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**Updating the guidelines for data transparency in the British Journal of Pharmacology – data sharing and the use of scatterplots instead of bar charts.**

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The oft-termed ‘crisis’ in reproducibility of pre-clinical investigations (Prinz *et al.*, 2011; Begley *et al.*, 2012) continues to grab the headlines, not only in scientific journals but also in the lay press (Guardian, 2015; Economist, 2017). Recent reports by eLife have confirmed that some attempts to reproduce ‘key’ cancer papers by the ‘Reproducibility Project: Cancer Biology’ were successful while others were not (Aird *et al.*, 2017; Horrigan *et al.*, 2017a; Horrigan *et al.*, 2017b; Kandela *et al.*, 2017; Mantis *et al.*, 2017). This has added further fuel to the fire that was first stoked up by the findings of Bayer and Amgen (Prinz *et al.*, 2011; Begley *et al.*, 2012; McGrath *et al.*, 2015a; Liu *et al.*, 2016; Ortuno *et al.*, 2016; Wang *et al.*, 2016). All of this activity has led to soul searching within the research community, prompting appraisal of the pre-clinical research implementation and publication processes. In particular, and relevant to this editorial, this appraisal has illustrated that ‘replication’ of pre-clinical research is not as simple a task as the word suggests. To replicate work, it is essential to have as much transparency, regarding the study and its results, as possible. Specifically, methods, tools, cells, animals, instruments, conditions, must be described in sufficient detail. This imperative underlies the need for initiatives that improve the design, interpretation and reporting of experimental data (CAMARADES; Kilkenny *et al.*, 2010). Journals and publishers have addressed many issues concerning the rigour and transparency of experimental design, and the British Journal of Pharmacology (BJP) is amongst these. As ‘starting points’, the BJP has begun to address concerns regarding the reporting of animal experiments through adoption of the ARRIVE guidelines and the development of a series of Design and Analysis guidelines for pre-clinical research (Curtis *et al.*, 2015; McGrath *et al.*, 2015a; McGrath *et al.*, 2015b). The eighteen-point Declaration (see Table 1 in (Curtis *et al.*, 2015)) has sought to strengthen the reporting and conduct of experimental design of research published in the BJP.

To further improve reproducibility of research findings, the BJP has examined two aspects of data reporting that are the subject of intense debate: 1) the extent to which raw data should be made accessible to readers and 2) the format for presenting the data in a way that reveals qualities of the datasets that underpin the validity of authors’ conclusions. Proceeding in parallel with this debate has been the stipulation by an increasing number of research councils and granting agencies that fundees comply with the FAIR initiative (i.e., that data should be **f**indable, **a**ccessible, **i**nteroperable and **r**euseable) (see ***a***). The SHERPA/JULIET database of funders’ research data policies shows that 42/60 (70%) now require authors to comply with their policy on accessible data. Here, we outline our views on how the BJP should respond to this debate.

Data Sharing

It is self-evident that any practice that increases transparency, rigour and accessibility of data will benefit both expert and non-expert communities, and should help mitigate the failures of reproducibility. However, the practicalities of data sharing are confusing and complex and the relative merits of freely-accessible data sharing, versus sharing on request, are unresolved. Moreover, what ‘data sharing’ means is often not explicitly defined: for example, how “raw” should the data be? Another problem is that standardization of data formatting and structuring will play a critical role in rendering data useful but barriers exist to achieving this. The successes in making available DNA sequences (e.g. GenBank, dbEST) and protein structures (e.g. PDB), for which the data lend themselves to standardised structuring and phylogenic profiling, will be difficult to replicate in other types of datasets, for example, the minute-or-so cellular patch-clamp traces acquired under a variety of experimental conditions or for that matter a complete set of 24-hours sleep EEG (polysomnography) recordings. Also, how might researchers be expected to accurately annotate and report the multitude of difficult-to-define determinants that contribute to particular experimental outcomes (e.g., the ‘nuisance variables’ (Button *et al.*, 2013; Krzywinski *et al.*, 2013; Voelkl *et al.*, 2016))? If a data sharing policy is to be of use, criteria must be much more explicitly defined than at present.

The natural extension to the use of Supplementary Data that often accompanies the published articles is to use digital repositories to archive and openly share research datasets (e.g. Open Science Framework, Open Microscopy, Figshare and Dryad; see ***b***). However, technological platforms that enable data sharing are not yet fully developed and few low- or no-cost repositories have been set up to make available the terabytes of data typically generated by contemporary platform technologies (e.g. high-throughput imaging systems). Making data available in accessible formats (e.g., those not requiring proprietary software files) poses a problem and, although journals should not be held accountable for ensuring that the data underpinning published content is shared, identifying which party carries this responsibility (authors, funders, publishers) remains a challenge.

The editors of the BJP and its publisher Wiley acknowledge that making available integrated platforms that link published data to the original component datasets for many types of common pharmacological data is presently not feasible. Such datasets include, for instance, ‘raw’ traces of electrophysiological measurements; large imaging files; and the reams of continuous telemetry recordings. For this reason, the BJP encourages but does notmandate data sharing. The BJP will update the readership and prospective authors of the Journal on these developments in due course.

Data presentation

Meanwhile, improvements in standards of data presentation and accessibility present a more immediately tractable issue to enhance the information in the ‘two-dimensional’ format of a research paper. In order to prepare publications, authors distil carefully compiled observations and readouts from multiple technical platforms into elements presented in tabulated or graphical form. The Editors of the BJP share concerns that this compaction may result in cardinal features of the dataset being masked, or lost altogether (Drummond *et al.*, 2011; Weissgerber *et al.*, 2015). Bar charts, typically of grouped data presented as means with a descriptor of experimental error, are the most common form of graphical visualisation in manuscripts submitted to this Journal, and are used to present results from diverse types of experiments, including measurements on humans, *in vivo* and *ex vivo* data from studies with animals, *in vitro* studies in tissues and cell lines and from the biochemical assessment of samples *(e.g. immunoblotting, RT-PCR)*. An illustrative example of a comparison of cell lines is described in the **Figure**, which shows that bar charts do not give the reader adequate information on the variability and distribution of each sampled ‘n’. This is because bar charts frequently do not adequately convey major features of the dataset. As explained below, the **Figure** illustrates why moving away from using bar charts to visualise the entire dataset is a necessary refinement that can increase the transparency and reporting of data.

The immediate conclusions that could be drawn from the data presented using bar charts in ***(i)*** are 1) that cell lines A, B and C exhibit identical mean values of receptor activation under baseline conditions (55 units); 2) there is negligible inter-population variation (inter-group Kruskal-Wallis statistic p > 0.9999); and 3) the drug has no effect in any cell line. Scrutiny of the error values may intuitively point to an increasing level of intra-group variability (A vs. B vs. C under baseline and drug-stimulated conditions) but plotting the data as a bar graph ***(i)*** masks the fact that the identical mean values of receptor activation in cell populations A, B and C are derived from values that differ considerably with respect to their ranges.

By plotting each individual ‘n’ in grouped scatter plots ***(ii),*** one sees that under baseline conditions receptor activation in cell line A is relatively homogeneous, in contrast with the broad normal distribution of activation in cell line B and the two entirely distinct sub-populations of cell line C. Note that in both ***(i)*** and ***(ii)***, the identical standard error of the mean (SEM) values before and after drug addition might (erroneously) suggest a highly uniform response of each cell line to the drug.

Presenting these data as scatter plots of paired measurements before- and after the addition of drug ***(iii)*** reveals very different responses. Visualising the data in this form leads one to conclude that the addition of drug has no effect in cell line A or cell line B. The data also corroborate the conclusion that the level of receptor activation under baseline conditions in sub-populations has no bearing on the response to the drug of those sub-populations. By contrast, the response to the drug by sub-populations of cell line C is large and depends on the extent of baseline receptor activation. This type of presentation is also valid (and useful) for data derived from human and animal experiments.

Given these issues, the Editors of the BJP now stipulate that, where possible, numerical data (whether categorical or continuous), particularly involving two sets or paired data, should be presented using scatter-plots, before-after graphs etc. in which each individual ‘n’ value is individually plotted, rather than using bar charts. Authors presenting data as bar charts should state that a scatter plot or before-after charts did not reveal unusual or interesting aspects of the data not obvious from the bar chart. We will update our Declaration with its checklist to acknowledge this change.

**Figure.**



**Figure legend.** The extent of activation of a receptor in three cell lines A, B and C under baseline (drug-naïve) conditions and following the addition of a drug is given in arbitrary units. The same data sets are presented in three different ways: ***(i)*** bar chart, ***(ii)*** grouped column scatter plot with means and error, ***(iii)*** before-after scatter plot. n=10 (i.e. biological replicates, not technical replicates). In this example data, error bars represent the SEM although authors should consider the sampling size and distribution of ‘n’ when choosing the most appropriate way of showing experimental error (e.g. standard deviation (SD) or confidence interval (CI) (Drummond *et al.*, 2011)).

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