Total Protein Analysis as a Reliable Loading Control for Quantitative Fluorescent Western Blotting

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Abstract

Western blotting has been a key technique for determining the relative expression of proteins within complex biological samples since the first publications in 1979. Recent developments in sensitive fluorescent labels, with truly quantifiable linear ranges and greater limits of detection, have allowed biologists to probe tissue specific pathways and processes at higher resolution than ever before. However, the application of quantitative Western blotting (QWB) to a range of healthy tissues and those from degenerative models has highlighted a problem with significant consequences for quantitative protein analysis: how can researchers conduct comparative expression analyses when many of the commonly used reference proteins (e.g. loading controls) are differentially expressed? Here we demonstrate that common controls, including actin and tubulin, are differentially expressed in tissues from a wide range of animal models of neurodegeneration. We highlight the prevalence of such alterations through examination of published “–omics” data, and demonstrate similar responses in sensitive QWB experiments. For example, QWB analysis of spinal cord from a murine model of Spinal Muscular Atrophy using an Odyssey scanner revealed that beta-actin expression was decreased by 19.3±2% compared to healthy littermate controls. Thus, normalising QWB data to β-actin in these circumstances could result in ‘skewing’ of all data by ~20%. We further demonstrate that differential expression of commonly used loading controls was not restricted to the nervous system, but was also detectable across multiple tissues, including bone, fat and internal organs. Moreover, expression of these “control” proteins was not consistent between different portions of the same tissue, highlighting the importance of careful and consistent tissue sampling for QWB experiments. Finally, having illustrated the problem of selecting appropriate single protein loading controls, we demonstrate that normalisation using total protein analysis on samples run in parallel with stains such as Coomassie blue provides a more robust approach.

Background

Biochemical analysis using Western blotting is an essential tool in determining relative protein expression in complex biological samples. It is often used in conjunction with mass screening technologies such as proteomics to confirm differential candidate expression in various models of disease. Together with increasingly sophisticated in vivo and in vitro biological models, quantitative protein expression analyses are frequently being employed in attempts to elucidate the molecular mechanisms regulating cellular form and function in health and disease.

Traditionally, Western blotting with ECL (enhanced chemiluminescence) has been referred to as a semi-quantitative technique due to the lack of cumulative luminescence linearity and limited quantitative reproducibility [1]. With the development of more sensitive fluorescent labelling, which demonstrates a greater quantifiable linear range, sensitivity and stability in comparison to conventional ECL detection [2], analysis of protein expression can be justifiably termed “quantitative”. It is therefore imperative to ensure uniformity of sample loading with an even greater degree of precision to avoid erroneous data acquisition when using these higher resolution tools [3]. The leading company in this market is LICOR and its Odyssey fluorescence imaging scanner appears to have the most significant market penetration at present with 206 instruments currently installed in the UK alone (Personal communication; LICOR technical consultants).

In order to accurately measure protein levels in a sample, “loading control” (LC) proteins are commonly used as internal standards. The loading controls are generally derived from ubiquitously expressed “housekeeping” genes and have been widely used due to their presumed consistent level of expression across a diverse range of samples. Actin and tubulin are two of most frequently used loading controls in biomedical research, however an increasing number of studies have suggested they may...
be differentially expressed in animal and experimental models [4–
0]. Furthermore, LCs may also differ in expression from tissue to
tissue or following exposure to infectious agents [9]. Therefore,
normalising data according to loading control protein expression
could further skew results leading to erroneous conclusions.

This study set out to specifically investigate the reliability of
loading controls as internal standards, and characterise a robust,
reproducible and simple method for normalising protein load
when practicing modern quantitative Western blotting. We
present data focusing on the expression levels of commonly used
loading control proteins in the nervous system, as this is our
specific area of research interest. However, we go on to
demonstrate the importance of accurate loading controls for
research on a broad range of biological tissues and demonstrate
that normalisation using total protein analysis (TPA) with stains
such as Coomassie and Instant blue provides a more robust
baseline for performing QWB experiments.

Results and Discussion
Expression Levels of Commonly used “Loading Control”
Proteins

In order to obtain an initial estimate for the variability of
expression levels of commonly used loading control proteins in
experimental studies, we first undertook an examination of
published protein expression data from a range of human and
animal model studies including four of our own published datasets
(Table 1; [6,10–16]). With the increased stringency requirements
for mass screening reporting, raw data sets are publicly available
for many experimental comparisons, allowing further examination
of the data for proteins of interest which may not have been
highlighted in the full manuscript. We began with a focus on the
expression levels of commonly used loading control proteins within
the nervous system, as this is our specific area of research interest.
We searched these datasets for expression data on a range of
cytoskeletal proteins (actin, actinin and various tubulin isoforms),
mitochondrial proteins (VDAC1 and VDAC2) and a nuclear
protein (H1C1), all of which are commonly used as a loading
control or internal reference proteins for expression value
normalisation in comparative protein quantitation experiments
(Table 1).

This initial analysis revealed evidence for differential expression
of all loading controls assessed using mass screening tools,
including both -array technologies and proteomics. Discrepancies
in expression of loading control proteins were detected across a
diverse range of conditions, including Alzheimer’s disease,
lysosomal storage disorders and the motor neurone disease Spinal
Muscular Atrophy (SMA). Notably, this re-interrogation of
published data also identified differential expression of loading
control proteins across an assortment of tissues sampled, an issue
of potential critical importance for subsequent data normalisation
and post-omics validation.

Actin and Tubulin are differentially Expressed in
Pathologically-affected Tissue from a Mouse Model of
SMA

Given that our analysis of raw data from published protein
expression studies revealed widespread differential expression of
common loading controls in various pathological conditions and
neurological diseases ([10,17], cf. Table 1), we next wanted to
demonstrate that such alterations can also be detected when using
modern QWB techniques. Using spinal cord tissue harvested from
an established mouse model of severe SMA [18–19], we next
quantified levels of β-actin and β-tubulin proteins. We found
altered expression levels of both β-actin and β-tubulin when
comparing the spinal cord of SMA affected mice (SMN1/SMN2
with littermate controls (figure 1a. and 1b. respectively) using
QWB. Both β-actin and β-tubulin expression were significantly
down regulated in SMA compared to control tissue, by 19.3±2% and
7.3±0.3% respectively (mean ± SEM). Moreover, determina-
tion of total protein load demonstrated a high level of uniformity
with a difference of 1.8±0.4% (mean ± SEM) between wild type
and affected mice (n=6) (figure 1c). Thus, altered LC protein
expression as highlighted by proteomic studies on tissues from
neurodegenerative conditions (including SMA; see Table 1) can also
be detected by quantitative Western blot.

Differential β-actin Expression is Detectable by QWB
across Multiple Tissues from Wild-type Mice

Given that significant alterations in common loading control
proteins can be detected in affected tissues sampled from a range
of disease models (Table 1), we next wanted to establish whether
similar differences in expression could be identified in healthy
(‘wild-type’) tissue, and also establish whether several tissue types
could be differentially affected within the same individual. This
latter issue is particularly pertinent for many neurodegenerative
diseases (including SMA), where multi-system pathology is now
being reported (e.g. [20]). QWB screening of multiple tissues for
expression of candidate proteins is common practice and is an
especially critical procedure when comparing systemic protein
expression profiles, identifying biomarkers for disease progression
in peripherally accessible tissues, or validating regulatory proteins
during genetic or pharmacological manipulation. It is therefore
essential for a loading control to exhibit stable expression across a
wide array of tissues. However, our studies using C57Bl/6J (‘wild-
type’) mice demonstrated that expression of β-actin varied
considerably across different tissues from the same mouse
(figure 2 A & B). In order to verify that the variability of β-actin
expression was not due to loading error, quantification of the
protein load of each tissue was carried out on a series of mass
ranges corresponding to protein electrophoretic migration
(figure 2C). These data demonstrated low individual variation
of protein load across the different tissue samples within the series
of molecular weight ranges measured. Variability ranged from only
1.87% (SEM) in the 40–80 kDa range up to 5.65% for the
broadest mass range of 10–160 kDa across all tissues. Therefore,
consistency of load across each of the tissues was demonstrated
and was independent of the mass range measured (figure 2C).

Expression of β-actin was strikingly different when comparing a
diverse range of tissues (figure 2B & D). An appreciable disparity
in β-actin expression was observed between heart and spleen tissue,
with a difference of 44 arbitrary fluorescence units (AFU)
contrastng with the consistent total protein load demonstrated in
the 40–80 kDa molecular weight range (figure 2D). Critically,
normalising data to β-actin expression for cross tissue comparisons
could therefore result in skewing of data by up to 22 fold when
comparing these tissues. However, there was some homology in β-
actin expression between certain tissues such as bone and fat
(5.33 AFU and 4.99 AFU respectively), and for these tissues it may
therefore be acceptable to use β-actin as an internal control, but
only when comparing these specific samples. Interestingly,
a similar issue regarding β-actin has been raised with cross-tissue
RNA expression profiling. For example, a qRT-PCR study in fish
concurs with our findings suggesting that β-actin is not an ideal
loading control for certain tissue comparisons particularly for
heart, muscle and brain, whereas its expression is more consistent
between kidney and spleen [9]. Our results therefore demonstrate
that it would be advisable to use total protein expression as an

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Table 1. Examination of publicly available mass screening data reveals alterations in expression levels of commonly used “loading control” proteins.

<table>
<thead>
<tr>
<th>Protein Name</th>
<th>Symbol</th>
<th>Protein ID</th>
<th>Localisation</th>
<th>Alteration</th>
<th>Model</th>
<th>Tissue</th>
<th>Technique</th>
<th>Reference</th>
</tr>
</thead>
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<tr>
<td>Actin</td>
<td>Actc1</td>
<td>IPI00654242.1</td>
<td>Cytoskeleton</td>
<td>Upregulated (3.07 fold)</td>
<td>SMA mouse</td>
<td>Cardiac Muscle</td>
<td>Label Free Proteomics</td>
<td>Mutsaers et al 2011 [6]</td>
</tr>
<tr>
<td>Actin</td>
<td></td>
<td>p60710</td>
<td>Cytoskeleton</td>
<td>Absent</td>
<td>Wlds mouse</td>
<td>Striatal Synapses</td>
<td>2D Gel Proteomics</td>
<td>Wishart et al 2007 [10]</td>
</tr>
<tr>
<td>actinin, alpha 1</td>
<td>Actn1</td>
<td>Q7TPR4</td>
<td>Cytoskeleton</td>
<td>Downregulated (0.6 ratio)</td>
<td>SOD1 mouse</td>
<td>Cortical Synapses</td>
<td>iTRAQ Proteomics</td>
<td>Flynn et al 2012 [11]</td>
</tr>
<tr>
<td>Histone Cluster 1</td>
<td>H1C1</td>
<td>NP_056601</td>
<td>Nuclear</td>
<td>Upregulated (1.3 ratio)</td>
<td>AOPE mouse</td>
<td>Sciatic Nerve</td>
<td>iTRAQ Proteomics</td>
<td>Comely et al 2011 [12]</td>
</tr>
<tr>
<td>Tubulin Alpha 18 Chain</td>
<td>ActA1B</td>
<td>IPI00117348.4</td>
<td>Cytoskeleton</td>
<td>Upregulated (1.32 fold)</td>
<td>SMA mouse</td>
<td>Skeletal Muscle</td>
<td>Label Free Proteomics</td>
<td>Mutsaers et al 2011 [6]</td>
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<tr>
<td>Tubulin Beta 4 Chain</td>
<td>TubB4</td>
<td>IPI00109073.5</td>
<td>Cytoskeleton</td>
<td>Downregulated (−1.16 ratio)</td>
<td>SMA mouse</td>
<td>Hippocampus</td>
<td>iTRAQ Proteomics</td>
<td>Wishart et al 2010 [13]</td>
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<td>Tubulin beta-4 chain</td>
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<td>Cytoskeleton</td>
<td>Downregulated (−0.28 ratio)</td>
<td>Ercc1 mouse</td>
<td>Hippocampal synapses</td>
<td>iTRAQ Proteomics</td>
<td>Vegh et al 2012 [14]</td>
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<tr>
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<td>P68366</td>
<td>Cytoskeleton</td>
<td>Downregulated (−1.13 fold)</td>
<td>FTLD patient</td>
<td>Cerebellum</td>
<td>Gel Separation &amp; Label free</td>
<td>de Souza et al 2012 [15]</td>
</tr>
<tr>
<td>tubulin, beta 2a</td>
<td>Tubb2A</td>
<td>Q9CWF2</td>
<td>Cytoskeleton</td>
<td>Upregulated (0.26 ratio)</td>
<td>SOD1 mouse</td>
<td>Cortical Synapses</td>
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<td>Prefrontal Cortex</td>
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<td>de Souza et al 2012 [15]</td>
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<tr>
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<td>VDAC1</td>
<td>q60932</td>
<td>Mitochondrial</td>
<td>Downregulated (−408 fold)</td>
<td>Wlds mouse</td>
<td>Striatal Synapses</td>
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<td>Wishart et al 2007 [10]</td>
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<td>Cortex</td>
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<tr>
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<td>Mitochondrial</td>
<td>Downregulated (−0.43 ratio)</td>
<td>SOD1 mouse</td>
<td>Cortical Synapses</td>
<td>iTRAQ Proteomics</td>
<td>Flynn et al 2012 [11]</td>
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Note: Alterations listed in table are in units published in the original manuscript or accompanying supplementary material.
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indication of protein load to accurately perform a comparative expression analysis across a wide range of tissue samples.

Loading Control Expression is not Homogeneous throughout Structurally Asymmetrical Tissues

Having demonstrated that expression of common loading controls such as actin was not necessarily consistent when comparing different tissues from the same animal, we next wanted to establish whether or not uniformity of protein expression was preserved throughout a single tissue from the same animal. Anecdotally it appears that researchers assume that loading control protein expression is stable within a given organ or tissue, with methodology sections of manuscripts routinely detailing gross, rather than specific, anatomical terms to describe tissue harvesting. To assess consistency within a biologically-relevant tissue, we measured and compared β-actin and neurofilament-light (NF-L) levels in proximal versus distal portions of the same mouse sciatic nerve using QWB. We found that levels of these LC proteins were not consistent throughout the two portions of the same nerve (Figure 3). Once again, we found that total protein load was consistent when quantified, however β-actin (a predominantly cytoplasmic isoform; [21] labelling was significantly higher (p = 0.0045), 52% greater ±14.3% (SEM), when comparing the proximal to distal portion of the sciatic nerve (figure 3C & E). In contrast expression of neurofilament (NF-L), a major component of the neuronal cytoskeleton [22], was appreciably higher, nearly 8 fold with up to a 4 fold SEM in the distal portion of the sciatic nerve (figure 3D & F). Therefore, these results stress that accuracy and consistency of dissection are crucial, even when evaluating the same tissues from a single animal for comparative analysis. Whilst this should be standard practice regardless, it is especially true for structurally asymmetrical tissues and failure to do so could have significant consequences for both -omics screens (such as proteomic comparisons) and for QWB analyses.

Limitations in Standard Loading Controls: β-actin and β-tubulin Working Range and Sensitivity Explored

The ideal internal loading control protein for QWB must be abundant with a wide linear range of detection to accommodate proteins of varying levels of expression. In order to evaluate the suitability of β-actin and β-tubulin as loading controls for QWB we tested both their working linear range and sensitivity by quantifying their expression throughout a dilution series, ranging from 1 to 40 μg of protein, produced from mouse whole brain tissue homogenate (figure 4a & b). The working range of β-actin where linearity was maintained was between 1 and 30 μg of protein loaded (figure 4a & b). Here, our quantitative Western blotting data conflicts with that of others who have reported that the linear range of β-actin was far smaller, only up to 2 μg of protein before saturation of the signal occurred [4,8]. However these studies used less sensitive ECL detection methodologies therefore this disparity is most probably caused by the limitation of ECL based imaging and “quantification”.

When examining β-tubulin, it was evident that this protein was so abundant in brain extracts that the signal began to saturate out at less than 10 μg of protein (figure 4 a & b). In order to accurately determine the precise linear range of β-tubulin a tighter dilution series of protein load, 0.5 to 14 μg, was employed (figure 4c & d). Saturation of the β-tubulin signal occurred at 8–10 μg of protein load. Again, our QWB results conflicted with previous ECL studies suggesting that the linear range of β-tubulin peaks at 5 μg [4] and it is assumed that this is also likely due to the limitation of ECL based imaging and “quantification”. Our results suggest that β-tubulin rather than β-actin would be a more appropriate loading control when detecting low-abundance proteins in homogeneous conditions.
extracts, however actin has a greater working range of sensitivity. In addition, our data has emphasised the superior sensitivity of fluorescent QWB in comparison to ECL as we demonstrate a far wider working range for both of these commonly used single protein loading controls.

Total Protein Analysis is an Accurate Measure of Protein Load

Our analyses detailed above have highlighted several important variables that need considering when choosing an internal control for QWB, including but not limited to the linear range of sensitivity and disparities in expression across different tissue samples or within portions of the same tissue. Finally, we wanted to establish whether a total protein analysis approach would provide a more reliable and accurate measure of protein load for QWB experiments. To begin, we assessed the detection sensitivity using a dilution series created using a bovine serum albumin (BSA) protein standard (Figure 4E & F). BSA has a single band at a molecular weight of 66.5 kDa. We were able to detect a linear fluorescence profile across a broad concentration range as determined by the coefficient of variation ($R^2$ value) of 0.998. Importantly, as the BSA standard dilution series effectively means that all of the protein loaded is represented by a single band, this not only validates the linear nature of detection using this system but also demonstrates the lack of saturation typically found with ECL based systems.

Finally, when applied to real biological samples using the LICOR Odyssey quantitative scanning system, total protein analysis (using coomassie) was linear in its detection across a broader range of protein loading than either actin or tubulin (1 to 40 µg; Figure 4G & H). The coefficient of variations for both Bicinchoninic Acid solution (BCA) assay and total protein analysis were 0.979 and 0.996 respectively. These $R^2$ values demonstrated a high degree of linearity in both assays. Moreover, total protein analysis correlated directly with BCA protein concentration data (Figure 4G & H) throughout the broad 1–40 µg protein load dilution series further indicating its reliability as a control for protein load using the Odyssey quantitative imaging system. We therefore suggest that the use of total protein analysis provides a measure of protein load that circumvents many of the problems associated with the use of single loading control proteins: it is unchanged when comparing tissues from different models (c.f. Figure 1); it is consistent across different tissue types (c.f. Figure 2); and different portions of the same tissue (c.f. Figure 3).

Conclusions

Western blotting has traditionally been a “semi-quantitative” technique using house keeping genes as internal reference standards. These standards are required to compensate for any technical errors that may have arisen due to issues such as poor transfer or unequivocal loading. However, our QWB studies have demonstrated a critical problem with the use of some common loading controls for this role. Differential expression of commonly
used loading control proteins including β-actin and β-tubulin, occurs when comparing a wide range of tissues, when examining different portions of the same tissue and when pathological conditions arise. Therefore, if normalisation (using single protein loading control expression as a correction factor) of quantitative Western blotting results is required, all of the resulting data could be skewed as a result of the differential expression of a single protein [23].

Total protein analysis is an alternative simple technique in QWB to accurately determine if equivalent protein loading has been achieved within a gel [24]. Data obtained by total protein analysis is independent of the pitfalls that can occur using “common” housekeeping genes as loading controls. That is not to say that housekeeping genes cannot be used as loading controls, but that they should only be used in a limited fashion once the researcher has fully investigated sample expression homogeneity for the gene in question, and if the protein load falls within the working range of that particular loading control. Consequently, we propose it would be prudent to use total protein analysis to save time, resources, increase sensitivity and accuracy as well as the working range of protein load for quantitative Western blotting. Total protein analysis should therefore be considered an alternative standard reference for data normalisation in modern quantitative fluorescent Western blotting.

**Materials and Methods**

**Tissue Harvesting and Protein Extraction**

**Ethics statement.** All animal experiments were approved by a University of Edinburgh internal ethics committee and were performed under license by the UK Home Office (project license number 60/3891).

**Preparation of severe model of Spinal Muscular Atrophy (SMA) spinal cord homogenates.** Spinal cord were harvested from Smn−/−/Smn2 severe model of SMA and wild-type controls at P5 and homogenised in RIPA buffer (Sigma, UK) containing 5% protease inhibitor cocktail (Roche) as previously described [13]. All SMA based comparisons were therefore Smn−/−/Smn2 (SMA) compared to wild-type (WT) litter mate controls unless otherwise stated. Protein was extracted and concentrations determined using a BCA assay (Ferment) according to manufacturers instructions, as previously described [10,13,25].

**Preparation of range of tissue samples from C57/black mice.** Quadriceps femoris muscle, gonadal fat, heart, calvaria,
Figure 4. Linear range and sensitivity of total protein stain is greater than the conventional loading controls β-actin or β-tubulin. (A) Representative LICOR image for a protein dilution series of whole brain homogenate 1, 5, 10, 20, 30, 40 µg demonstrating the working range of β-actin and β-tubulin when using QWB. (B) Quantification of protein dilution series showing the linear ranges of β-actin (black circle) and β-tubulin (open triangle). Note that tubulin expression appears to saturate at less than 10 µg of brain homogenate. (C) In order to pinpoint the saturation level a tubulin specific protein dilution series over a smaller range (0.5, 1, 2, 4, 6, 8, 10, 12 and 14 µg) establishing the saturation point of β-tubulin when using QWB. (D) Quantification of β-tubulin linear range. (E) Total protein stain of dilution series 2, 10, 20, 40, 80 µg made using the bovine serum albumin standard (2 µg/µl) from the Pierce BCA kit (see methods). BSA molecular weight is 66.5 kDa. Imaging of this dilution series demonstrates imaging of a broad concentration range without saturation at a single protein mass. (F) Graphical representation of quantification from BSA dilution series in panel E. This demonstrates wide linear detection and high correlation (0.998) validating the use of total protein measurements as a viable method for detecting protein load using the LICOR system. (G) Total protein stain of whole brain homogenate dilution series 1, 5, 10, 20, 30, 40 µg
spleen and tibial bone tissue were dissected from 10 day old C57Bl/6 mice. Tissues were processed as outlined above.

**Preparation of sciatic nerve tissue.** Proximal and distal sections of the sciatic nerve from C57/ black mice were dissected in 1xPBS (phosphate buffered saline) and frozen immediately on dry ice. The samples were homogenised in iTRAQ buffer (6 M urea, 2 M thiourea, 2% CHAPS, 0.5% SDS), sonicated 5×10 secs on ice, and centrifugated at 14 K for 30 minutes. 1:100 dilutions of supernatant: dH2O were used in a BCA assay following manufacturers instructions.

**Quantitative Western Blotting**

Samples were denatured in NuPage® LDS Sample buffer 4X (Invitrogen, UK) at 98°C and 15 μg of protein loaded (with the exception of the protein dilution series) and run on commercially produced pre-cast 4–12% Bis-Tris gels (Invitrogen). Gels were run in duplicate in parallel in the same electrophoretic tank at the same time. One gel was stained using Instant blue (Expedeon) or coomassie (see total protein stain below) and one was used to transfer the protein to the polyvinylidene fluoride (PVDF) membrane using the I-Blot® transfer system (Invitrogen, UK) using programme 3 for 8.5 minutes. Membranes were incubated with Odyssey blocking buffer (Li-Cor) prior to incubation with rabbit polyclonal antibodies directed against β-actin (1:1000, Abcam 8226), β-tubulin (1:1000, Abcam 8226) and mouse monoclonal anti-NF-L (Millipore AB9568) overnight at 4°C. Goat anti-rabbit IgG (H+L) 800 CW, goat anti-rabbit (680 RD) and/or goat anti-mouse (H+L) was applied for 90 minutes at room temperature (1:5000, LI-COR) prior to washing with PBS. Visualisation and quantification was carried out with the LI-COR Odyssey® scanner and software (LI-COR Biosciences). Blots (and gels) were imaged using an Odyssey Infrared Imaging System Scan resolution of the instrument ranges from 21 to 339 μm, and in this study blots (and gels) were imaged at 169 μm. Quantification was performed on single channels with the analysis software provided as previously described [10,13,25].

**Total Protein Gel Stain**

All total protein stains within this manuscript have been carried out on gels and not membranes unless otherwise stated. As such there are caveats which should be taken into consideration if comparing total protein stained gels with membranes due to variables accompanying membrane transfer which are not accounted for when using this approach (see below). Post electrophoresis gels (see above) were stained using either Instant Blue (Expedeon) or Coomassie (0.1% Coomassie R250, 40% methanol, 10% acetic acid) solution. Gels were left in Instant blue for 1 hour and washed in dH2O prior to visualisation. Coomassie stained gels were left in Coomassie solution for 1 hour and de-stained using several washes in de-stain solution (40% methanol, 10% acetic acid) and then washed in dH2O prior to visualisation. Stained gels were imaged directly on the Li-COR Odyssey® scanner using the 700 channel and quantified using the Odyssey® software. See above. Incidentally the instant blue stain can be visualised in both the 700 and 800 channels however it has greater resolution in the 700 channel. For total protein stains to be of use as loading controls, ideally the membranes to be probed should be stained directly for load. However, there are limitations with this in modern fluorescent imaging systems. Coomassie is not as effective on most PVDF membrane types when compared to stained gels as the background auto-fluorescence is naturally higher. If used directly on a membrane coomassie does not strip to allow for re-probing in the same imaging channel. Ponceau stains are reversible but staining is difficult to visualise and remains to be proven linear in its adherence. Moreover, any stain which is not blue in presentation or does not fluoresce in the 680 or 800 wavelength channels used by the LICOR Odyssey scanner can not be imaged using this system and measurement scaling will therefore differ from the quantitative fluorescent blots for candidates of interest. Parallel commercially produced precast gels should therefore be used to reduce polyacrylamide matrix variability; they should be loaded at the same time using the same “master mix” (i.e. protein/water/loading dye) in the same tank (multi gel capacity) in order to be run from the same powerpack under the same conditions. A further variable when comparing a probed membrane to a stained gel are alterations introduced by variable transfer efficiency. Potential inconsistency in transfer efficiency generally occurs according to differential molecular weight rather than inter lane variability i.e. higher efficiency of transfer with lower molecular weight. The use of commercially procured transfer packs (such as the I-Blot® stacks; see above) coupled with a rapid semi-dry fast transfer system as detailed above should further improve reproducibility. By being aware of the steps where possible error could be introduced and taking the appropriate precautions such as those listed here, inter gel variability and transfer variance should be kept to a minimum, and appropriate data interpretation can be expected within the limitations of the system.

**Calculation of Linear Ranges of β-actin and β-tubulin**

Concentration of protein extracts can be determined in a variety of ways. The most commonly used may be the Bicinchoninic Acid assays (BCA). BCA assays involve reduction of copper ions in a temperature dependant fashion with the level of reduction correlating with protein concentration. Reduced copper ions bind to BCA forming a purple product which can be detected at 562 nm. Each run includes a dilution series of a known protein standard – bovine serum albumin (BSA) as a reference curve to allow determination of absolute protein concentrations. As each reaction set can be subtly influenced by incubation time and temperature, samples which will be grouped together for analysis should routinely be assayed together against the same standard curve. Here we employed a series of protein dilutions (1, 5, 10, 20, 30, 40 μg) and (0.5, 1, 2, 4, 6, 8, 10, 12, 14 μg) which were produced from mouse whole brain homogenate. Preparation of the dilution series was carried out after the concentration of protein had been determined using a micro BCA assay (Pierce). Briefly, two 4–12% Bis-tris gels were loaded; one stained for total protein (Coomassie or Instant Blue) and the other was transferred for QWB as above. Visualisation and quantification was carried out using the LI-COR Odyssey imager and software. See above.

**Photography**

Photographs of sciatic nerve dissection and organs were taken using a Nikon D200 camera with 105 mm micro NIKKOR F2.8 lens.
Data Analysis and Figure Production

QWB data was analysed using Odyssey software as per manufactures guidelines and as previously described [10]. Data was graphed and statistical comparisons carried out using GraphPad Prism as previously described [23]. Image overlays were produced using Adobe photoshop to overlay 700 & 800 channels obtained from the Odyssey imager (LICOR Biosciences).

References


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Author Contributions

Conceived and designed the experiments: TMW SLE. Performed the experiments: SLE SLR ML KO. Analyzed the data: SLE SLR ML KO CF THG TMW. Contributed reagents/materials/analysis tools: CF THG TMW. Wrote the paper: SLE THG TMW.