Visual Abstract

C-bouton Changes in ALS

- Reduced C-bouton number and density on motoneurons only noted at end-stage
- No change in C-bouton size noted throughout G93A disease pathogenesis

The possible presence of pathological changes in cholinergic synaptic inputs [cholinergic boutons (C-boutons)] is a contentious topic within the ALS field. Conflicting data reported on this issue makes it difficult to assess the
roles of these synaptic inputs in ALS. Our objective was to determine whether the reported changes are truly statistically and biologically significant and why replication is problematic. This is an urgent question, as C-boutons are an important regulator of spinal motoneuron excitability, and pathological changes in motoneuron excitability are present throughout disease progression. Using male mice of the SOD1-G93A high-expresser transgenic (G93A) mouse model of ALS, we examined C-boutons on spinal motoneurons. We performed histological analysis at high statistical power, which showed no difference in C-bouton size in G93A versus wild-type motoneurons throughout disease progression. In an attempt to examine the underlying reasons for our failure to replicate reported changes, we performed further histological analyses using several variations on experimental design and data analysis that were reported in the ALS literature. This analysis showed that factors related to experimental design, such as grouping unit, sampling strategy, and blinding status, potentially contribute to the discrepancy in published data on C-bouton size changes. Next, we systematically analyzed the impact of study design variability and potential bias on reported results from experimental and preclinical studies of ALS. Strikingly, we found that practices such as blinding and power analysis are not systematically reported in the ALS field. Protocols to standardize experimental design and minimize bias are thus critical to advancing the ALS field.

**Key words:** ALS; amyotrophic lateral sclerosis; C-bouton; cholinergic synapse; motoneuron; SOD1-G93A

**Introduction**

Amyotrophic lateral sclerosis (ALS) is a progressive neurodegenerative disease that is characterized by selective degeneration and death of upper and lower motoneurons in the cortex, brainstem, and spinal cord, leading to muscle weakness, atrophy, paralysis, and death (Charcot, 1874; Rowland and Schneider, 2001). After decades of research, there are reports of many disease changes and many hypotheses on pathogenesis; yet, the field lacks the clear mechanistic understanding needed to prevent or impede motoneuron degeneration in ALS. The only Food and Drug Administration-approved treatment for ALS patients is riluzole, which extends survival marginally (~3 months; Bensimon et al., 1994).

Several mutations have been identified in ALS patient populations and recreated within transgenic animal models. Among those is the G93A superoxide dismutase 1 (G93A) model (with >25 copies; Gurney et al., 1994), which is the most widely characterized and extensively studied model of ALS (Turner and Talbot, 2008). G93A mice develop a phenotype similar to that of ALS patients, including motor impairment, axonal loss, motoneuron death, muscle atrophy, and limb weakness (Fischer et al., 2004); and riluzole was developed through studies of the G93A line. We used this disease model, with wild-type (WT) transgenic controls, in the present study.

Cholinergic bouton (C-bouton) size changes represent an important, but disputed, topic within the field. Recently, a large number of studies have examined potential changes in C-bouton size during ALS disease progression in male mice of the G93A line (summarized in Table 1). However, the reported results have been conflicting: various studies report enlargement (Pullen and Athanasiou, 2009; Herron and Miles, 2012; Saxena et al., 2013), no change (Pullen and Athanasiou, 2009; Herron and Miles, 2012; Milan et al., 2015), or shrinkage (Milan et al., 2015) of C-boutons at various disease stages. Because C-bouton changes could potentially contribute to excitability dysregulation of motoneurons in ALS, reconciling these findings is a critical issue. If these changes are verified, they would improve our understanding of the mechanisms underlying disease pathogenesis and could represent an important therapeutic target for restoring normal excitability to degenerating motoneurons. Thus, our objectives were (1) to determine whether variations in study design or analysis practices could underlie the conflicting results reported in literature and (2) to resolve this research question by examining whether we can replicate any of the changes reported in male mice of the G93A line on C-bouton size. Our hypotheses were (1) that variations in study design and data analysis would impact our study results and (2) that a large-sample study with high statistical power would demonstrate C-bouton size changes in a mouse model of ALS, compared with WT mice. Both objectives are important because conflicting reports of study results impede understanding of this and other ALS pathophysiology, thus delaying the development of effective treatments.

We used immunohistochemistry and confocal microscopy to examine C-boutons in great detail at several stages throughout disease progression. We performed histological analysis to test whether changes in C-bouton size took place, using standard published techniques.
Table 1. Recent literature examining C-bouton size in ALS mouse models

| 1. Authors | 2. Age | 3. Change | 4. Magnitude of change | 5. Calculated effect size | 6. Grouping unit | 7. Blinding | 8. Power analysis | 9. Size parameter | 10. Sex/no. of animals (WT, G93A) |
|------------|--------|-----------|------------------------|--------------------------|----------------|-------------|----------------|----------------|-----------------------------|
| Pullen and Athanasiou (2009) | P42 | NC | N/A | N/A | By bouton | Not reported | Not reported | Positional length | M² |
| | P70 | Increase | 25% | 0.71 | | | | | |
| | P156 | Increase | 56% | 1.42 | | | | | |
| Herron and Miles (2012) | P8 | NC | N/A | Inadequate information | By animal | Blinding included | Not reported | Positional length (Feret’s diameter) | M (25 mice total for all time points and groups)² |
| | P30 | Increase | 7% | | | | | | |
| | P120 | Increase | 14% | | | | | | |
| | P140 | Increase | 17% | | | | | | |
| Saxena et al. (2013) | P37 | Increase | 100% | Inadequate information | By bouton | Not reported | Not reported | Volume | M (3, 3) |
| Milan et al. (2015) | P10 | Decrease | 9% | 0.15 | By | Blinding | Not reported | Area | M (6, 6) |
| | P21 | Increase | 7% | 0.17 | cell | included | | | M (7, 6) |
| | P40 | NC | N/A | N/A | | | | | |
| | P75 | NC | N/A | N/A | | | | | |
| | P100 | Decrease | 5% | 0.43 | | | | | M (5, 8) |
| This article | P10 | NC | N/A | N/A | By | Blinding | Included | Largest cross-sectional area | M (3, 3) |
| | P30 | NC | N/A | N/A | cell | included | | | M (4, 4) |
| | P90 | NC | N/A | N/A | | | | | M (3, 3) |
| | P120 | NC | N/A | N/A | | | | | M (3, 3) |

1. Authors; 2. age of G93A mouse studied; 3. whether C-bouton size increases, decreases, or there was no change were reported for each age studied; 4. magnitude of change; 5. calculated effect size, if possible; 6. the sampling strategy (grouping unit) used; 7. the reported blinding status, if reported; and 8. the power analysis, if reported. M, Male; N/A, not available; NC, no change.

¹Inadequate information. SD values or sample size (n) data were not included in the published work, making the calculation of effect size impossible.

²No information on animal number was provided.

³Information on each time point was not provided. Female data not included in the table comparison.

(Pullen and Athanasiou, 2009; Herron and Miles, 2012; Saxena et al., 2013; Milan et al., 2015). We then performed additional examinations of C-boutons while varying methods of experimental design and data analysis to identify potential sources of research design variability and bias that might impact the results. Last, we examined the possible contribution of research design variability and bias to the conflicting results reported in experimental and preclinical studies of ALS.

Our results show that C-bouton size does not change throughout the disease. Additionally, our results show that variations in the methods of data breakdown and sampling strategy, as well as whether blinding was used during analysis, has significant impact on the outcomes of statistical analysis, such that a statistically significant increase or decrease, or no change, in C-bouton size could all be produced from the same dataset. We concluded that these issues likely contribute to conflicting reports on C-bouton changes in the ALS literature, and that the standardization of experimental design and data analysis will benefit the field.

Materials and Methods

All animal procedures were performed in accordance with the regulations of the Wright State University Laboratory Animal Care and Use Committee (LACUC).

Animal genetic background

All mice were either purchased from The Jackson Laboratory or bred from these mice to produce male mice with a B6SJL-TG genetic background (Tg(SOD1-G93A)1Gur). Briefly, B6/SJL hybrid females were bred with male hemizygous mice expressing the human SOD1 gene with a glycine-to-alanine mutation at amino acid 93 (SOD1-G93A). Male offspring of this pairing were used for all experiments and compared with their noncarrier littersmates. Because a major goal of our study was to compare our results with a large body of literature, we used males in the present study because ALS studies looking at C-bouton changes have largely used males, and in humans it appears that males are affected more by ALS than females (McCombe and Henderson, 2010). Genotyping using tail clippings was performed by Transnetyx. All mutant hemizygous mice expressed a high copy of the mutated gene (>25 copies). Mutant mice and their noncarrier littersmates were killed at four time points [postnatal day 10 (P10), P30, P90, and end-stage], which were predefined at full hindlimb paralysis (~P120–P140).

Animal surgical procedures

All animals were anesthetized with EUTHASOL solution (pentobarbital sodium and phenytoin sodium) and transcardially perfused with a vascular rinse (0.01 M phosphate buffer with 0.8% NaCl, 0.025% KCl, and 0.05% NaHCO3, pH 7–8), followed by 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7–8.

Tissue preparation

The lower lumbar spinal cord was quickly removed and postfixed in 4% paraformaldehyde fixative for ~2 h or overnight. Tissue was then stored in 15% sucrose at 4°C overnight. Transverse sections of L4–L6 spinal cords were then cut on a cryostat at a thickness of ~50 μm and collected in 0.1 M PBS, pH 7–8.

Spinal cord immunohistochemistry

Sections were rinsed with PBS-T (0.01 M PBS containing 0.1% Triton-X, pH 7.3), blocked with normal horse serum (10% in PBS-T), and then incubated free floating in cocktails of primary antibodies overnight at 4°C. All anti-
bodies were diluted with PBS-T. Nissl immunocytochemistry was performed using a 435/455 blue fluorescent Nissl stain (1:100; catalog #N-21479, Neurotrace, Life Technologies) to visualize cell bodies. Labeling of C-boutons was performed with anti-vesicular acetylcholine transporter (VACHT; 1:1000 dilution; goat, Abcam; RRID: AB_956453). All primary antibodies were diluted in PBS-T 0.1%, pH 7.4, and incubated overnight at 4°C. The sensitivity and specificity of the primary antibodies against VACHT have been demonstrated previously (Alvarez et al., 1998, 1999; Deng and Fyffe, 2004; Muennich and Fyffe, 2004; Deardorff et al., 2013). Immunoreactivity was detected with a goat–specific secondary antibody conjugated to Alexa Fluor 647 (Jackson ImmunoResearch) diluted 1:50 in PBS-T 0.1%, pH 7.4, and incubated at room temperature for ~3 h. Sections were then mounted onto slides and coverslipped in Vectashield mounting medium (Vector Laboratories).

**Confocal microscopy and bouton analysis**

Images were obtained on a Fluoview 1000 (Olympus) confocal microscope with a 60× oil-immersion objective in 1 μm steps. Alpha-motoneurons were differentiated on the basis that soma size measurements fell within a previously published range of >300 μm$^2$ (Ishihara et al., 2013), that they were located in Rexed lamina IX, and that they received synaptic input from large cholinergic boutons, as evidenced by VACHT immunoreactivity (VACHT-IR). A 2-D analysis of motoneurons was then performed. This process allowed for the relatively efficient analysis of the somatic morphology and neurochemistry of a large sample of motoneurons from each age and genotype group. Bouton areas were measured using Fluoview software. Regions of interest were drawn around the largest cross-sectional area of boutons on every cell.

**Bouton size measurements**

Bouton areas were obtained at four time points (P10, P30, P90, and P120+/end-stage) in G93A mutants and their wild-type littermates. One to seven boutons were randomly analyzed per motoneuron, and their properties were averaged by cell for statistical analysis. This method allowed us to measure a large number of en fosse boutons from every cell (i.e., large-sampling strategy), which we maintained throughout our work except when otherwise noted. Two to 16 spinal cord sections per animal were analyzed ($n = 2–16$) from three to five animals per age per genotype ($N = 3–5$). Our data analysis was blinded by coding image genotype information throughout the study except when mentioned otherwise.

**Bouton number and density measurements**

Density measurements were performed according to modified protocols from the study by Alvarez et al. (2011). Briefly, Nissl-stained cells stained for VACHT-IR were randomly sampled from both WT and G93A motoneurons. Each cell was imaged with a separation by 1 μm z-steps. From these image stacks, a mid-somatic region was identified by the presence of a well defined nucleolus; and from this center image, three optical sections separated by 2 μm in the z-axis (to avoid resampling the same terminals) were chosen for quantification. The files containing the sections were then labeled with a letter identifier, and the analyzer was blinded to the genotype. The number of VACHT-IR clusters on the surface of labeled Nissl cell bodies was counted, and the largest cellular perimeter (at the mid-somatic region) for each cell was measured, excluding the origins of primary dendrites. Counts and perimeter measurements were obtained with Fluoview software. Densities were estimated as the number of contacts per 100 μm of linear perimeter. An average density estimate was obtained for each motoneuron sampled.

**Statistical analyses**

Statistical analyses were experimental, not descriptive; random sampling was performed. Statistica and Graph-Pad Prism 6/7 were used for all statistical analyses. Two-way ANOVA, followed by Tukey’s post hoc tests, was performed to examine the effects of genotype and age on C-boutons. For ANOVA, $F$ values are provided to indicate the significance of the effects. For Mann–Whitney $U$ test analysis, $U$ values are provided to show the difference between the two rank totals. Significance for all tests was set at $p \leq 0.05$.

**Analysis of ALS literature**

A PubMed search was conducted for all articles containing the term “G93A” between May 1, 2015, and May 1, 2016. Any articles that were written in English and available for download through Wright State University libraries was included for analysis. Review articles were excluded, and research articles containing a significant preclinical component were noted. A database was compiled to note the presence or absence of various components of experimental design, including blinding, power analysis, and effect size. The journal name was denoted for all articles used in the analysis.

**Results**

To examine C-bouton size in ALS, we used the VACHT-IR to label and measure C-boutons in the ventral horn of spinal cords of G93A versus WT mice at various disease stages. We analyzed the following four time points: P10, P30, P90, and end-stage (P120+). At P10, many electrical and morphological motoneuron abnormalities have been observed in the G93A model, yet neurodegeneration has not started (Quinlan et al., 2011; Leroy et al., 2014). P30 is early adulthood and early disease stage; and P90 is full adulthood and late disease stage. Specifically, neurodegeneration of fast-type motoneurons starts at P30 followed by the slower types at ~P90 in this model (Pun et al., 2006; Hegedus et al., 2007, 2008). The end stage of disease in this model, when mice have developed full paralysis of both hindlimbs and fail to right themselves, occurs between P120 and P140, which is designated as P120+. Importantly, these four time points also parallel comparable disease stages of several C-bouton studies in ALS, allowing us to compare our results to the G93A line literature (Table 1).
C-bouton size is not different between WT and G93A 
motoneurons

For each time point, we compared data from G93A mice to data from age-matched, littermate WT mice. We found no significant differences in the mean cross-sectional area of WT versus G93A C-boutons (Fig. 1A). We used two-way ANOVA to examine the effects of both genotype and age on bouton size. This analysis revealed a significant effect of age ($F_{(3,262)} = 7.066; p = 0.0001$; Table 2) but no genotype or interaction effects. The age effect reflects the normal development of these boutons with age and is in agreement with the published literature (Wetts and Vaughn, 2001). Tukey’s post hoc analysis showed a significant increase in mean bouton size between motoneurons at P10 and P30 versus motoneurons at P90 in WT mice (Fig. 1A). However, Tukey’s post hoc analysis revealed no significant changes in mean bouton size between G93A and WT at any time point (Fig. 1A). Therefore, we increased the sample size of WT and G93A cells ($n$ ranged between 80 and 110 cells/group), which increased statistical power to 99.4%. Importantly, we saw no differences in C-bouton size between G93A and WT at any time point ($p = 0.21$) (Fig. 1C), confirming the analysis conducted in Figure 1A at lower statistical power. Statistical power analysis is performed to determine the sample size needed for a statistical test to detect a statistically significant difference when such a difference actually exists; a statistical power of 80% is generally accepted as sufficient.

We have also examined the number and density of C-boutons at different disease stages. The two-way ANOVA showed no effects of genotype or age but showed a significant effect of the interaction between them on bouton number and density. Fisher’s least significant difference (LSD) post hoc analysis revealed a significant decrease in C-bouton number (data not shown) and density (Fig. 1D) in WT versus G93A motoneurons only at end stage, which is in agreement with data from ALS patients (Nagao et al., 1998) and with data from two G93A studies (Chang and Martin, 2009; Gallart-Palau et al., 2014), but is opposite to data from other studies reporting either an increase or no change at end stage (Pullen and Athanasiou, 2009; Herron and Miles, 2012).

Figure 1. The size of a C-bouton synapse does not change in ALS (all sizes presented in square micrometers). A, Mean C-bouton size values per cell at four time points in WT vs G93A motoneurons, with the number of cells sampled per group inside each bar. B, Median C-bouton size values per cell at four time points in WT vs G93A motoneurons, with the number of cells sampled per group inside each bar. C, Mean C-bouton size values per cell at four time points in WT vs G93A motoneurons, using a larger sample size with $>99\%$ statistical power and with the number of cells sampled per group inside each bar. D, Mean C-bouton density values per cell at four time points in WT vs G93A motoneurons, with the number of cells sampled per group inside each bar. See Results.

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Table 2. Statistical table

| Statistical values from text | Data structure | Type of test | Power |
|------------------------------|---------------|-------------|-------|
| a. Significant effect of age, mean: $\mu_{G93A} = 7.066; \rho = 0.0001$ | Normal distribution (8/8 datasets) | Two-way ANOVA | 70% |
| b. Significant effect of age, median: $\mu_{G93A} = 7.027; \rho = 0.0001$ | Mostly normal distribution (7/8 datasets) | Two-way ANOVA | 70% |
| c. No significant effect of genotype in large-size study, $F_{G93A} = 1.575; \rho = 0.21$ | Mostly normal distribution (5/8 datasets) | Two-way ANOVA | 99.4% |
| d. Significant effect of interaction of age and genotype, mean bouton number: $\mu_{G93A} = 3.346; \rho = 0.0199$ | Normal distribution (7/8 datasets) | Two-way ANOVA with Fisher’s LSD post hoc | Genotype: 68% |
| e. Significant effect of interaction of age and genotype, mean bouton density: $\mu_{G93A} = 3.346; \rho = 0.0199$ | Normal distribution (7/8 datasets) | Two-way ANOVA with Fisher’s LSD post hoc | Genotype: 68% |
| f. Per cell analysis shows C-bouton decrease by $-8\%$; effect size, $0.51; \rho = 0.0337$ | Normal distribution (2/2 datasets) | Mann–Whitney U test | 85% |
| g. Per bouton analysis shows C-bouton decrease by $-8\%$; effect size, $0.33; \rho = 0.0001$ | Non-normal distribution (2/2 datasets) | Mann–Whitney U test | 99.9% |
| h. Large-sampling strategy shows C-bouton decrease by $-8\%$; effect size, $0.51; \rho = 0.0337$ | Normal distribution (2/2 datasets) | Mann–Whitney U test | 85% |
| i. Small-sampling strategy shows C-bouton increase by $-16\%$; effect size, $0.36; \rho = 0.0039$ | Normal distribution (one-half datasets) | Mann–Whitney U test | 99.9% |
| j. Small-sampling strategy, reduced cell count, shows C-bouton increase by $-28\%$; effect size, $0.89; \rho = 0.0038$ | Non-normal distribution (2/2 datasets) | Mann–Whitney U test | 70% |
| k. Unblinded analysis of C-bouton size shows decrease by $-20\%$; effect size, $1.22; \rho = 0.0017$ | Normal distribution (2/2 datasets) | Mann–Whitney U test | 94.4% |

Column 1 supplies the reference letter for those used in the Results section, a description of the effect measured, and the value of the statistical analysis conducted; column 2 states the data structure for that dataset; column 3 states the type of analysis; and column 4 states the statistical power for that dataset analysis.

In sum, our data could not replicate reported differences in the G93A line (Pullen and Athanasiou, 2009; Herron and Miles, 2012; Saxena et al., 2013; Milan et al., 2015) by showing no change in C-bouton size, and also showed a decrease in C-bouton number and density only at end stage.

C-bouton data breakdown, sampling strategy, and blinding influence the statistical analysis outcome

In this section, we examined the potential reasons that could (1) underlie our failure to replicate the published changes on C-bouton size and (2) explain the inconsistency in published data on this topic.

Grouping unit

Data on C-bouton size have been reported and analyzed in the literature by different units (average area per animal, average area per cell, or average area per bouton; Table 1). Thus, we examined the possibility that different methods of data breakdown could contribute to the discrepancy in published results. We further analyzed the P90 time point (an advanced stage of disease, where we might expect more detectable disease changes) and compared C-bouton cross-sectional area in WT versus G93A cells. Data were broken down and averaged in three different ways, as follows: (1) by animal; (2) by cell; or (3) by bouton (Fig. 2A). In these analyses, we used the Mann–Whitney U test because: (1) only a single time point is being considered; and (2) the U test does not require a normal distribution of data, making it more general than a Student’s t test. Generally, statistical analysis generated different results among the three breakdown strategies, despite comparable average areas being examined in each breakdown (Fig. 2A, all blue bars have similar magnitudes, as do all red bars). Although the Tukey’s post hoc analysis of the two-way ANOVA discussed in first section...
of Results indicated no statistical difference in WT versus G93A cells at the P90 time point (Fig. 1A), in the per cell analysis, the Mann–Whitney U test showed a significant decrease in G93A bouton size (by ~8%; effect size, 0.51; p = 0.0337; statistical power, ~85%; Fig. 2A, middle bars). This difference was not detected by the stringent post hoc Tukey’s test in Figure 1A, probably due to its small magnitude. The “per bouton” analysis showed a similar significant decrease in G93A bouton size (by ~8%; effect size, 0.33; p = 0.0001; statistical power, 99.9%; Fig. 2A, right bars). Conversely, the “per animal” analysis indicated no statistical difference in C-bouton size in WT versus G93A motoneurons (statistical power, 13%; Fig. 2A, right bars). This discrepancy in statistical analysis outcome is due to the large difference in sample size (i.e., the n) for each group based on the method of data breakdown (the n for each group is shown in the bars of Fig. 2). These results demonstrate that the method of data breakdown could strongly influence the outcome of statistical analysis and, thus, the reported results.

**Sampling strategy**

We also examined whether the strategy of bouton sampling from cells influences statistical analysis, again using the Mann–Whitney U test for the following analyses. We examined two strategies of sampling boutons per cell: (1) large sampling of boutons per cell (three to seven boutons measured per cell; average, 5 boutons/cell; Fig. 2B, right bars); and (2) small sampling of boutons per cell (one to three boutons measured per cell; average, 2.5 boutons/cell; Fig. 2B, left bars). Because the small-sampling strategy generated a much smaller number of boutons than the large-sampling strategy, we collected more cells using the small-sampling strategy to ensure rigorous statistical
Blinding

Blinding is the process of having the experimenter analyze data without prior knowledge of whether the data belong to the control group or the experimental group. Although blinding is recommended in data analysis to minimize potential bias, we found that this practice is not commonly reported in ALS literature. For instance, Table 1 shows that on the topic of C-bouton size change, only two among five recent studies explicitly mentioned that they blinded their analysis. It is unknown whether the other three studies performed a blinded analysis. It is interesting to observe that the magnitude of the reported changes and the effect size were generally much smaller when blinding was reported (Table 1). Therefore, we examined whether blinding the analyzer would have an effect on the outcome of analysis. We asked one experimenter to analyze a random subset of cells of the P30 time point data while blinded to animal genotype. No statistical difference was observed in WT versus G93A C-bouton size using the Mann–Whitney U test (Fig. 2C, left). However, when the same experimenter was asked to analyze the same slices, among other slices, while knowing which slices belonged to which genotype, we observed a statistically significant reduction in C-bouton size (by ~20%; effect size, 1.22; \( p = 0.0017 \)) in G93A data relative to WT data (Fig. 2C, right). Importantly, the experimenter did not know that the impact of blinding was being studied when these analyses were conducted. Notably, the magnitude of the detected decrease in C-bouton size and its effect size was largest under unblinded conditions versus our other analyses that showed a decrease in C-bouton size (20% decrease of effect size of 1.22 under unblinded conditions (Fig. 2C, right) vs 8% decrease of effect size of 0.51 with a large-sampling strategy (Fig. 2B, right) vs 8% decrease of effect size of 0.33 with per bouton analysis (Fig. 2A, right)]. These results demonstrate that blinding status can have significant impact on the outcome of data analysis. Taken collectively, these results demonstrate that methods of data breakdown, sampling strategies, and blinding can significantly influence the outcome of statistical analysis, which could contribute to the discrepancy in published results on C-bouton changes in ALS.

Reporting of blinding status is not the norm in ALS experimental and preclinical studies

Given our results on the strong impact of study design variability and the potential bias on reported data, we wanted to examine whether these issues are prevalent in the experimental and preclinical studies of ALS. To achieve that, we analyzed published experimental and preclinical articles between May 2015 and May 2016 that studied ALS using the G93A model, which resulted in 105 articles (Fig. 3A). The analysis showed that 6 of 105 experimental and preclinical articles reported blinding in all of their measurements (5.7%). Of those, 4 of 46 articles included a preclinical component and reported blinding of all of their measurements (8.7%), whereas 2 of 59 articles did not include a preclinical component or were solely experimental and reported blinding of all of their measurements (3.4%; Fig. 3A, green block). Importantly, 29 articles reported blinding in some of their measurements (27.6%; Fig. 3A, orange block). Of those, 22 articles included a preclinical component (47.8%), whereas 7 articles did not include a preclinical component or were solely experimental (11.9%). Strikingly, a large percentage of the articles (70 of 105 articles) had no mention of blinding in any of their measurements (66.7%; Fig. 3A, red block). Of those, 20 articles (43.5%) had a preclinical component, whereas 50 articles did not include a preclinical component or were solely experimental (84.7%). This analysis surprisingly suggests that blinding might not be a common practice in the ALS literature.

Reports of significant results and datasets without reported power analysis are prevalent in ALS experimental and preclinical research

Our analysis also revealed that 100 of 105 articles reported a significant result in a major outcome measure (95.2%; Fig. 3B), whereas 5 articles (4.8%) reported no significant result from a major outcome measure. In addition, 99 of the 105 articles (94.3%) reported no statistical power analysis in support of the sample sizes used to determine significant results (Fig. 3C). Six of the 105 articles (5.7%) reported a power analysis; and 1 article reported effect size (0.95%) as well.

Discussion

The present study examines potential factors that might contribute to the conflicting data reported on the disputed topic of C-bouton size changes in ALS and additionally examines this topic in our own large-sample, statistically robust study. We examined (1) C-bouton size, number, and density in WT versus G93A mice at several time
Figure 3. The majority of ALS studies do not report blinding and/or power analysis. A, The number of ALS studies on the G93A model conducted between May 2015 and May 2016 that reported blinding of their analysis vs those that did not report blinding [6 studies (5.7%) reported all elements blind, 29 studies (27.6%) reported some elements blind, 70 studies (66.7%) did not report blinding]. The analysis is further broken down by studies containing a preclinical component vs purely experimental studies. B, The number of ALS studies that reported statistically significant results vs those that reported no changes [100 studies (95.2%) reported significant results, 5 studies (4.8%) did not]. C, The number of ALS studies that reported power analysis and/or effect size calculations vs those that did not report these data [6 studies (5.7%) reported power analysis, 1 of these (<1%) reported effect size as well, and 99 studies (94.3%) reported neither). See Results.
points over the full span of disease progression in G93A mice, which serves as a model for ALS in human patients; (2) collected large WT and G93A data samples and conducted power analysis and effect size analysis; (3) examined different methods of sampling and data analysis; and (4) examined the impact of blinding. Our data showed that different statistical outcomes (enlargement, no change, or diminution) in C-bouton size could be produced from the same dataset of WT and G93A groups, depending on which experimental approaches and data analysis practices were used. We also examined effect size and power analysis, where possible, in the studies we assessed.

C-boutons are cholinergic synapses that have been suggested to increase spinal motoneuron excitability through the regulation of ion channels responsible for changes in firing rate (Wilson et al., 2004; Brownstone, 2006; Miles et al., 2007). Thus, any putative changes in these inputs (e.g., size, density per cell, activity) would appear to have tremendous implications for the firing behavior of motoneurons during disease conditions in ALS. It seems appropriate, then, that a great deal of importance was placed on the repeated detection of morphological abnormalities in these boutons on motoneurons from G93A mutant mice in past studies (Table 1). In our opinion, the failure of our own study to detect any changes in C-bouton size does not in any way decrease their significance to motoneuron excitability or their possible role in pathological states.

### C-bouton size does not differ between WT and G93A motoneurons

In contrast to a number of published reports, our analysis of C-bouton size did not show differences between WT and G93A motoneurons. This was surprising, as bouton enlargement has been reported in the literature (Pullen and Athanasiou, 2009; Herron and Miles, 2012; Saxena et al., 2013; Milan et al., 2015) and has been interpreted as an example of an ALS disease change that influences motoneuron excitability. It is important to mention that our investigation, as opposed to most studies on this topic (Table 1), was based on large samples of WT and G93A motoneurons supported by statistical power analysis and was conducted with blinding to avoid potential bias. We also followed a large-sampling strategy in collecting bouton data per cell.

### C-bouton number and density do not differ between WT and G93A motoneurons except at end stage

Similar to the size of C-boutons, their number and density is another disputed topic in the ALS field in which inconsistent results of increase (Pullen and Athanasiou, 2009; Herron and Miles, 2012; Vinsant et al., 2013; Milan et al., 2015), decrease (Chang and Martin, 2009; Casas et al., 2013; Gallart-Palau et al., 2014; Milan et al., 2015; Vaughan et al., 2015), and no change (Chang and Martin, 2009; Pullen and Athanasiou, 2009; Herron and Miles, 2012; Casas et al., 2013; Gallart-Palau et al., 2014; Milan et al., 2015; Vaughan et al., 2015) have been reported at different disease stages, sometimes all in the same study. Importantly, postmortem data available from ALS patients show a decrease in C-bouton number at end stage (Nagao et al., 1998). Our data agree with the human results and constitute one of three datasets that suggest there is a depletion of C-boutons only at the end stage (Chang and Martin, 2009; Gallart-Palau et al., 2014). Additionally, we do not see any change (increase or decrease) at earlier time points, contrary to a number of nonhuman studies. While there may be some inherent shortcomings with the animal models, inconsistent results on the same topic in the same animal model raise serious questions about the rigor of experimental design and methods of data analysis that probably contribute to unsuccessful translation (Gordon and Meininger, 2011). It is possible that widespread adoption of consistent experimental design and data analysis methods will help to clarify the comparative value of various animal models to human pathology.

### Experimental design considerations

Our results indicate that several experimental design factors could influence statistical analysis sufficiently to produce different outcomes from the same dataset.

#### Grouping unit

C-bouton size data have been reported in the ALS literature using different grouping units (by animal, by cell, or by bouton; Table 1). Our analysis indicates that changing how the data are grouped has a large impact on the significance level and outcome of the statistical analysis, even with no change in the raw averages. This effect comes from the fact that, for a given dataset, the grouping unit determines the sample size (Fig. 2A, the sample size per bouton > sample size per cell > sample size per animal), which goes into the calculation of the significance level and, therefore, influences whether a difference could be detected between the control and experimental groups. Accordingly, this issue raises the question of the proper grouping unit to be used when analyzing and comparing data. We suggest using the cell as the grouping unit and as the basis of comparison between the control and experimental groups. Cells are known to be of different types (e.g., slow vs fast motoneurons) and could be affected differently in diseases. Thus, it is plausible to expect boutons of different cell types to be affected differently, and, therefore, it becomes important not to pool boutons of all cell types. On the other hand, grouping data by animal averages lacks the adequate statistical power needed for rigorous statistical analysis.

#### Sampling strategy

Our analysis indicates that the method for sampling boutons per cell has a significant effect on the outcome of statistical analysis, such that a statistical increase between the control and experimental groups could be reversed to a decrease (Fig. 2B, an example). This factor is challenging, because most published studies do not include information on how they sampled their measurements. This issue, therefore, raises another question on the proper sampling strategy to be used when collecting bouton data from cells. We recommend collecting many boutons per cell (i.e., a large-sampling strategy) because this approach considers any potential intracellular variability among the boutons and makes the data less sensitive to errors and outliers. One potential explanation for
the conflicting data in ALS literature on C-bouton size could be due to differences among studies in the way boutons have been sampled from motoneurons. If a small-sampling strategy was followed (i.e., collecting few boutons per cell), this could underlie and explain the increase in C-bouton size that has been observed in several, but not all, C-bouton studies in ALS (Table 1). Similarly, unbiased stereology practices (e.g., random sampling of regions of interest and precise rules for marking and quantifying samples) could potentially influence analyses and should be considered and strictly applied in order to ensure unbiased quantification.

Blinding
A key issue that impacts potential bias in measurements is blinded versus unblinded data analysis. A comparison of both conditions led us to the conclusion that a lack of blinding can result in false-positive data. While raw measurements were not dramatically different between blinded and unblinded datasets, we saw statistically significant differences between groups when the assessor knew which group was experimental and which group was the control; while there were no statistically significant differences between groups when the same assessor performed the analysis in blinded conditions (Fig. 2C). This suggests that bias might affect results if the data analysis is performed without blinding. Although a number of C-bouton studies have blinded their analysis and have reported that explicitly, we found that many studies have not included any information on whether blinding was followed in analyzing their data (Table 1). Interestingly, the magnitude of reported changes and the effect size were generally much smaller in studies in which blinding was reported versus those in which blinding was not reported (Table 1). Importantly, our analysis showed that 94.3% of the examined ALS studies, preclinical and basic, do not report blinding or other procedures to limit bias. While there are no data on under-reporting that provide statistics on what percentage of studies do not report, but do perform, blinding, it is puzzling why authors would omit this important detail from their published work, if implemented. The data from our analysis do show a correlation between studies that report blinding and studies that report a small magnitude of significant results and a small effect size (Table 1).

**Effect size and power analysis**

The effect size and power analysis are two important statistical parameters that quantify the magnitude of a given change and whether the sample used is adequate to detect this change with confidence, and we recommend that these be consistently calculated and reported. Despite the importance of these parameters, we found that all C-bouton studies listed in Table 1 did not include these data. We therefore attempted to calculate the effect size of these studies from their published data; then we compared this information to our effect size data to assess the magnitudes of the reported differences. Importantly, we could not calculate the effect size for three of five studies due to a lack of information on either the SD or the sample size (Table 1). Of the two studies for which we were able to calculate the effect size, one had small effect size values, indicating a small effect of the reported changes (Milan et al., 2015; Table 1), and the second study had relatively high effect size values, although the sample size was a less than one-quarter of our dataset (sample sizes of 17–22 in Pullen and Athanasiou, 2009 vs our sample size of 80–110 in the large dataset seen in Fig. 1C). Strikingly, none of the studies included any power analysis to assess whether the sample size was adequate to detect differences with confidence. The lack of power analysis in all studies, combined with either a small effect size or an unsupported large effect size makes it difficult to assess the scientific significance of the reported findings on C-bouton size in ALS. This conclusion is in agreement with similar observations by Scott et al. (2008) on ALS preclinical studies. Our results, supported by a power analysis >99%, indicate no changes in C-bouton size in G93A motoneurons.

**Other considerations**

There are several other factors among ALS studies that could still impact the reproducibility of results: (1) background strains: although the data we collected in this study and the literature against which we compared our data were all obtained from G93A mice of the high-expressor line (with a copy number >25) to normalize the G93A expression level, studies in the literature have used different background strains of mice, which have different disease/survival timelines and might have different disease mechanisms (Table 1); (2) mutations: it is possible that transgenic animal models of ALS with different gene mutations (e.g., FUS, TDP-43, G85R, G37R) or a different copy number (e.g., G93A with a low copy number <8) could have disease mechanisms that differ among models or differ from human pathology, leading to inconsistencies in results; (3) size parameter: the C-bouton size has been assessed in literature using different measures (e.g., surface area, largest cross-sectional area, volume, appositional length; Table 1), and notably, in our study, these differing measures produced comparable results (i.e., similar differences and percentage changes between WT and G93A mice across methods) in our datasets (data not shown); and (4) sex: ALS studies looking at C-bouton changes have largely used males for their studies, and it appears that males are affected more by ALS than females (McCombe and Henderson, 2010). Because a major goal of our study was to compare our results to the larger body of literature, we preferred to use males in the present study in order to allow closer comparison. It is noteworthy that Herron and Miles (2012) found no difference in C-bouton size in female mice. These factors are important to be considered in order to have successful replication of results.

It is also noteworthy that the experimental design issues discussed here are not unique to ALS research. Similar issues have been observed in the design of studies in cancer, stroke, Parkinson’s disease, and multiple sclerosis (Sena et al., 2007; Hess, 2011). Additionally, a positive correlation was found between studies that do not report the use of practices such as blinding and power analysis with data that is not reproducible (Scott et al.,

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**Table 1**

| Study                         | Sample Size | Effect Size | Power Analysis |
|-------------------------------|-------------|-------------|----------------|
| Milan et al., 2015            | 17–22       | Low         | >99%           |
| Pullen and Athanasiou, 2009   | 80–110      | High        | <99%           |

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2008; Landis et al., 2012). Accordingly, guidelines on best practices for conducting ALS research (including recommendations on mouse model and strain use, colony management, sample sizes, blinding, and statistical methods) have been developed (Leitner et al., 2009; Ludolph et al., 2010). Also, funding agencies, such as the National Institutes of Health, now require explicit description of the use of blinding in the proposed research design as well as reporting of the statistical power of the proposed sample sizes and statistical analyses.

Conclusion

The conflicting results on C-bouton size in the ALS literature makes it difficult to assess the role of this synaptic input in the disease. Our analysis showed that factors related to experimental design, such as the grouping unit, sampling strategy, and blinding, could contribute to and explain the failure in replicating results as well as the discrepancy in published data on this topic. Furthermore, the lack of power analysis and effect size data makes it difficult to assess the scientific significance of the reported findings on this topic. Our analyses, backed by blinding practices, large samples, and power analysis, do show that the size of C-boutons does not change in G93A motoneurons throughout the disease. The number and density of C-boutons were found to be reduced only at end stage, which is in agreement with data from ALS patients. We expect that widespread adoption of consistent practices, such as those proposed here, will help to clarify many such disputed topics within both the field of ALS research and in other fields of neuroscience, leading to improved clinical translation of results.

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