# Response Letter

### Reviewer #1 (Josh’s review)

### -- Ref 1.0 Clarifying terminology–--

|  |  |
| --- | --- |
| Reviewer  Comment | This is a very important and novel angle on interpreting the PCAWG dataset. I expect it will be read with much interest. I do find the terminology to be very confusing to follow. The definitions of passengers and drivers get adorned and blurred. I appreciate that it would seem very tricky to find the correct term to describe “impactful passengers,” which itself seems tautological. There seems to be a spectrum between drivers and passengers and, if we believe the results here, another 3rd class in the middle of the two. I found myself wondering what the difference really is between a weak driver and an impactful passenger…? The authors might play with it a bit more until they find terminology that sounds a bit more sensical. I admit that I also do not have a clear idea on what terms should be used. |
| Author  Response | We thank the reviewer for pointing out issues related to our terminology. Majority of terminology used in our manuscript is borrowed from prior literature and we cite these references. For the remaining terminologies, we define them upfront and use these consistently throughout the text to avoid any confusion. We also provide detailed definition of all these terms in our supplement section.  Mutations with weak effects on fitness are assumed to have a negligible impact on tumor growth and are termed here as “nominal passengers” (i.e. all mutations other than drivers in a cancer). In contrast, an “impactful nominal passengers” here refers to a subset of “nominal passengers”, which have high predicted molecular impact scores and thus might play a role in tumor growth. We suggest that, through aggregated effects, such mutations can play weak driver roles and thus be subject to weak positive selection (or negative selection in the case of deleterious passengers).  In this manuscript, we considered weak drivers as a subset of impactful passengers. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.1 – Background model--

|  |  |
| --- | --- |
| Reviewer  Comment | The major sticking point of this work is the definition of the background model that determines their noncancerous set. The authors state that these are created by shuffling the mutations while preserving various features such as overall burden and mutation signatures. While this is convincing, one wonders if there are other possible confounders since the genome is so architected. Could the authors at least address the role different chromatin states might (or might not?) influence the null model? It could be that if they accounted for these effects that the entire signal would drop away. But even then, perhaps the authors could argue that the architecture itself somehow influences the accumulation of such passengers? |
| Author  Response | We thank the reviewer for identifying potential limitations in the background model. In the previous version of this work, we applied a nonparametric null model, which preserves the signature and local burden. We believe this to implicitly reflects chromatin states as well as many other covariates whose influence clusters locally. This null model has been applied across the PCAWG project. However, based on suggestions by all reviewers, we now also apply a null model that considers additional covariates *explicitly* such as chromatin openness, replication timing, and GC content. We observe a slight increase in the variance explained by our model with these explicit covariates, suggesting the signal is robust to the effects mentioned.  [[SK2MG: We still need to figure out if we are going to stick with Inigo’s simulation or the moat-sim simulation, which has better signal for additive variance analysis.]] |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.2 – Comments on the setup of the paper--

|  |  |
| --- | --- |
| Reviewer  Comment | Abstract and Introduction are very well written and intriguing. Pitched with the right amount of background, motivation, controversy, and reservation. |
| Author  Response | We thank the reviewer for this positive comment. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.3 –Clarifying random effects model parameters--

|  |  |
| --- | --- |
| Reviewer  Comment | I believe the phenotype variable y[j] records if the sample is a sample from PCAWG (y[j]=1) versus a randomly generated sample (y[j]=0). The authors should make this just a little more explicit. |
| Author  Response | As per suggestion, this is now explicitly defined in the updated text. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.4 – Rationale for using random effects model --

|  |  |
| --- | --- |
| Reviewer  Comment | Page 3. Not clear why the particular model used was chosen. Is this standard from the GWAS community or is it the idea of the authors? Either provide a citation or refer the reader to the appropriate part of the supplement that gives justifications for the form chosen. |
| Author  Response | This particular model is commonly used in complex trait analysis – a recent approach (PMC3232052) which used it to explain variability in human height has been adapted for many studies. The first supplemental note referred to on page 3 addresses suitability of this for somatic mutation in cancer. The model is appropriate for detecting the cumulative effect of variants on tumorigenesis which may be individually weak. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.5 – Related to signature analysis --

|  |  |
| --- | --- |
| Reviewer  Comment | I don’t quite follow the argument and interpretations under the mutational signatures section. Are there certain signatures that will lead to stop codons? Which ones are these and can they be noted in Fig 3? Or are the authors looking at associations of any mutational signature? I lose sight here of how their nominal passengers have been used in this analysis if at all. |
| Author  Response | In the mutational signature section, we analyze the role of mutational signatures to the differential burdening of genomic elements by nominal passengers. We perform this analysis for the coding LoF mutations, as well as for non-coding mutations leading to TF motif break events.  Reviewer’s interpretation is correct. For LoFs, we exclusively look at mutational spectrum in the context of “nominal passenger” mutations altering stop codons. In Figure 3a, mutational spectrum is plotted for mutations leading to LOF events.  Furthermore, we also compare the difference in mutational signature for “nominal passengers” with low and high impact scores for various PCAWG cancer cohorts. Categorization of nominal passengers into low and high impact score groups was done based on functional impact score threshold. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 1.6 – Clarifying comparative statements --

|  |  |
| --- | --- |
| Reviewer  Comment | There are quite a few comparative statements where one of the classes being compared is implied and this leads to an ambiguity and lack of clarity in following the logical arguments of the manuscript in many places. For example, statements like “As expected, we observe lower mutational heterogeneity among high impact nominal passenger SNVs,” are hard to decipher because its not clear what is lower? To random? To low-impact nominal passengers? To drivers? What? |
| Author  Response | The reviewer makes a good point. In the updated version, we explicitly state these comparisons. In the particular example the reviewer mentions; the intended comparison is to low-impact nominal passengers rather than to random set. |
| Excerpt From  Revised Manuscript |  |

### Reviewer #2 (Peter’s comments)

### -- Ref 2.0 Overall comment–--

|  |  |
| --- | --- |
| Reviewer  Comment | The paper is considerably improved from earlier versions we have seen. In particular, I like the general concept of estimating the size of the set of unobserved driver mutations using random effects models. This is potentially very powerful, but requires rigorous attention to detail in its specification. |
| Author  Response | We thank the reviewer for the encouraging comments and highlighting the importance of this work. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.1 – Clarifying the model--

|  |  |
| --- | --- |
| Reviewer  Comment | I didn’t have the Supp Notes in reviewing this, but if I understand correctly, the model fitted is basically a GLMM? That is, basically a random effects logistic regression (in which case the formula y\_ij = … should have the logit function specified)? |
| Author  Response | The Model is a GLMM (General linear mixed model) with random effects, where the random effects (SNV effect sizes) are modeled as Gaussian distributed, and are not estimated directly but integrated across when estimating the variance explained by the SNVs. As in previous analyses which use this model for complex trait analysis (PMC3232052), we do not use a logit linking function as would be used in logistic regression, but use the linear outputs directly to predict the binary phenotype. The additive variance thus estimated is referred to as using the ‘observed scale’. We also quote results on the ‘liability’ scale, which uses a probit link function. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.1 – Clarifying the model--

|  |  |
| --- | --- |
| Reviewer  Comment | It is difficult to assess without the Supp Methods, but how well have the authors tested / controlled for over-fitting? That is, the statement of 64.5% variance explained with all mutations versus 52.5% with drivers only could be due to the vastly larger numbers of data-points in the former analysis – even small mis-specifications could accumulate to lead to apparent better predictive power. Separate test-retest cohorts are essential here. |
| Author  Response | The issue of overfitting on the SNV effect sizes does not arise in the random effects model, since the model is not trained as a predictive model. Instead, these parameters are integrated across in order to estimate the single hyper-parameter which controls their variance. In its simplest form, the model thus uses only one degree of freedom to model the SNV effects regardless of the number of SNVs, and hence does not suffer from overfitting as more SNVs are added. We verified this directly through simulations, which show that adding Poisson-distributed SNVs to the model lowers the estimated additive variance.  In addition, we also performed sensitivity analysis to cross-check the issue of over-fitting in our random effects model. We computed the additive variance of two random samples. If this model were overfitting the data, one would get non-zero additive variance for such a double random dataset. In contrast, we observe 0% additive variance in all cancer cohorts, suggesting no overfitting in our analysis. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.2 – Improving background model --

|  |  |
| --- | --- |
| Reviewer  Comment | I remain concerned about the generation of the null model samples. Any factor that influences true passenger mutation distribution that is not accounted for in the null model redistribution will have the potential to get picked up by the additive model as containing predictive power, but not for the ‘functional impact’ reasons, but rather for ‘uncorrected information in mutation signatures’ reasons. In particular, the following factors could well play a role and should be included in the null model redistribution:   * 1. Replication timing (especially likely to have an effect)   2. Intergenic versus intragenic; Gene expression   3. Chromatin openness   4. Replication and transcription strand (less concerned about this)   5. Nucleotide context beyond trinucleotide (especially problematic for POLE hypermutators and also the UV light signature). |
| Author  Response | We update our null model to include some of suggested covariates. We also use penta-nucleotide context for the melanoma and liver cancer cohorts. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.3 – Background model related issues--

|  |  |
| --- | --- |
| Reviewer  Comment | Many of the downstream analyses depend critically on the accuracy of these models – even small inaccuracies in inferences can lead to quite large numbers of, for example estimated weak drivers or negatively selected mutations (Figure 5) when multiplying up by the total number of mutations and/or number of samples. This is particularly concerning for the melanoma and liver cancer findings in Figure 5. The authors will need to be scrupulous in reassuring the reader that these calculations are valid. |
| Author  Response | We updated our downstream analysis based on the new null model, which corrects for the covariates suggested by the reviewer. In addition, we apply additional filters (used in the main driver paper) on our putative list weak drivers and deleterious passengers. This provide a very conservative estimate on the number of weak drivers and deleterious passengers. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.4 – Terminological inconsistency --

|  |  |
| --- | --- |
| Reviewer  Comment | The Introduction and much of the paper suffers from terminological difficulties – for example, the phrase “…and large numbers of passengers (with weak or neutral fitness effects) is analogous to…” [Page 2] is not accurate. By definition, passengers cannot have weak fitness effects – this would make them drivers. It is, however, perfectly reasonable to ask whether mini-drivers exist, and what they might look like if they do. |
| Author  Response | We agree with the reviewer that there are challenges in defining terminology. We explicitly use the term “nominal passenger” throughout the text. Nominal passengers correspond to all non-driver variants. These potentially include weak drivers, neutral passenger and deleterious passenger. In Figure5, we explained this in detail. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.5 – Missing heritability relevance --

|  |  |
| --- | --- |
| Reviewer  Comment | I didn’t find the GWAS analogy in the Introduction especially helpful. Whatever the truth about the driver – passenger – deleterious mutation landscape in cancer, it is clearly vastly different to the complex diseases studied by GWAS. Also, whether there is the somatic equivalent of “missing heritability” is moot – it is not at all clear to me that we are missing anywhere near the same proportion in cancers as in GWAS. |
| Author  Response | The Driver discovery exercise in PCAWG suggest that we don’t find many non-coding drivers in PCAWG cohorts as well as on pan-cancer level. This can be potentially attributed to the lack of power for identifying non-coding driver due to limited sample size. In this context, our additive variance analysis provides an alternative approach to address this issue.  In complex trait analysis, it was strongly felt that there must exist many SNPs of low-penetrance or low effect because of the missing heritability problem. This motivated the development of the random effects model to measure the aggregate effect of variants in a statistically rigorous way. If there are variants of low effect in cancer, the random effects model is a powerful way to find their aggregated, where the additive variance here measures the collective predictive power of the variants against an appropriate null model.  In our updated text, we emphasize that additive variance in our model does not directly measure heritability as in the GWAS case, but rather the combined effects of SNVs on tumorigenesis. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.6 – Suggestion on modifying introduction --

|  |  |
| --- | --- |
| Reviewer  Comment | Instead, in the Introduction, I would sharpen the discussion about the methodological constraints of current approaches to driver-passenger dichotomisation and whether there might be fundamental limits to recurrence-based methods (especially given the long tail of cancer genes). Make the point that such limits might mean that an undefined proportion of rare drivers might be being mis-annotated as passengers, and with current sample sizes, we are likely only to be able to detect a global signal of their existence (but this is hugely challenging), rather than identify each one specifically. |
| Author  Response | We thank the reviewer for this suggestion. We include some of these points in our updated text. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 2.6 – Suggestion on modifying introduction --

|  |  |
| --- | --- |
| Reviewer  Comment | Furthermore, make the point that selection acting on somatic cells is dynamic, and that just because something is a passenger now, doesn’t mean that it couldn’t become a driver when treatment is given or the clone spreads to another organ – therefore of value to study how many of them have functional effects, even if these functional effects are not of selective consequence currently |
| Author  Response | We thank the reviewer for this suggestion. We include some of these points in our updated text. |
| Excerpt From  Revised Manuscript |  |

### Reviewer #3 (Gaddy’s review)

### -- Ref 3.0 Issue with background model–--

|  |  |
| --- | --- |
| Reviewer  Comment | In their paper “Passenger mutations in 2500 cancer genomes: Overall molecular functional impact and consequences”, Kumar et al. present a very intriguing analysis of putative passenger mutations and their potential functional impact in cancer. Overall, the paper presents a very interesting analysis with important implications to our understanding of positive and negative selection in cancer.  My main concern about this paper remains the uncertainty in our background models. The use of the additive effect model is very interesting. However, as opposed to GWAS studies where one uses cases and controls, here the authors use a simulated dataset of non-cancer “neutral” cases using a mutation randomization scheme. |
| Author  Response | We agree with the reviewer’s comment regarding potential limitations in the background model. As noted earlier, we update our background model to address some common issues brought up by all reviewers. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.1 Sensitivity analysis of additive variance model–--

|  |  |
| --- | --- |
| Reviewer  Comment | Calculate additive variance for double randomized sample set, i.e., use two iteration of random sample to calculate additive variance for different cancer cohort. |
| Author  Response | As per suggestion, we performed sensitivity analysis to cross-check the issue of over-fitting in our random effects model. We computed the additive variance of two random samples. Across cancer types, we observe ~0% additive variance suggesting no overfitting in our analysis. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.2 Sensitivity analysis of background model–--

|  |  |
| --- | --- |
| Reviewer  Comment | Calculate additive variance with different randomized sample set. Vary the length of local window to generate multiple random set and compute the additive variance. |
| Author  Response | As per suggestion, we performed sensitivity analysis to cross-check the influence of background model on additive variance. We generated two distinct randomizations set within a local window length of 50kb and 100kb. Overall, there is a slight variation in the total additive variance calculated based on these distinct randomizations set. This can be attributed to small effect due to various genomic co-variates.  We also note that in our updated analysis, we perform additive variance on a separate background model, which corrects for various covariates suggested by all the reviewers. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.3 – Some limitations in current background models--

|  |  |
| --- | --- |
| Reviewer  Comment | My concern is that we (the community) are far from understanding the background model and even our best randomization scheme does not truly reflect the underlying processes which can vary at the single-base level (as opposed to the 10-100kb resolution of the randomization scheme). As we are seeing in the PCAWG drivers paper, even after controlling for many effects, there are still significant hits that are false positives due to many effects, such as (i) ignoring the lack of repair due to DNA binding proteins (most prominent in melanoma); (ii) inaccurate modeling of AID and APOBEC processes; (iii) Not taking into account the amount of DNA available for mutagenesis during tumor development (e.g. if a chromosomal region is lost early on, the rate of mutations in that region will be lower since there is only one copy to mutate); (iv) our ability to detect mutations as a function of sequence coverage (ie. GC-content affects coverage and our ability to detect mutations). Promoters typically have high GC-content, hence we will detect fewer mutations in them (whereas the simulated datasets will not have the same decrease in density); (v) ignoring local DNA structure such as palindromes and other sequence motifs; (vi) ignoring the association between signatures and timing; and (vii) ignoring different repair processes that operate different at various scales and in different genomic regions (introns, exons, intragenic regions, early vs. late replication timing).  The uncertainty and inaccuracy of the background model can lead to incorrectly reaching the conclusion of positive selection or negative selection (depending on the genomic regions and mutational processes that operate in the specific tumor type). |
| Author  Response | Thanks to these comments, our updated background model now incorporates many of these features explicitly. We present results from both models in the supplement and main text. By comparing the results from the two models, the reader can appreciate how further, yet undiscovered covariates might continue to confound the results. Furthermore, we apply some of the suggested filters on our putative list of weak drivers and deleterious passengers to provide a more conservative estimate. In addition, we also mention these limitations explicitly in our discussion section. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.4 – Consistent use of terminology--

|  |  |
| --- | --- |
| Reviewer  Comment | The authors define “nominal passengers” but it is unclear exactly how these are defined. It is critical for the rest of the paper. I would remove any event in a very inclusive list of cancer genes and not only the ones that we detected as drivers using the PCAWG dataset. I think the term “potential passengers” is better.  It is crucial that the term is used throughout the manuscript. There are several places where they are referred to as just “passengers”. This is confusing since true passengers cannot be weak drivers, but potential/nominal passengers can. |
| Author  Response | “Nominal passengers” include all mutations that are not identified as drivers in PCAWG, or previously known to be cancer driving events.  We agree with reviewer’s comment to consistently use the terminologies across the manuscript. We have updated the manuscript accordingly.  We also would like to point out that some of our terminologies are borrowed from prior literature and we cite them accordingly. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.5 – Additive variance in coding region--

|  |  |
| --- | --- |
| Reviewer  Comment | If the authors focus only on coding regions, are the results of the random effects model consistent with significance analyses such as MutSig and dN/dS ? |
| Author  Response | We now perform this comparison directly using the new nested model results. We compare our list of weak driver genes with the gene set curated by the driver group. We specifically look for overlap between our weak driver genes and driver discovery gene set, which didn’t satisfy the statistical significance criterion during the driver discovery process.  Presumably, some of these genes are weak drivers and failed the statistical significance threshold due to limited cohort size and thus insufficient power. We find good overlap between these two-list validating our approach to certain extent. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.6 – Impact score and additive variance model --

|  |  |
| --- | --- |
| Reviewer  Comment | It is not exactly clear how the authors use the FunSeq score to derive the z\_ij. How would the results change if only evolutionary conservation is used? (Even evolutionary conservation has its own problems since some of the same processes that affect cancer shape evolution)What are the underlying features that contribute to the peaks in the impact score (conservation? specific chromatin marks? alteration of TF binding sites?). |
| Author  Response | We have attempted to explain this more clearly in the manuscript. The FunSeq threshold is treated as an additional optimization parameter. We also compared our results with a model optimized by conservation score (e.g. GERP) and found no significant difference in total additive variance explained.  The different peaks in the impact score distribution can be attributed to combinations of features (conservation, chromatin marks, and TF motif alterations). |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.7 – SV impact score definition --

|  |  |
| --- | --- |
| Reviewer  Comment | It is unclear how the impact score for SVs was defined. |
| Author  Response | We have provided a detailed description in the supplemental methods section. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.8 – Singletons and additive effect model --

|  |  |
| --- | --- |
| Reviewer  Comment | How many of the variants are singletons (ie. occur only in one sample across the cancer and “neutral” cases)? How do they affect the model and its predictive power? |
| Author  Response | Singletons cannot increase the additive variance, since they do not contribute to genetic relatedness. As noted in response 2.1, we do not train a predictive model, so the issue of overfitting does not apply to the SNV effects. We verified directly through simulations that adding singletons decreases the estimated additive variance. Hence, we remove singletons for all additive variance calculations in the paper. We note that, for the SNV-level model, singletons are SNVs occurring in only one sample, while in the gene-level model, they are genes which have non-zero mutation burden only in one sample. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.9 – Signature and impact score distribution --

|  |  |
| --- | --- |
| Reviewer  Comment | Not clear what is the contribution of different mutational signatures to the 3 peaks on functional impact. |
| Author  Response | We address this question in Figure3d, which shows the signature differences between nominal passengers with high impact score (mutations present in second and third peak) and low impact score (mutations present in the first peak). As noted, these differences vary between different cancer types and we highlight some of these differences in updated text more explicitly. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.10–Potential role of signature driving some observation--

|  |  |
| --- | --- |
| Reviewer  Comment | It is concerning that many of the results are in tumor types with specific signatures that we are probably not modeling accurately, such as UV, lymphomas (canonical and non-canonical AID), lung cancer, liver cancer, and esophageal cancer. |
| Author  Response | We would like to emphasize that the main aim of our work is to characterize “nominal passenger” landscape in PCAWG. For this purpose, we specifically look at their overall burden among different genomic elements and their predicted molecular functional impact.  As the reviewer points out, for certain cohorts it’s very likely that signature plays an important role in influencing our observations. We think this is perfectly fine as this provide mechanistic understanding to some of our observations. Moreover, this is one key reason why we look at mutation spectrum and signature differences among “nominal passengers”.  We would also like to point out that additive variance analysis is a subset of many analyses performed in this work. This analysis only addresses a specific question related to “nominal passengers (i.e. whether their cumulative effect has any role in cancer progression). For this purpose, we use an updated random model, which takes into account of many mutational processes mentioned by the reviewer.  In the updated manuscript, we clarify this point more explicitly. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.11 – comparison of germline and somatic SVs --

|  |  |
| --- | --- |
| Reviewer  Comment | I am not sure that the comparison to germline for SV is meaningful — there are very different patterns of SVs in germline and somatic. It is also unclear what is the random model (is it uniform?) |
| Author  Response | The goal of this analysis was to highlight the observation that both germline and somatic large deletions prefer to engulf genomic elements rather than partially break it. This is very interesting and provides mechanistic insight into how different categories of SVs burden various genomic elements. As the reviewer points out, we use a uniform randomization scheme to perform this analysis. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.12 – Essential gene and LOF analysis --

|  |  |
| --- | --- |
| Reviewer  Comment | What is the list of essential genes that is used? Keep in mind that essential genes can be expressed at higher levels and therefore subject to different DNA damage and repair mechanisms (such as transcription-coupled repair and transcription-coupled damage (in liver cancer)). |
| Author  Response | The essential gene list was based on previous publication “Essential genes - CRISPR knockouts in four cancer cell lines”.  The reviewer suggests a good point, which we explicitly mention in the updated draft. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.13 – Consistency in PCAWG wide terms --

|  |  |
| --- | --- |
| Reviewer  Comment | We need to be consistent across PCAWG papers. We typically use SCNA (somatic copy number alteration) rather than CNV. |
| Author  Response | The reviewer makes a good point here. We update the text accordingly to keep the nomenclature consistent with other PCAWG papers. |
| Excerpt From  Revised Manuscript |  |

### -- Ref 3.14 – TFBS related analysis --

|  |  |
| --- | --- |
| Reviewer  Comment | I am skeptical about the analysis of hits in different binding sites of a TF. There is no correction for signatures. I believe this is mostly mechanistic, e.g. CTCF binding sites in liver cancer. |
| Author  Response | We concur with reviewer that TF related analysis is mechanistic and not trying to highlight any role of selection. This analysis is aimed towards highlighting the differential burdening of various TFs and the corresponding gene regulatory network. We have modified the main text to clarify this point more explicitly. |