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30th November 2015

*Nature Communications*

*75, Varick Street*

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Dear Dr. Cho,

Thank you for the invitation to revise and resubmit our manuscript. We believe we have completely addressed all the remaining comments of the referees. In doing this -- in this and the previous re-submission -- we have expended significant efforts to the extent of modifying our algorithm and reprocessing and re-analyzing *hundreds* of datasets.

Overall, we were heartened that Reviewers #1 and #3 endorsed our manuscript for publication in *Nature Communications*. However, we are rather surprised by Reviewer #2’s continued criticisms. Fundamentally, we feel that the remaining criticisms are very minor, bordering deliberate obstructionism. Nonetheless, we have tried to do all potential calculations to address the criticism. We demonstrate in our response and manuscript that the effects he/she claims are largely inconsequential to the results we report in our piece. Moreover, Reviewer #2’s comments suggest that there is a universally accepted standard for reporting allelic effects, which is simply not the case and we hope to make this clear in the response.

Now in detail, Reviewer #2 had cited two major concerns in both rounds of reviews: (a) mapping and (b) overdispersion in the datasets.

For (a), the reviewer suggests that we have missed an important source of allelic mappability bias. However, our use of the personal diploid genome is explicitly designed to compensate for the majority of this bias, which is termed the 'reference bias'. The personal genome approach accounts for additional issues beyond the reference bias, leading to better read alignment due to its ability to incorporate variants beyond just SNVs, such as indels and structural variation (as shown by Sudmant et al.) [5]. Various studies have a different take on how to account for these mappability biases (Supplementary Table 1), with many agreeing that the alignment to a personal genome, as we have done here, is a conservative and effective method for mitigating a large amount of potentially confounding bias [1, 2, 3].

Nonetheless, in this round of revision, we have strived to quantify and compensate for the remaining bias suggested by Reviewer #2, which we termed the ‘ambiguous mapping bias’ [1,4]. We show that this has a much smaller effect than the reference bias when using the personal genome approach and does not change the main results of our previous submission. In response to Reviewer #2 we have added this filter and make our approach fully compliant with what he/she sees as the ‘standard’ for the field. While small, this addition required substantial engineering effort, involving many month-long computes to reprocess all 1,263 datasets.

For (b), in Reviewer #2’s previous comments, he/she mentioned that “the correct analysis must use *some* strategy to estimate the overdispersion parameter and take it into account when testing for ASE”. Based on just this very general description, we first responded by explaining that there is indeed a range of perspectives and methods to account for the issue of overdispersion [4,6-9] (please also refer to Supplementary Table 1). We then went to great lengths to implement a novel two-step procedure to account for overdispersion in the context of our approach. In response, he/she commented that the previous methods were “mistakes” and that they “got it wrong”. We would like to point out that these methods are some of the *most current* work by *authorities* and *peer-reviewed* by colleagues in the field. More importantly, the key point that we are trying to make is *not* to show the ‘correctness’ of these methods, but to point to the broader reality that there is currently a diversity of methods in the community. For example, most recently, Castel *et al.* from *Genome Biology* [1] describes a new tool in the GATK software package and discussed the best practices for allele-specific analyses that do *not* take overdispersion into account. Van de Geijn *et al.* from *Nature Methods* [4] introduced a new allele-specific detection tool that takes into account overdispersion on a per-individual basis (similar to our pipeline; not site-specific as suggested by Reviewer #2)*.* We have also cited at least five other methods in the response that advocate for a global estimation.

Given the plurality of current approaches, the fact that the reviewer has been insisting on his/her points of view suggests his/her prejudice for a particular ‘right’ approach, when there is simply no firm consensus. Furthermore, our current approach has already been extensively discussed and utilized in the ENCODE [10], and the Epigenomics Roadmap consortia. It has also been implemented in the recent *Nature* publication by the 1000 Genomes Project Structural Variants (SV) group [5], which was the reason we initially submitted this manuscript as a companion to the 1000 Genomes paper, as the methods were extensively used by the consortium, particularly in the SV and Functional Interpretation groups.

We have made considerable efforts to modify our manuscript to take into account Reviewer #2’s criticisms while preserving our main themes. We are encouraged by the other two reviewers’ endorsements of our current manuscript and indeed believe that our approach and resource will generate considerable interest in the community. Hence, we do hope to seek your understanding and consideration of this cover letter when making your decision.

Yours sincerely,

Mark Gerstein

Albert L. Williams Professor of Biomedical Informatics,

 Co-chair of 1000 Genomes Project Consortium Functional Interpretation Group

**References:**

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