**Title:** The “*F-*word”: Next-generation data *F*iltering in the genomics era

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**Abstract**:

Genomic data are now ubiquitous in studies across disciplines including human health, agriculture, biodiversity, molecular ecology, and evolutionary biology. While genome-wide data can improve parameter estimates and resolve previously intractable questions, data filtering choices can significantly influence downstream analyses. Here we review recent literature across disciplines to assess the commonly applied filter types and thresholds, including minor allele frequency/counts, missing data, linkage disequilibrium, and Hardy-Weinberg deviations. Surprisingly, we found that a large percentage of studies did not report key filtering parameters. We also illustrate large effects of filtering thresholds on statistics such as Tajima’s D, *F*ST, and effective population size (*N*e). Given the current state of affairs, we propose two key principles for investigators to keep in mind when analyzing genomic data: 1.) different filtering choices can result in different results such that two researchers analyzing the same dataset, but using different filters, could reach entirely different conclusions, and 2.) distinctively filtered versions of the same dataset should be used to address specific questions and to quantify the effects of filtering. We conclude by providing recommendations for standardizing filtering approaches to usher in the next-generation of data filtering.

**Introduction:**

Recent and rapid advances in both short and long-read sequencing technologies (*e.g.*, Illumina and PacBio) have resulted in the proliferation of large genomic datasets1–4. All high-throughput (“next-generation”) sequencing methods result in large numbers of reads that, importantly, do not all have low error rates5,6. Errors can be further inflated when these sequences are aligned to a reference genome or transcriptome, particularly if the reference is incomplete or assembled ***de novo*** from the reads themselves7–9. Therefore, most investigators perform multiple types of data ***filtering*** for quality control prior to data analysis, including filtering on read quality, mapping quality, read depth/coverage, minor allele frequency or minor allele counts, missing data (across loci and/or individuals), and deviations from Hardy-Weinberg or linkage equilibrium. Despite the ubiquity of filtering in genomics, no comprehensive review currently exists on trends, standardization, or best practices for data filtering.

Genomic filtering is not a trivial problem. Indeed, it has been aptly termed the “*F*-word” by geneticists due to how *F*lipping challenging it can be to conduct and understand effects of *F*iltering on downstream conclusions10,11. Here, we define ***filtering*** as the intentional removal of bases, reads, individuals, and/or ***genetic variants*** from a genomic dataset with the explicit goal of improving data quality for downstream analyses. Filtering is distinct from other forms of data processing in that it centers on the removal of data (as in “filtering out”), whereas other approaches, such as ***imputation***, modify the data directly without removal. Filtering is an issue of paramount importance because: 1) every genomic dataset must be filtered, often repeatedly, 2) the same dataset filtered in different ways often yields entirely different results12, and 3) filtering choices can be confusing, are often subjective, and currently lack standard or agreed-upon guidelines13.

Filtering approaches are often inadequately described and vary widely among studies. We reviewed 173 recent empirical genomic research articles from the fields of agriculture/domestication, biodiversity studies, and human health, and found that the use of crucial and common filters were not mentioned in 17-56% of studies, depending on the filter (Box 1). Importantly, it was often not possible to tell if these papers conducted certain filtering steps and simply neglected to mention them, or if they skipped the filtering steps entirely. Furthermore, even when filters were mentioned, the methodological details used were rarely reported (see Box 1). Among papers that recorded filtering thresholds, the values used ranged several-fold, suggesting limited standardization of filtering efforts. Qualitatively, we also observed that the vast majority of studies did not provide justifications for their filtering choices, often using the default program settings with no further comment. Our literature review indicates that: 1) no comprehensive sequence data filtering standards or best practices exist, 2) the impacts of filtering are rarely quantified (and thus are poorly understood), and 3) many published studies are not reproducible due to insufficient methodological details.

To help investigators improve their filtering methodologies, we review different strategies for filtering genomic datasets, examine their downstream effects, and provide practical recommendations including a detailed flowchart illustrating a typical order of operations. We also provide example tables demonstrating how to document filtering steps and a checklist to consult before publication, which should be useful for both investigators and biology journals.

**Common filters: complexity and importance**

Today, investigators have many different instrumentation options for obtaining DNA, RNA, and epigenetic sequencing data14,15, which subsequently yield reads of different lengths, configurations (*e.g.,* single- or paired-end), and qualities16. These sequences are then either aligned back to a ***reference*** (such as a reference genome or transcriptome), or *de novo* assembled and aligned to consensus sequences (*i.e.*, contigs or stacks) for subsequent variant calling or analyses17,18. Without some form of reference-guided alignment, downstream analyses can be limited, since it can be difficult to estimate relevant parameters (*e.g.*, runs of homozygosity19) or determine the genomic context for loci of interest (*e.g.*, linkage)20 (but see21 and 22 for best practices in metagenomics and eDNA, respectively). Lacking a quality reference genome therefore represents a challenge for researchers working in non-model systems, where alignments are often made to low quality reference genome assemblies (*e.g*., high contig counts with low N/L50 scores23) or even to different, often distantly-related species24,25. Given these many challenges and technical limitations, errors are always introduced to some degree during both sequencing and alignment, and these errors must be accommodated – usually via filtering – to avoid downstream bias.

Many types of population genetic or statistical analyses should be conducted only after applying specific filters, and the results from certain analyses can suggest the need for additional or modified filtering strategies. For example, unexpected results from exploratory methods such as a principal component analysis (PCA) can be indicative of experimental or laboratory errors (*e.g.*, mislabeling), sequencing bias, sex-linked loci, selection, or other phenomena26–28 and thereby suggest the need for further filtering steps. Thus, the process of filtering begins immediately after sequence data collection and typically may not end until all analyses are completed. An exhaustive list of filters, their descriptions, and the typical stage in the workflow where they are applied can be found in Table 1.

In this section, we introduce and review several of the filters that often have the largest downstream effects. Specifically, we discuss filters for: read and mapping quality, ***read depth***, ***missing data***, ***Minor Allele Frequency (MAF)/Minor allele Count (MAC)***, ***Hardy-Weinberg proportions***, and ***linkage disequilibrium***. These filters can be generally divided into two sequential categories: pre-variant filtering (*e.g.*, read/mapping quality and read depth) and post-variant filtering (*e.g.*, minor allele counts, MAF/MAC, missing data; see Table 1). Pre- and post-variant stages generally coincide with pre- and post-VCF-file stages (or genotyping).

*Pre-variant filtering:*

Prior to ***de novo assembly*** or alignment to a ***reference***, data are usually filtered via the removal of low-quality reads (see Table 1). This initial filtering may have downstream effects, and researchers across disciplines often use different and potentially arbitrary definitions of what constitutes a low-quality read (see Box 1). Low-quality reads can also be removed during the process of alignment itself, since different alignment algorithms can prevent sequences from mapping back to a reference if their quality scores are below user-defined thresholds29–31.

Conversely, high-quality reads may also be excluded if the reference does not contain the sequence or if their sequence is highly repetitive (*e.g.*, transposable elements) or found in multiple genomic locations (*e.g.*, paralogs; see Figure 1A). To aid reviewers and other researchers in understanding the impacts of these filtering decisions, investigators should always report: 1) the percentage of reads that were removed prior to alignment and 2) the percentage of reads that mapped successfully and uniquely (to one location) on the reference.

Reporting these statistics can help reviewers assess the quality of the data underlying the study and allow future investigators to determine whether alternative filtering could address additional questions (*e.g*., re-filtering to not remove reads that map to multiple locations could identify paralogous loci or transposable elements). Note, however, that even a ***strong alignment*** of putatively high-quality reads does not always mean a *correct alignment,* since reference bias32, genome assembly errors33, ***structural variants***34, and challenging alignments (*e.g.*, transposable elements35, PCR duplicates36, and copy-number variants37) are present in most genomic datasets (Figure 1).

*Post-variant filtering:*

After pre-variant filtering, genomic work-flows typically proceed with **genotyping**, whereby genetic variants such as **single nucleotide polymorphisms** (SNPs), insertions or deletions (indels), andstructuralvariantsare algorithmically identified with software such as GATK38, ANGSD39, STACKS9, ipyRAD40, or LUMPY41. During this process, the read depth (or coverage) of each locus must be considered, since high depth of coverage allows for more confidence in genotyping (and subsequently downstream inferences). These algorithms typically either mark genotypes as missing if below a certain read depth or if they have poor quality scores caused by low read depths42. The depth and/or quality filters used at this step vary substantially across studies (see Box 1). Note, however, that well-developed approaches to make use of low coverage sites do exist39, and so filtering out such loci is not *always* necessary.

While many different ways to filter genotypic data exist (see Table 1), we will focus here on four widely used filters (Box 1; Figure 2). Below, in the “Solutions and Best Practices” section, we suggest how to best implement these filters.

***Missing Data***: filtering out loci or individuals with a user-defined amount (or proportion) of missing data. An excess of missing data can indicate something awry with sample collection or preservation, genomic library preparation, or alignment, all of which can obscure patterns of variation43. The filtering choices used for missing data vary *widely* among studies and the downstream consequences are rarely evaluated (Boxes 1, 2).

***Minor Allele Frequency (MAF)***: filtering out loci (typically SNPs) where the less frequent allele (*i.e.*, the minor allele) occurs below a certain frequency. MAF filtering is often based on the assumption that **singletons** or **other rare variants** that occur at a frequency of less than ~5% are the product of genotyping errors. MAF filtering is therefore typically done to this specific threshold (Box 1). Depending on the analysis and objectives, this filter can be applied study-wide (*e.g.,* globally across populations) or within each population.

***Minor Allele Count (MAC)***: an alternative to MAF filtering wherein loci are removed based on the absolute count of the minor allele rather than its frequency, allowing for more consistent filtering across samples of different sizes (although arguably producing an uneven MAF filter across those same samples). A MAC may be preferable over a MAF to remove only singletons (or doubletons, *etc.*) or when small samples of individuals exist while filtering within populations or sample groups.

***Hardy-Weinberg Proportions (HWP)***: filtering out loci based on statistically significant (for a given alpha/*p*-value; see Box 1) deviations from Hardy-Weinberg proportions. HWP is a common assumption of many downstream analytical tools (*e.g.,* STRUCTURE44), and removing loci that violate HWP can help ensure unbiased results for downstream analyses45. Correcting *p*-values for multiple tests in datasets with many loci is rarely performed46. Researchers who do correct for multiple tests should explicitly report the reasoning behind the correction method they use, which can vary in stringency (from Benjamini–Hochberg47 to sequential or simple/stringent Bonferroni48). Note, however, that more stringent multiple testing correction for HWE paradoxically *decreases* filtering stringency overall, since more loci will be found to be non-significant and thus retained. Ultimately, different approaches and alpha thresholds should vary depending on the questions being asked and the tolerance for including problematic loci49 (Box 3). HWP deviations can often reflect sequencing, assembly, or alignment errors (such as a heterozygote deficit caused by allelic dropout or a heterozygote excess caused by paralogous regions; see50–52). However, loci out of HWP can be indicative of real, biological phenomena, such as cryptic population sub-structuring (Figure 2), or balancing selection. As a result, it is crucial to filter for HWP within sample groups (*e.g.*, populations) rather than study-wide (*e.g.*, globally on all samples; see *Study-wide versus within sample-group filtering*, below)53 and to do so with a low stringency if loci under selection or those that differ between populations are of interest.

***Linkage disequilibrium (LD)***: pruning clusters of loci that are in substantial LD with each other down to a single locus to ensure statistical independence among loci - a common assumption made by many downstream methods. For example, methods based on the site frequency spectrum (SFS) of a population may be biased if allele frequencies in a variant-rich region differ from the genome-wide average. Similarly, failure to remove non-independent (linked) loci can bias estimates of parameters like effective population size (*N*e)54. However, filtering out SNPs based on LD could also strongly influence diversity estimates (such as the number of segregating sites across genomic regions) or inadvertently cause investigators to overlook important structural variants (Figure 2). Studies that lack a high-quality reference may also require LD filters to ensure independence of loci or contigs55, which can be accomplished through pairwise correlation measures such as Pearson’s *r*2. Alternatively, many investigators working with *de novo*-assembled datasets simply extract a single SNP from each contig to mitigate the effects of linkage (though this assumes distinct stacks/contigs are themselves unlinked). Corrections for multiple testing are also important for LD filtering, but seldom reported46.

**Effects of filtering:**

The effects of filtering are often unappreciated and unknown in genomic studies. While *concerning*, this is not particularly *surprising*, since1) many different filtering approaches exist, 2) filtering requires non-trivial time and computational resources to perform, and 3) many individual filters can be applied with different thresholds and at different data processing stages (Table 1). Indeed, circumstances exist where the same type of filter can, and possibly should, be applied at multiple stages of the bioinformatic process (Table 1). Furthermore, many types of filtering occur during the “black box” of alignment and genotyping, leading many investigators (ourselves shamefully included) to use default settings and not think about the downstream consequences. This “ignorance is bliss” strategy occurs because the added complexity of filtering can be overwhelming, time-consuming to properly address, and seemingly distract from the main goals of the study. Nevertheless, we propose two guiding principles that investigators should adopt while filtering genomic datasets:

1) *Different filtering choices can result in different results* such that two researchers, analyzing the same dataset but using different filters, could reach entirely different conclusions.

2) *Distinctively-filtered versions of the same dataset should be used to address specific questions and to quantify the effects of filtering.* Investigatorsshould filter their data in multiple ways, report the results of such filtering on downstream analyses, and think critically to ensure that the filtered datasets used to answer specific questions are appropriate and do not themselves create a significant source of bias.

The thoughtful application of these principles should help investigators avoid biased conclusions and produce more consistent and biologically meaningful results.

To illustrate how simply changing filtering thresholds for two key filters (MAF and missing data) leads to different results, we re-filtered three pre-existing empirical datasets (Box 2). While MAF filters are often applied to remove **singletons or other rare variants**, which can reflect sequencing or genotyping errors, these rare variants (e.g., MAF < 0.05) are critical in several analyses including demographic estimation and tests for selection. Most notably, ***Tajima’s D***, a commonly used indicator of both demographic history and response to selection56, is *substantially* biased by a MAF filter choice, leading to widely differing biological inferences depending on filtering stringency (Box 2). In this case, our recommendations are straightforward: because low-frequency alleles heavily influence Tajima’s D57, researchers should apply both *no* MAF filter and a *very minor* one (such as a singleton filter to catch the most blatant sequencing errors) and compare the results when using the statistic57. The effects of additional filters (including MAF) can be substantial for diversity estimators, demographic inference58,59, *F*ST and population structure estimates60,61, estimating the distribution of locus effects on phenotypes61, and allele frequency spectra62–65. Other filtering choices therefore require similar levels of care.

With respect to principal #2 above, investigators should become more comfortable using distinctively-filtered datasets to answer different questions, even within the same study. For example, low-stringency MAF filters should be used for several demographic inferences (*e.g.*, Tajima’s D; SFS) whereas a relatively stringent MAF filter is useful for delineating populations60, planning genomically informed breeding strategies66, or for estimating individual relatedness67–69. We have created a detailed flow chart to assist investigators in matching research questions with methods and filters (Figure 3; see also Solutions and Best Practices). Pragmatically, however, this concept requires a fundamental shift in the way genetics data are analyzed: *investigators must realize that no single “best” filtering strategy or filtered dataset exists for every question, method, or objective*.

*Study-wide versus within sample-group filtering:*

Many filtering methods can be applied to all individuals in the study or separately within each sample group, which can represent different populations, geographic or temporal sampling units, or experimental treatments. When filtering occurs across all samples (*e.g.*, all individuals) within a study jointly and simultaneously, we refer to this process as ***study-wide*** (or “global”) ***filtering***. When filtering occurs within each sample group separately, we refer to this process as ***within-group filtering***. The effects of this filtering decision can be surprisingly large. For example, when a study-wide MAF filter of 0.01 was applied globally to a yellow perch (*Perca flavescens*) ***whole-genome sequencing (WGS****)* dataset, each sample group was constrained to 714,000 SNPS (Figure 2). However, when the same 0.01 MAF filter was applied separately within each sample group, the number of SNPs per sample group varied by a factor of 3.370. In this case, some populations in the study had radically different SFS,likely caused by recent population expansions in some but not all populations56,57. Study-wide filtering therefore led in this case to the removal of critically informative, globally rare but locally common alleles. While filtering MAF globally instead of within study groups is common, it should generally be expected to have serious effects whenever SFS vary between study groups, such as when demographic histories differ (like above) or when local adaptation has occurred.

Study-wide versus within-group filtering will also impact ***Genome-Wide Association Studies (GWAS)***, where it is common to perform study-wide MAF filtering where the MAF threshold is dictated by sample size (which can often be quite large, particularly in human or agricultural work71). The implications of these standardized pipelines are often not given much consideration, but the effects may be non-trivial. When populations with different SFS are compared, for example, a study-wide MAF filter can introduce ascertainment bias by removing more segregating loci from specific study groups. Human populations (and any others with complex biogeographic histories) may be prone to this bias, since populations with African ancestry tend to have more sites with low-frequency alleles than those with European ancestry72. Using a study-wide MAF filter will therefore remove more segregating loci from the African ancestry sample group and could result in the preferential detection of large effect loci in European populations. While we have focused on MAF filtering here due to its near universal implementation (see Box 1), other filtering approaches can be similarly biased by study-wide versus within-group filtering. Differences in downstream outcomes from filtering HWP53 and LD73 within-groups vs. study-wide, for example, have been previously documented.

In light of these findings, it is crucial to consider *why* results may differ when applying filters globally or within groups, particularly if sample groups include individuals from different populations, locations, or time points74. Tests for HWP should be conducted on each local group or population (deme) separately. If genetically-distinct groups are pooled, there will be an excess of homozygotes (positive *F*IS) across loci genome-wide (i.e., a ***Wahlund effect***) and their removal can mask population structure (see Figure 2)45,75. Genome-wide heterozygote deficiency can also result from low sequencing coverage and allelic dropout, which can be difficult to differentiate from Wahlund effects (although the latter influences all loci independent of read depth49,63). If a specific locus shows a consistent HWP deviation (*e.g.,* positive or negative *F*IS) in multiple different groups, it may indicate genotyping error (*e.g.*, allelic drop-out) or alignment or genome assembly errors (beyond natural/biological processes76); whereas consistent, mutilocus within-group deviations may instead indicate inbreeding, underlying cryptic population structure, and/or assortative mating.

**Solutions and best practices:**

Filtering is a powerful tool that should be applied thoughtfully, early, and often throughout genomic dataset construction and analysis to test for the effects of broad-ranging problems (*e.g.*, experimental, sample collection, labeling, or library preparation errors; batch-effect sequencing and genome assembly errors), and interesting biological phenomena (*e.g.*, natural selection, structural variants, Wahlund effects). Even in highly-studied species, such as humans, thoughtful and multi-faceted filtering is important as novel structural and genetic variants occur within every population77, and failing to account for them may curtail power to identify causal associations or even lead to incorrect inferences42.

To facilitate next-generation filtering, we have created a detailed flow chart that can be applied generally across disciplines and study systems (Figure 3; see also Table 1 and Box 4). We recognize that some alignment-free methods exist that use high throughput sequencing data, particularly for metagenomics and other phylogenomic analyses78, and they are not considered extensively here. Likewise, filtering for RNA-seq is not thoroughly reviewed here–while many concepts hold true (especially for SNPs called from RNA-seq data), many of the specific filters may not apply (see 79 and 80). For all other studies, we here highlight the salient features of a genomics study workflow.

First, we note that most genomic workflows differ depending on the question and researcher decisions, and that the documentation of filtering decisions is therefore paramount for reproducibility and research efficiency. As a first step, we recommend that raw data be immediately archived in independent, non-local repositories created for genomics data (*e.g.*, the NCBI Short-read Archive, the European Variation Archive, the DNA DataBank of Japan) prior to any analysis (see81,82 for reviews on genomics data management best practices). Given that filtering, by definition, requires manipulating data, the importance of archiving raw data cannot be understated. To this point, we refer the reader to83 for information on dataset and study organization.

After archival, reads should be filtered for general quality control (base calling quality, adapter removal, poly-G tails, sequencing artifacts, *etc.*) and trimmed when appropriate and useful84,85. For most workflows, the alignment of reads to a reference or *de novo* assembly is the next step (Figure 3). Depending on the goals of the study, it may be useful to create multiple datasets at this stage for downstream analysis. This practice is particularly relevant to *de novo* reference assembly, since assembly decisions can result in very different references and thus very different filtering and analytical outcomes. The *m* and *M* STACKS parameters and their impacts on *de novo* reference construction, for example, have been well-studied86–89.

After alignment, the data should be filtered for technical (*e.g.*, PCR) duplicates. Note that while removing PCR duplicates has been suggested to often be of little consequence90,91, this is unlikely true for every study, such as those with low coverage11,36,92 (in conflict with common practice, see Supplementary Materials). The remaining reads should then be filtered for mapping and read quality, and researchers should ensure that they record and eventually report the number of reads that passed these pre-variant filters.

Next, researchers can call variants, first filter the resulting data set to remove loci (for MAF/HWP/LD/*etc*.), and then remove poorly sequenced individuals (and/or samples with other quality or analytical concerns; see Figure 3). Note that it may be beneficial to reverse the last two steps and filter across individuals first (and loci second) in instances where retaining as many loci as possible is needed or where data quality varies widely among individuals8. An iterative approach, where individuals and loci are first removed with low stringency filters and then subjected to additional rounds of filtering stringencies may also improve data quality by removing poor individuals that reduce overall call-rates in high-quality loci and vice versa8. As with pre-variant filters, the percentage of reads, sites, and individuals retained at each post-variant filtering step should be reported (Box 4).

We generally recommend at this stage that a *minimum* of two (divergently-filtered) datasets be created - one with low filtering stringency (*e.g.*, allowing more missing data, a low MAF, and few loci removed due to HWP and LD deviations), and one with high filtering stringency (*e.g.*, many loci/individuals removed due to missing data, and a higher MAF, etc.). The precise filters and thresholds used should reflect the specificities of the study: for example, studies interested in transposable elements may want to vary alignment thresholds (uniquely versus multiply mapped reads) but keep other filters stringent to strike a balance between sensitivity and accuracy93,94. After the initial filtered datasets are created, investigators should proceed with their parameter estimation, statistical analyses, and modeling with these datasets in parallel to answer their key questions of interest. Note that we are not the first to suggest comparing outcomes from different filtering strategies13,59,63, and we suspect that this recommendation will become more common over time. Some stand-out papers exist that already follow this recommendation13,61,95,96, although they are in the minority (Box 1).

As data analysis proceeds, we suggest that re-filtering should be part of most genomics workflows. For example, PCA may reveal individuals that were mislabeled, or mis-classified into an incorrect sample group or were contaminated during sample preparation and should be removed. Similarly, the addition of new analyses (*e.g.*, Tajima’s D, transposable element annotation) that were not initially considered and may also require a careful re-filtering of the data. We suggest that authors and journals implement a supplementary table that describes the final datasets, the specific filters and thresholds employed, the name of the final VCF files, and the specific analyses for which each distinctly filtered dataset was used. We provide an example of such in Table 2.

After the analysis is completed, the resulting data and analytical tools should again be archived and/or recorded, including all relevant meta-data and the exact filtering decisions and genotyping pipelines (*e.g.*, bioinformatic scripts, software versions). Given that recreating the exact filtering and genotyping pipelines requires a considerable amount of resources (and may actually be impossible given limited data or analytical tool availability), we strongly recommend that post-project archival include all filtered genotypic/variant data in the form of carefully annotated ***VCF files*** that include detailed filtering descriptions in the header (see 97 for a detailed description of VCF files).

We recognize that a proper and thorough examination of filtering will mean extra time and work for researchers. However, these changes to current workflows are generally necessary to achieve high-quality, reproducible research and a better understanding and quantification of filtering effects. Following reproducible research guidelines may help: reproducible research is reproducible not just for other researchers, but also for the investigators themselves. A reproduction-friendly pipeline script that runs a suite of analyses when given a dataset and a set of filtering parameters is also easy to re-run a second time with a new (re-filtered) dataset. For examples of studies with well-documented methods, easily accessible data, and that would be relatively straightforward to reproduce with new filters and thresholds, see60,98,99.

**Conclusions and future directions:**

In this new and exciting era of genomics, a systematic, thoughtful, and transparent approach to filtering sequence data should be an integral part of every publication and data analysis pipeline. Investigators should strive to filter with a focus on reproducibility and aim to match the filters they employ to the questions they hope to answer. While technological advances have exponentially increased sequencing quantities, advances in filtering and its subsequent documentation or reporting have not kept pace. Instead, many investigators are unfamiliar with or ignore filtering issues during the quality control and analysis of large genomic datasets, for example, using default program settings without critical thought or explanation of their filtering decisions.

Here, we reviewed the different types of data filtering, illustrated the effects of divergent filtering choices and thresholds, and presented a flow chart and checklist to simplify, streamline, and potentially standardize and improve the filtering process. Critically, we highlight that for the same dataset 1) different filtering thresholds can create different downstream results for the same dataset, and 2) most analyses should be run on multiple datasets produced with different filters and thresholds to allow for the quantification of filtering effects on results and improve conclusion certainty. As the technology behind genomic sequencing continues to advance, we will likely see longer, higher-quality reads, which, alongside improvements in reference quality100–102 and the burgeoning field of pangenomics103,104, will increase the accuracy and power of genomic data analyses. Nonetheless, no genomic dataset will be error-free, and filtering will undoubtedly remain a central part of all genomic analyses for decades to come. We hope this review ushers a new era of next-generation filtering in genomic analyses that sparks improvements in our understanding and applications of filtering, data interpretation, reproducibility, and drives production of new data analysis tools to make it easier to re-filter and quantify filtering effects on questions across disciplines.

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# Figure Legends:

**Figure 1:** Challenges and potential solutions related to filtering that occurs prior to variant calling (“pre-variant” filtering). In the first row, individual 1 has both sufficient read depth (here illustrated with 9 reads, but a higher read depth would be better) and read alignment quality (“quality scores”) to allow for successful variant calling (a SNP is circled in orange). Despite this seemingly successful read alignment, challenges still exist; genome assembly errors (such as misplaced scaffolds on reference genomes), and structural variants (such as inversions) can cause issues downstream. Filtering for linkage disequilibrium can resolve some of these concerns. Individual 2 has a weak alignment both across an entire read and at a single base pair position across all reads, both of which should be filtered out prior to variant calling. Individual 3 has too few reads; this individual should be removed, re-sequenced, or, if this low-quality is expected and occurs across all individuals in a study, genotype likelihoods could be calculated. Individual 4 has too many reads, which can be caused by paralogs, copy number variants (CNVs), or technical (*e.g.*, PCR) duplicates. These excess reads should be filtered out and then analyzed carefully to determine the underlying causes and to facilitate answering questions of interest.

**Figure 2:** Challenges associated with four common filters that can occur after variant discovery (“post-variant” filtering). Panel A illustrates missing data, which can occur across loci and individuals. Data from monarch butterflies95 are used to show that when missing data filtering occurs within sample-groups the percentage of missing data can be high (21-100% per locus), but if filtering is performed across all samples the proportion of missing data per locus is much lower (19-56%) and quality differences between populations is obscured. In panel B, we illustrate different minor allele frequencies and use a perch study70 to show that if a MAF filter of 0.01 is applied within sample groups—but not study-wide—the number of SNPs can vary 3-fold among populations. In panel C, we illustrate how loci can be out of Hardy-Weinberg Equilibrium (HWE) due to homozygote or heterozygote excesses. Using the same monarch dataset, we show that combining two divergent sample-groups results in higher *F*IS and many more loci out of HWE. Filtering study-wide would therefore cause the erroneous removal of loci due to Wahlund effects (see74). Lastly, in panel D, we illustrate linkage disequilibrium (LD) with haplotypes that are perfectly correlated across individuals. We use a stickleback dataset105 to illustrate that LD thinning (orange points) obscures an inversion (blue points).

**Figure 3:** Flow chart to facilitate thoughtful, systematic, and reproducible filtering. Typical filtering proceeds through raw sequence QC filtering, alignment, mapped-read filtering, and variant discovery. After variant discovery, investigators must decide whether to apply filters study-wide or within sample-groups and whether to filter first by locus or individuals (see main text for recommendations). Regardless of the study objectives, multiple datasets should be constructed to examine the effects of various filtering decisions. Data should be carefully archived before filtering, and all filtering methods and results carefully reported. See Table 1 for a complete list of filters, Table 2 for a simplified example of how to report filtering results, and Box 4 for a checklist.

Glossary

ALIGNMENT

Mapping of sequencing reads and/or contigs to either each other (pairwise/multiple alignment) or to a **reference**. Alignments can vary in the strength of the evidence that supports them. Strongalignments, for example, usually have a **base (Phred) quality score** of > 20 (99% certainty); weak alignments may have a Phred score of 10 or less (<90% certainty).

BASE QUALITY SCORE

Value in a logarithmic, Phred scale given to each base on a sequencing read that indicates a quantitative degree of confidence in a nucleotide called from the sequencing instrument.

DE NOVO ASSEMBLY

**Reference** free alignment of reads into overlapping stacks or contigs for subsequent use in variant discovery and genotyping.

DISCORDANT READ-PAIRS

Paired-end reads that do not match either the expected relative directions (5’ to 3’) or the physical distance (base pairs) between reads.

FAMILY STRUCTURE

Non-independence among individuals in a study caused by direct recent shared ancestry (*e.g.*, parentage, siblings, cousins) between groups of individuals.

FILTERING

The use of quality control steps to remove errors, reads, individuals, loci, or genotypes from a dataset to improve data quality for specific analyses. This procedure differs from other forms of data processing in that it focuses on the *removal* (or “filtering out”) of data specifically rather than other forms of quality improvement such as **imputation**.

GENETIC VARIANT

A polymorphism or change in a DNA (or RNA) sequence. Includes both sequence variants (such as single-nucleotide polymorphisms) and structural variants (such as chromosomal inversions, indels, and copy-number variations). Used interchangeably with “loci” in this review.

GENOME WIDE ASSOCIATION STUDY (GWAS)

A test for statistical relationships between a phenotype (including disease) and the allelic/genotypic state of individuals across the entire set of sequenced loci. GWAS is “genome wide” in that associations are tested at many loci spread throughout the entire genome.

GENOTYPING

Calling the allelic states at a locus (*e.g.*, A/A, A/C, or C/C at a biallelic SNP in a diploid organism) or at many loci based on the underlying sequence data. Genotyping algorithms often consist of multiple steps during which filtering can occur.

HARDY-WEINBERG PROPORTIONS (HWP)

The expected frequencies of the genotypes at a given locus under Hardy-Weinberg Equilibrium (HWE). Filtering on HWP is often executed via an exact test, with loci that deviate significantly from HWP removed from subsequent analyses.

IMPUTATION

The filling in of missing data for specific genotypes and/or loci by leveraging linkage disequilibrium between missing genotypes and the called genotypes at other loci. Imputation can use reference panels of well-described haplotypes to improve performance when available, usually in well-studied model organisms.

LINKAGE DISEQUILIBRIUM (LD)

When the genotypic state of individuals in a dataset/sample group at one locus are predictive of the genotypic state of individuals at other loci. This can be caused either by physical linkage, when alleles are co-inherited due to non-independent assortment driven by physical proximity on a genome or other factors such as inversions, or gametic phase disequilibrium**,** when underlying population or family structure makes certain alleles at different loci more likely to co-occur.

MAPPING QUALITY

Score given to a read or other DNA sequence indicating the uniqueness of the alignment to a reference sequence; mapping quality algorithms vary between alignment programs.

MINOR ALLELE COUNT (MAC)

The number of gene copies or individuals carrying the minor (*i.e.* least frequent) allele at a locus.

MINOR ALLELE FREQUENCY (MAF)

The proportion (frequency) of the least common allele at a locus across a study or sample group. For example, if one diploid individual out of 50 had one copy of a unique allele (*e.g.*, an A instead of a T), the MAF would equal 0.01 In this review, we often refer to filtering out loci with MAFs below a given threshold as “MAF filtering”.

MISSING DATA

Missing genotype calls at a specific locus or individual. Missing data can be caused by many reasons, most commonly the absence of a sufficient number of reads covering a locus to call a genotype in an individual with any degree of confidence.

PARALOG

A duplicated genomic region that has arisen via either the duplication of that specific region or the entire genome (*e.g.,* genome duplication events). A type of homolog (loci identical by descent) distinct from orthologs, which arise instead due to speciation events.

PCR DUPLICATE

A technical duplicate resulting in spurious, usually identical read copies caused by repeatedly sequencing the same piece of template DNA multiple times.

POPULATION STRUCTURE

Also known as population subdivision. Non-independence among individuals in a study area/region caused by within-group biases in reproduction, usually created by spatial, temporal, or behavioral separation and often responsible for creating allele frequency differences systematically across loci.

REDUCED-REPRESENTATION SEQUENCING

The sequencing of either random or targeted subsets of a genome. Common examples include Restriction-Associated Digest (RAD) sequencing, Genotyping-By-Sequencing, and targeted sequence/exon capture.

REFERENCE

A known, often well-annotated sequence to which reads can be aligned. Can be either a published reference genome or transcriptome for a species or a ***de novo* reference**.

SAMPLE GROUPS

Groups of samples that are not independent of each other created by natural populations, geographic or temporal variation in sampling, and/or experimental treatments.

SINGLE-NUCLEOTIDE POLYMORPHISM (SNP):

A genetic variant where the allelic state of the population varies at a single base-pair.

SEQUENCING DEPTH/COVERAGE

The number of reads that were aligned to and overlap a particular genomic locus. More simply, the number of times a locus was sequenced.

SINGLETON

An allele sequenced only one time across all individuals. Sometimes alternatively defined as an allele sequenced in only one individual (which may be homozygous for that allele).

SITE-FREQUENCY SPECTRUM (SFS)

The distribution of allele frequencies across loci within a study or sample group. Can be either an “unfolded” or “polarized” derived allele frequency spectra which describe the frequency distribution of derived alleles or a “folded” or “unpolarized” minorallele frequency spectra which describes the frequency distribution of the minor alleles. Also known as the allele frequency distribution.

STRUCTURAL VARIANTS

Genetic variants which are caused by underlying variation in the order, number, and/or arrangement of loci. Copy number variations, paralogs, and indels all fall into this category. Sequence variations, where individual or series base-pairs change without changing genomic positions or counts, do not.

STUDY-WIDE FILTERING

Applying a filtering threshold globally (across the entire data set) rather than separately within each sample group.

WAHLUND EFFECT

A reduction in observed homozygosity () at many/most loci caused by underlying population structure. When multiple (sub)populations are included in a sample, any differences in allele frequency between (sub)populations will cause there to be considerably more homozygous individuals at those loci than would be expected under HWE (causing an elevated *F*IS, the reduction in heterozygosity within individuals relative expected heterozygosity). Note that this observation is what underlies the differentiation measure *F*ST when populations are split and could thus be described as *F*ST improperly calculated as *F*IS.

WITHIN-GROUP FILTERING

Applying a filtering threshold within groups separately rather than globally.

WHOLE-GENOME SEQUENCING

Sequencing the entire genome without any attempt to target specific regions (in contrast to reduced-representation sequencing). Usually entails “shotgun” sequencing and results in many more sequenced loci than reduced representation sequencing.

**Box 1: Literature Review of Current Filtering Practices**

Researchers are often confronted with a multitude of choices when filtering genomic data, not only with respect to which filter(s) to use, but which values (or thresholds) to use for each filter (Fig. 3). We conducted a literature review of 173 papers (mostly published between 2018 and 2023 years; Table S1) that analyzed genomic data (in the fields of agriculture/domestication, biodiversity studies, and human health) to quantify the reporting of filters and values chosen for each filter (see Supplemental Materials for a description on how the literature search was conducted and additional analyses).

We quantified the parameter choices of five commonly applied filters in genomic datasets: minor allele frequency (panel B), Hardy-Weinberg proportions (panel C), linkage disequilibrium (panel D), missing data per individual (panel E), and missing data per locus (panel F). Depending on the specific filter, 17-56% of studies did not mention or provide details on how or if the filter was implemented (panel A). For each of the filters we surveyed, a wide range of values were implemented among studies. Filters for MAF values ranged from 0.001 - 0.10, per-locus missing data levels varied from 0 - 90%, and Hardy-Weinberg proportion significance levels ranged from *p* < 1x10-7 to 5x10-2 (often without correction for multiple testing, see panels B-F). In several cases, authors provided details that were ambiguous or insufficient. For example, “missing data” level was often mentioned without specification of per-locus or per-individual. Oftentimes, default parameter settings were used for each filter. The high numbers of studies not reporting their filtering choices indicates that many studies would be irreproducible given the details provided in their materials and methods sections.

We suggest researchers list all filters and their specific thresholds implemented, along with a (brief) justification for the selection of that threshold because default values are often uninformed and/or inappropriate. Researchers should also explain if they corrected for multiple testing along with a (brief) justification for the correction method used (*e.g.*, Bonferroni, FDR, etc.; see Sethuraman et al. 201946).

A collage of graphs

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**Box 2: Effects of Filtering**

Genomic filtering choices can have substantial effects on downstream analyses that are not necessarily consistent across samples, populations, or statistical methods. We used three existing datasets (monarch butterflies95; cutthroat trout106; yellow perch70 to demonstrate how different MAF (panel A) or missing data filters (panel B) can affect estimates of pairwise *F*ST63–65, observed heterozygosity, the number of segregating sites (polymorphic loci; Pseg), and Tajima’s D57. We also show how a MAC filter can affect population structure estimates using the same monarch data filtered with a MAC of between 0 and 11(reproducing methods from60 , see panel C)107,108.

Our principal finding is that changing MAF and missing data thresholds can impact different datasets and different populations within datasets unequally. For example, while the monarch and perch data were not strongly affected by increased missing data stringency, outcomes from the cutthroat data changed unpredictably past a certain threshold. The impact of MAF threshold likewise differed between studies and populations: note that the number of segregating sites plummeted with even a mild MAF filter (MAF > 0.0125) in one specific population of monarch butterflies. Tajima’s D deserves particular attention: even a relatively low MAF filter (e.g., < 0.01) causes every population in every dataset to flip from an “expansion” signal (D < 0) to a “bottleneck” signal (D > 0)56,57. While not shown here, MAF filtering is also of particular note in that it almost uniformly decreases estimates of effective population size (*Ne*) derived from LD based approaches109

In contrast, population structure resolution (panel C) increased with a higher MAF filter, consistent with previous findings60,63,110. Some other methods, such as *F*ST outlier detection111, GWAS 112,113, and genomic selection66 can also see improvements with higher MAF threshold, although not without risk of bias.

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**Box 3: Filtering Trade-offs**

Different filtering choices result in trade-offs described here as false-positives and false-negatives (Type-I, α, and Type-II, β, errors, respectively; see diagram below). In the case of variant calling (*i.e.*, genotyping) we here hold that an incorrectly called genotype is the null hypothesis: while this is not the only reasonable interpretation, it simultaneously allows for a more conservative philosophical approach towards genotyping and allows for power to equal the proportion (or percentage) of correctly called genotypes that are retained in a data set. Within this framework, a false positive occurs when an incorrectly called genotype (at a single locus) is retained within a data set and a false-negative occurs when a correctly called genotype is incorrectly filtered out of the dataset (see diagram below)114. False-positives occur most frequently when filters are not strict (for example, no MAF filtering is performed) and/or when the read depth at a locus is low and thus incorrectly genotyped loci are allowed to proceed into downstream analyses. By contrast, false-negatives will be more likely when strict filters are used (such as a high MAF filter), since more loci are assumed to be errors (and thus removed) even though many of those sites may represent real, correctly called genotypes.

To answer specific questions, certain trade-offs will invariably arise. For example, when calculating Tajima’s D, many or most low frequency sites must be retained (see Box 2), however, this procedure will invariably allow more false-positives into the data set (affecting the precision of the estimator). Alternatively, when performing GWAS or outlier *F­ST* detection low frequency sites are often removed71,111, which creates data sets with few genotyping errors but may exclude real, causal variants that segregate at low frequency. Investigators should be cognizant of the very real trade-offs associated with certain filtering choices and should consider practical solutions such as creating two or more datasets with different filtering thresholds, sequencing loci of interest to higher depth/confidence, and re-sequencing select samples.

A diagram of a scientific experiment

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**Box 4: Filtering checklist**

Throughout dataset assembly (*e.g.*, from raw sequencing reads to genotypes), researchers should perform various steps to aid in reproducibility and explore the effects of alternative filtering strategies on downstream analyses. To ensure a robust examination of filtering effects and study reproducibility, the example checklist below should be consulted before and during a research project and checked-off before submitting a manuscript for peer-review.

|  |  |
| --- | --- |
|  | Data archival |
|  | Decide on filtered data sets given *a priori* study questions |
|  | Create filter recording/reporting table (see Table 2) |
|  | Filter on raw sequences (*e.g.*, read quality, poly-G tails; see Table 1) |
|  | Report exact filters used for filtering on raw sequences |
|  | Report total number of reads in study |
|  | Report total number of reads filtered out |
|  | Perform sequence alignment; report alignment parameters |
|  | Report total number of reads that aligned successfully |
|  | Report total number of reads that mapped uniquely |
|  | Report total number of reads that were filtered out |
|  | Perform filtering on successfully mapped reads |
|  | Filter on mapping quality, PCR duplicates, read pairs |
|  | Report number of reads retained/filtered at each step |
|  | Variant discovery |
|  | Begin or continue creation of multiple data sets |
|  | Decide on study-wide versus within-sample group filters |
|  | Decide on filter values to employ and order of filters |
|  | Locus filtering (see text for when individual filtering should go first) |
|  | Filter for MAF, HWP, paralogs, coverage etc. (see Table 1) |
|  | Individual filtering |
|  | Missing data; Mislabeling/contamination |
|  | Dataset construction; continue reporting of all filters and reads filtered |
|  | Data analysis and parameter estimation |
|  | Report effects of filters on parameters/questions of interest |
|  | Perform re-filtering and/or re-sequencing if necessary |
|  | Final filter recording (report reads, loci, individuals lost at each step) |
|  | Archive all filtered data sets as VCF files |

**Table 1:** Extensive list of the different types of filters available for genomic sequencing data. We briefly describe the type of filtering performed and the approximate stages of a data filtering pipeline that each filtering step would occur where: *i* = sequence QC (Quality control), *ii*= alignment to a reference, *iii* = variant discovery, and *iv* = data analysis. Note that stages *i* and *ii* constitute pre-variant filtering and stages *iii* and *iv* constitute post-variant filtering.

|  |  |  |
| --- | --- | --- |
| **Filter** | **Stage** | **Description** |
| Base quality scores | *i* | Removal of reads with many poor-quality (likely mis-read) bases. |
| Poly-G tails | *i* | Removal of spurious guanines (“G”s) that can be added to the ends of reads on certain sequencing platforms. |
| Adapter/Barcode/Cut-site trimming | *i* | Removal of adapter, barcode, or cut-site sequences from the reads. |
| Adapter/Barcode/Cut-site mismatches | *i* | Removal of reads with mismatches in the adapter, barcode, or cut-site sequences. |
| Read K-mer distribution | *i, ii* | Removal of reads with too many very common or rare runs of base-pairs (K-mers). |
| Technical/PCR duplicates | *i, ii* | Thinning of technical or PCR duplicates down to a single representative read. |
| Alignment/Mapping scores | *ii* | Removal of reads that have poor mapping scores. |
| Improperly paired reads (orientation and distance) | *ii* | Removal of paired-reads that are improperly paired (too far apart or incorrectly oriented) |
| Stack depth of coverage | *ii* | Usually for reduced-representation sequencing, removal of loci "stacks" that have too low of a sequencing depth across samples. |
| Stack mismatches | *ii* | Usually for reduced-representation sequencing, removal of loci "stacks" that have too many mismatched base-pairs across samples. |
| Number of Alleles | *ii, iii* | Removal of genotypes, haplotypes, or "stacks" with too many possible alleles. Usually for computational efficiency, but also to remove potential errors. |
| Low coverage/Quality by depth | *iii* | Removal of individual called genotypes that had poor coverage. Joint "Quality-by-depth" often alternatively used. |
| Genotype Quality/Confidence | *iii* | Removal of individually called genotypes with poor genotyping confidence. Joint "Quality-by-depth" often alternatively or additionally used. |
| High coverage | *iii* | Removal of individual called genotypes with very high coverage (usually indicating errors in the reference, paralogs, or copy-number variants, all of which require additional investigation). |
| Insertion-deletions (Indels) | *iii* | Removal of insertions or deletions (indels), often required by many down-stream applications |
| Non-biallelic loci | *iii* | Removal of non-biallelic loci (*e.g.*, monomorphic or tri-allelic SNPs); required by many down-stream applications. |
| Allow/deny listed variants | *iii* | Removal or inclusion of variants from pre-defined loci. Common for methods that target specific loci but may also sequence some additional off-target loci or where specific variants are known to be problematic. |
| Variant Read Position | *iii* | Removal of variants that tend to occur in biased positions on shotgun-sequenced reads. |
| Missing data - per individual | *iii, iv* | Removal of individuals with called genotypes at too few loci. |
| Missing data - per locus | *iii, iv* | Removal of loci with called genotypes in too few individuals. |
| Minor allele frequency | *iii, iv* | Removal of loci with low minor allele frequencies. |
| Minor allele count | *iii, iv* | Removal of loci with a low count of the minor allele across samples. |
| Hardy-Weinberg proportions | *iii, iv* | Removal of loci significantly out of Hardy-Weinberg proportions. |
| Strand Bias | *iii, iv* | Removal of loci where specific alleles are detected primarily on only the forward or reverse DNA strand. |
| Copy number variation | *iii, iv* | Removal of copy number variants. Often remain undiscovered. |
| Structural variants | *iii, iv* | Removal of structural variants, such as inversions. Often remain undiscovered. |
| Sex-linked loci | *iii, iv* | Removal of sex-linked loci, which may have biased statistical outcomes due to sex-specific sampling or behave in unexpected ways. |
| Paralogs - allelic imbalance/depth/heterozygosity | *ii, iii, iv* | Removal of reads aligned to paralagous loci, where for recent paralogs it can be unclear from which of the gene copies the read was sequenced. Additional analyses are required. |
| Mislabeling/Contamination | *iv* | Removal of individuals or loci that are likely mislabled, contaminated, or have similar issues. Can often be identified via PCA and other comparative analyses. |
| Transition-transversion bias | *iv* | Removal of loci from genomic regions with unexpected transition:transversion ratios. |
| *F*ST/selection Outliers | *iv* | Removal of outlier loci likely to be under selection. Useful for cases where putatively neutral processes specifically are of interest (*e.g.*, gene flow). |

**Table 2:** A simplified example of how investigators could report the types of datasets (from different filtering) produced during filtering, containing several examples of filtered sets with suggested standardized nomenclature/notation. In this example we assume that all filtering was performed on the same dataset. Filtering should be first guided by the types of questions investigators wish to answer, but we also strongly recommend that different filtering thresholds should be used to address the same question for comparison. For population structure, for example, investigators may want to examine the effects of several different MAF thresholds, and therefore create several different similar datasets. We show this below, although in practice several thresholds should be used for most questions, and more questions should be tested at multiple filtering thresholds. Notice that different questions suggest that different levels of filtering, types of filters, and filter thresholds should be applied. As a final note, we also suggest that filtered VCF files be retained and archived for each dataset used in the analysis, and the names of those files clearly listed in this table or in the supplementary material for reproducibility purposes.

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Questions** | **Level** | **Mapping Quality** | **Genotype Quality** | | | **MAF/MAC** | | **HWE\*** | | **LD** | **Missing Data - Individuals** | | **Missing Data - Loci** | | | ***FST* Outliers** | | **# SNPs\*\*** | **# Individuals** | | |
| Population structure | Study-wide | >20 | >14 (95% confidence) | | | > 0.05 | | p > 0.01\* | | Yes | < 20% | | < 20% | | | Yes | | 400,000 | 190 | | |
| Population structure | Study-wide | >20 | >14 (95% confidence) | | | > 0.1 | | p > 0.01\* | | Yes | < 20% | | < 20% | | | Yes | | 300,000 | 190 | | |
| GWAS | Study-wide | >30 | >20 (99% confidence) | | | > 0.05 | | p > 0.01\* | | No | < 20% | | < 20% | | | No | | 200,000 | 100 | | |
| Tajima’s D; SFS based estimators | Within Sample-Group | >20 | >14 (95% confidence) | | | MAC > 1 | | p > 1e-6 | | No | < 50% | | < 50% | | | No | | 1,500,000\*\* | 200 | | |
| Genetic diversity | Within Sample-Group | >20 | >14 (95% confidence) | | | MAC > 5 | | p >1e-4 | | Yes | < 75% | | < 75% | | | Yes | | 500,000\*\* | 170 | | |
| Selection; Selective sweeps | Within Sample-Group | >20 | >14 (95% confidence) | | | Any within group MAF > 0.05 | | p > 1e-4 | | No | < 75% | | < 75% | | | No | | 700,000\*\* | 180 | | |
|  |  |  | |  |  | |  | |  | | |  | |  |  | |  | | |  |  | |  |  |  |  |

\*Filters for loci out of HWE should only be applied at a within sample-group level (see Pearman *et al* 202253).

\*\*When filtering SNPs within sample-groups, different sample groups will contain different numbers of polymorphic SNPs. The numbers here reflect a hypothetical mean across all sample groups.