internal loading control (such as total protein staining).
- Normalization mathematically corrects for small, unavoidable variations in sample loading and transfer by comparing the target protein to an internal loading control. More information is found in Guidelines.

- See these resources to learn more about QWB normalization:

Western Blot Normalization Handbook (licor.com/handbook)

Western Blot Normalization: Challenges and Considerations White Paper

(licor.com/normalizationreview)

- Perform the ratiometric analysis, as described in Guidelines.
- The normalized signal intensity of the target band in each sample should be divided by the normalized intensity of the target band in the control sample.

The resulting ratios, expressed as **fold change** or **percentage (%) change**, are used to compare relative protein levels across the samples on your blot. Because all samples are compared to the control, these measurements are proportional and are independent of raw signal intensity.

Fold change is a unitless value that compares the relative abundance of a target protein to the control sample on the same membrane.

• A value above 1.0 indicates an increase in abundance relative to the control; a value below 1.0 indicates decreased abundance (Table 1).

Fold change =
$$\frac{\text{normalized signal}_{\text{SAMPLE}}}{\text{normalized signal}_{\text{CONTROL}}}$$

Percentage (%) change is a unitless value similar to fold change that expresses changes in relative abundance as a percentage.

• A positive percentage indicates increased abundance relative to the control; a negative percentage indicates decreased abundance (Table 1).

normalized signal_{CONTROL} Table 1. Relative abundance of target protein, expressed as fold change and % change. Fold Change % Change 1.0 0% 1.2 +20% 15 +50% 2.0 +100% 8.0 -20% 0.5 -50% -90% 0.1

Citation: LI-COR Quantitative Western Blot Analysis with Replicate Samples. protocols.io

dx.doi.org/10.17504/protocols.io.k6sczee

Published: 30 Jan 2018

Guidelines

Keys to Success

 Accurate quantitative Western blot (QWB) analysis requires a robust, well-characterized detection system. Data should be captured within the linear range of detection, with careful attention to limits of detection and saturation of strong bands. Methods should be sufficiently robust that the outcome is unaffected by trivial or unintended changes in the reagents and protocol [7-8].

More Info: This protocol provides guidelines for characterizing the linear range of detection for your system: Determining the Linear Range for Quantitative Western Blot Detection (<u>licor.com/LinearRange</u>)

- Relative changes in abundance can only be measured if they substantially exceed the variability of your measurement. See Section VI, Data Interpretation, for guidelines.
- Do not compare signal intensities between blots. Experimental variations in sample loading transfer conditions, antibody reagents, incubation times, and many other factors affect the raw signal intensities on each blot. Band intensity should only be analyzed by relative, ratiometric comparison to the control samples.
- Every blot should include control samples (untreated, baseline, etc.) and be processed for detection with the same reagents, protocol, and imaging method. Fold change values can then be compared between blots loaded with the same control samples.

Experimental Design

- 1. Plan an appropriate strategy for technical and biological replicates in your experiment. It may be helpful to consult a statistician before you begin to generate data.
- 2. Collect replicate data that represent the desired experimental conditions (drug treatments, time course, dose-response, etc). A minimum of three replicates should be performed for each sample, including the controls.

Important: Samples should be loaded in **random order** on each blot to minimize position effects. Sample placement on the blot (for example, in edge lanes vs. interior lanes) may affect signal intensity and introduce variability. Random sample placement will reduce the impact of position effects [4].

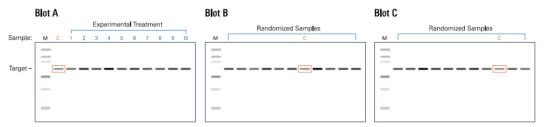


Figure 1. Replicate blot layout described in this protocol. Each gel is loaded with MW marker (M), control sample (C), and experimental samples (1-10). On blots B and C, control and experimental samples are loaded in random order.

- 3. Use this protocol to analyze and compare replicate samples on separate Western blots.
- a. This protocol describes the analysis of three replicate blots (designated A, B, and C; see Figure 1), each loaded with a molecular weight marker, control sample, and 10 experimental samples.
- b. This layout can be used to analyze either technical or biological replicates, depending on the nature of the samples you load. Appropriate interpretation of the resulting data will be

specific to your samples and experimental context.

• Technical replicates: Load the same group of control and experimental samples repeatedly, onto three (or more) separate gels, to generate replicate blots.

• Biological replicates: Perform three (or more) independent experiments. Load each set of control and experimental samples onto its own separate gel, to generate replicate blots.

Quantitative Western Blot Normalization

1. Normalize your Western blot data, using an appropriate internal loading control.

a. Normalization mathematically corrects for small, unavoidable sample-to-sample and laneto- lane variation by comparing the target protein to an internal loading control. Internal loading controls are endogenous reference protein(s) that are present in all samples at a stable level, and unaffected by the experimental conditions or treatments. The loading control serves as an indicator of sample protein loading, and is used to verify that observed changes in target protein abundance represent actual differences in the protein samples.

b. Common internal loading controls include total protein staining, "housekeeping proteins" (such as actin, tubulin, or GAPDH), and analysis of protein modifications with pan- and modification-specific primary antibodies.

• Total protein staining: The membrane is treated with a total protein stain to assess sample protein loading in each lane. After immunoblotting, the combined signal of all sample proteins in each lane is used as a loading indicator for normalization. This method is recommended by the Journal of Biological Chemistry [9].

Protocol: REVERT[™] Total Protein Stain Normalization (licor.com/RevertNormalization)

Housekeeping protein (HKP): This popular method employs a single, unrelated endogenous
protein as a readout of sample loading. Because a single indicator is used, changes in HKP
expression will introduce error. Validation is required to verify that HKP expression is constant in
all samples and unaffected by experimental conditions.

Protocol: Housekeeping Protein (HKP) Validation (<u>licor.com/HKP-Validation</u>)

Protocol: Housekeeping Protein Normalization (licor.com/HKP-Normalization)

 Pan- and modification-specific antibodies: This specialized form of normalization is used to analyze protein phosphorylation and other post-translational modifications. A modificationspecific primary antibody is multiplexed with a pan-specific antibody that recognizes the unmodified target protein. Signals are normalized to the actual level of target in each lane, using the target protein as its own internal control.

Protocol: Pan/Phospho Analysis for Western Blot Normalization (<u>licor.com/PanProteinNormalization</u>)

2. Use the normalized band intensity values for analysis of technical or biological replicates, as in the next Section.

Calculations for Replicate Analysis

Please see steps for detailed Calculations for Replicate Analysis.

Data Interpretation

Important: General guidelines are presented in this protocol. Appropriate replication and data interpretation are specific to each experiment, and beyond the scope of this protocol. Consult your local statistician for assistance.

- 1. Use the fold change and % change values for relative comparison of samples.
- 2. % change and CV can be used to evaluate the robustness of your QWB results, and determine if the magnitude of observed changes is large enough to be reliably distinguished from assay variability.
- a. As described in Section II, % **change** expresses the fold change as a percentage.
 - A positive number indicates an increase in relative abundance of target protein.
 - A negative number indicates decreased abundance.

- b. The **coefficient of variation (CV)** describes the spread or variability of measured signals by expressing the standard deviation (SD) as a percent of the mean. Because CV is independent of the mean and has no unit of measure, it can be used to compare the variability of data sets and indicate the precision and reproducibility of an assay.
 - A low CV value indicates low signal variability and high precision of measurement.
 - A larger CV indicates greater variation in signal and reduced precision.
- 3. On a Western blot, a change in band intensity is meaningful only if the magnitude of the % change substantially exceeds the CV (Table 3).
- a. Generally speaking, the magnitude of the reported % change should be at least 2X greater than the CV of that measurement.

For example, to report a 20% difference between samples (0.8-fold or 1.2-fold change in band intensity), CV < 10% would be recommended for replicate samples. For a specific measurement, this threshold for the magnitude of change would correspond to the mean \pm 2 SD.

- In Table 3, the change observed in sample 2 is not meaningful (grey shading). A small change in target abundance was observed (-3%), but the CV of the measurement (22%) was larger than the effect.
- Samples 8, 9, and 10 (orange shading) each showed a moderate increase in target abundance. In each case, the reported % change was considerably greater (>2.5X) than the CV of the fold-change measurement.

Table 3. CV and reported % change. The observed change in sample 2 (grey) is smaller than the CV of the measurement, and is not meaningful. In samples 8, 9, and 10 (orange), the reported change exceeds the CV by at least 2.5X. These changes are likely to be significant. Technical replicates of samples 1-10 are shown (n = 3).

Sample	Blot A	Blot B	Blot C	Mean	% Change	StDev	CV
	fold change	fold change	fold change	fold change			
С	1.00	1.00	1.00	1.00	0%	0.00	0%
1	0.90	1.00	1.10	1.00	0%	0.10	10%
2	1.20	0.90	0.80	0.97	-3%	0.21	22%
3	1.50	1.80	2.30	1.87	87%	0.40	22%
4	1.70	1.50	1.20	1.47	47%	0.25	17%
5	2.30	2.60	3.00	2.63	163%	0.35	13%
6	1.60	1.30	1.80	1.57	57%	0.25	16%
7	1.80	1.60	1.50	1.63	63%	0.15	9%
8	1.40	1.20	1.30	1.30	30%	0.10	8%
9	1.30	1.60	1.20	1.37	37%	0.21	15%
10	1.20	1.50	1.80	1.50	50%	0.30	20%

- b. Faint bands or subtle changes in band intensity are more difficult to detect reliably. In these situations, QWB analysis requires more extensive replication and a lower CV.
 - Technical replicates: If the CV of your technical replicates is high, you may need to identify and reduce sources of error in your protocol to increase the precision of your assay. Technical replicates help you characterize the variability of your assay, so you can determine if the experimental response is reliable and meaningful.
 - Biological replicates: Interpret your technical replication data in the context of your biological replicates. A strong experimental response may be easily demonstrated, even with substantial technical variation. However, a weak or variable response may be very difficult to discriminate from the assay noise, and may require more extensive technical replication [1].

Note: These are general guidelines only. Replication needs and data interpretation are specific to your experiment, and you may wish to consult a statistician.

References

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Please see attached full manuscript for further reading.

Protocol

Step 1.

Prepare an analysis spreadsheet with the **normalized band intensity values** for each replicate (as described in Guidelines).

P NOTES

Margaret Dentlinger 07 Dec 2017

Western blot data must be normalized BEFORE replicate analysis is performed. See Guidelines for more information.

Step 2.

Using the normalized values, calculate the fold change (ratio of the experimental sample to the control) for each sample on replicate blot A (depicted in Figure 2).

$$Fold\ change = \frac{\text{normalized signal}_{\text{SAMPLE}}}{\text{normalized signal}_{\text{CONTROL}}}$$

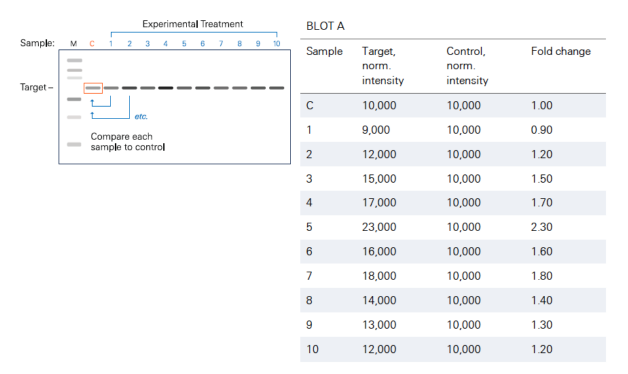


Figure 2. Fold change calculations: example data for Blot A. (*left*) In the blot diagram, C indicates the untreated control sample. Samples 1-10 received the experimental treatments. M indicates molecular weight marker. (*right*) Fold change was calculated for each band, using the normalized intensity values from Blot A. Each experimental sample is compared to the control, to generate a ratio. Example values are shown for illustration.

Step 3.

Repeat the fold change calculations with the normalized values from replicate blots B and C (depicted in Figure 3).

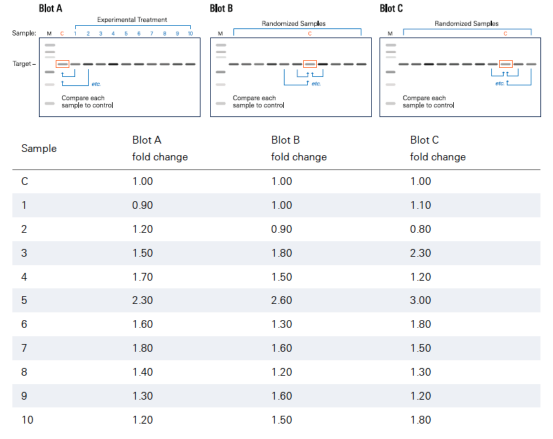


Figure 3. Fold change calculations. Example data are shown for Blots A, B, and C. *(top)* Strategy for calculating the fold change. *(bottom)* Fold change values for the three blots. In this illustration, samples 1-10 were loaded on replicate blots (Blots A, B, and C are technical replicates).

Step 4.

Using the fold change values from each replicate blot, calculate the mean fold change, overall % change, standard deviation, and coefficient of variation (CV) for all replicates (Table 2) (Steps 5-7).

EXPECTED RESULTS

Table 2. Mean, % change, standard deviation, and CV calculations. Example data are shown for analysis of samples 1-10 on Blots A, B, and C (n = 3 technical replicates).

Sample	Blot A fold change	Blot B fold change	Blot C fold change	Mean fold change	% change	StDev	CV
С	1.00	1.00	1.00	1.00	0%	0.00	0%
1	0.90	1.00	1.10	1.00	0%	0.10	10%
2	1.20	0.90	0.80	0.97	-3%	0.21	22%
3	1.50	1.80	2.30	1.87	87%	0.40	22%
4	1.70	1.50	1.20	1.47	47%	0.25	17%
5	2.30	2.60	3.00	2.63	163%	0.35	13%
6	1.60	1.30	1.80	1.57	57%	0.25	16%
7	1.80	1.60	1.50	1.63	63%	0.15	9%
8	1.40	1.20	1.30	1.30	30%	0.10	8%
9	1.30	1.60	1.20	1.37	37%	0.21	15%
10	1.20	1.50	1.80	1.50	50%	0.30	20%

Step 5.

Calculate the **mean fold change** of each replicate measurement.

Mean fold change =
$$\frac{\text{sum of replicate values for an experimental sample}}{\text{number of replicates for that sample}}$$

Step 6.

Calculate the **standard deviatio**n of the fold change for each replicate measurement.

Step 7.

Calculate the **coefficient of variation** (CV) of the fold change for each replicate measurement.

$$CV = \frac{\text{standard deviation of fold change for replicates}}{\text{mean fold change of replicates}} X 100\%$$

Step 8.

Plot the fold change in protein expression as a function of the treatment (Figure 4).

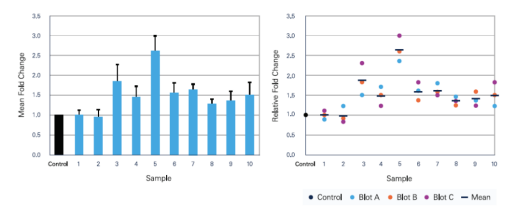


Figure 4. Bar graph and scatterplot of the fold change data in Table II. (left) Mean fold change was plotted for each sample on a bar graph. Error bars indicate the standard deviation for each measurement (n = 3). The control is indicated by a black bar. (right) Fold change values from all blots were displayed as a scatterplot. The mean fold change for each sample is indicated by a horizontal line. The control sample is represented by a black dot. Technical replicates of samples 1-10 are shown (n = 3).

NOTES

Margaret Dentlinger 07 Dec 2017

Some journals now prefer that small datasets be presented in scatterplot format, rather than the more typical bar graphs (9 - 10).

- Scatterplots clearly show the spread and distribution of your data points, making it easier for peer reviewers and readers to fully understand your results.
- Bar graphs are more suitable for large datasets. Many journals now discourage the use of error bars that indicate the standard error of measurement (SEM). Error bars that indicate the standard deviation of your measurements may be more appropriate (11, 12, 13).
- Figure legends should clearly state what the error bars on the graph represent, and the value of n for the experiment. Sufficient detail should be provided to indicate if replicates are technical or biological.