Benchmarking small-variant genotyping in polyploids

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Genotyping from sequencing is the basis of emerging strategies in the molecular breeding of polyploid plants. However, compared with the situation for diploids, where genotyping accuracies are confidently determined with comprehensive benchmarks, polyploids have been neglected; there are no benchmarks measuring genotyping error rates for small variants using real sequencing reads. We previously introduced a variant calling method – Octopus – that accurately calls germline variants in diploids and somatic mutations in tumors. Here, we evaluate Octopus and other popular tools on whole-genome tetraploid and hexaploid datasets created using *in silico* mixtures of diploid Genome In a Bottle samples. We find that genotyping errors are abundant for typical sequencing depths, but that Octopus makes 25% less errors than other methods on average. We supplement our benchmarks with concordance analysis in real autotriploid banana datasets.

Polyploidy is common in many plant species, including important agricultural crops such as wheat, potato, oat, coffee, rapeseed, cotton, banana, and sugar cane¹. In mammals, polyploidization regularly occurs during tumorigenesis, but has also been shown to be a normal part of development in some mouse and humans tissues². Molecular markers have been widely used for decades in artificial polyploid crop breeding to assist selection of more desirable traits such as better resilience to climate change and disease. More recently, genotyping by sequencing has been applied for marker assisted and genomic selection^{3–5}, and the assembly of high-quality plant reference genomes^{6–10}, together with developments in resequencing, promise new strategies for quantitative trait analysis with a wider variety of genetic variants and better linkage information than is currently possible^{10–13}.

Despite these advances, methods for genotyping polyploids from sequencing data has received little scrutiny in comparison to those for diploids $^{13-16}$. Variant calling and genotyping in polyploids is more difficult than in diploids primarily because the number of possible genotypes at a given loci is combinatorial in the ploidy and number of distinct alleles, but a sequencing read cannot distinguish identical alleles, in isolation. It therefore becomes harder to determine the copy number of a particular allele for a fixed read depth as the ploidy increases. The lower per-allele coverage also makes it harder to distinguish sequencing error from true variants. Haplotype-based methods increase power to genotype individual alleles by jointly evaluating combinations of several proximal alleles (haplotypes). They are now standard for diploid calling ^{17–22}, and are becoming more common for somatic mutation calling in tumours ¹⁷. Unfortunately, only a minority are capable of polyploid call ing^{17-19} , and none have been rigorously tested for this purpose. Specialised methods for polyploid genotyping have been developed ^{23–25}, but are only suitable for biallelic SNPs. Crucially, existing benchmarks of polyploid calling methods fall short of

the standard demanded for diploid calling 12,16,26,27 . In particular, we are not aware of any that consider indels, genotyping errors in real sequencing data, or representation differences between callers 16 . Polyploid genotyping error rates from sequencing are therefore highly uncertain, undermining developments that depends on them.

We sought to address some of these issues by conducting an in-depth assessment of polyploid small variant calling using an independent and comprehensive ground truth, real sequencing data, and haplotype-aware comparisons. Our analysis, implemented in Snakemake²⁸, is made available online at https://github.com/luntergroup/polyploid.

Results

genomes. We created synthetic Synthetic biolgylog tetraploid and hexaploid samples with high quality truth sets by merging Genome In A Bottle¹⁴ (GIAB) v4.2 GRCh38 variants for human diploid samples HG002, HG003, and HG004. We chose HG003 and HG004 for the tetraploid sample - the two unrelated parents of HG002. Evaluation regions were defined by intersecting 29 the GIAB high confidence regions for each sample, resulting in 2.50Gb (86% non-N primary reference) confident tetraploid bases containing 5,095,314 variants, and 2.49 Gb (85% non-N primary reference) confident hexaploid bases containing 5,028,566 variants. We constructed polyploid Illumina NovaSeq whole-genome test data by mixing reads generated independently for each sample with consistent library preparation and depths (Methods). Each individual sequencing run (both Illumina and PacBio) targeted 35x coverage, resulting in 70x coverage tetraploid samples and 105x coverage hexaploid samples. We confirmed total read counts were similar for each contributing sample, ensuring realistic heterozygous allele frequencies. We then randomly downsampled the full datasets, starting from 10x in 10x intervals to the full coverage, resulting in $6+2\times10=26$

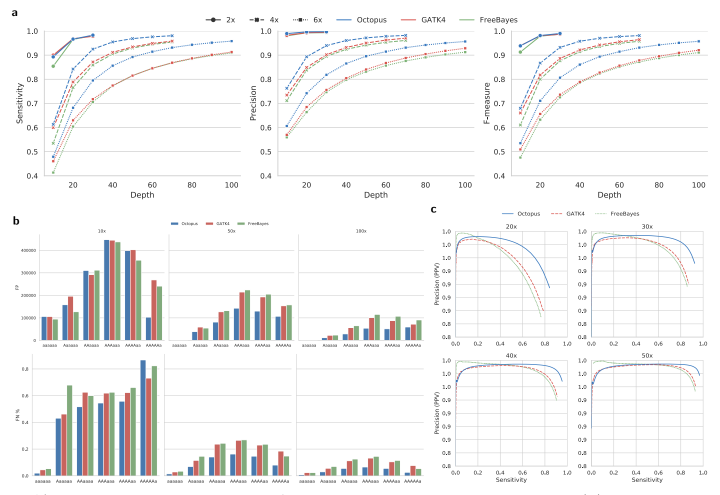


Fig. 1 | Genotyping accuracy in synthetic polyploids. a Sensitivity and precision by depth for each caller on real diploid (2x), and synthetic tetraploid (4x) and hexaploid (6x) Illumina datasets. b Counts of false positive biallelic genotypes stratified by depth and copy-number (top). Proportion of false negative biallelic genotypes stratified by depth and copy-number (bottom). c Precision-recall curves for a various tetraploid sequencing depths. Score metrics used to generate the curves were RFGQ (Octopus), GQ (GATK4), and GQ (FreeBayes).

polyploid datasets. All reads were mapped to GRCh38 with ${\rm BWA\text{-}MEM^{30}}$ (Methods).

Polyploid genotyping accuracy from short-read WGS. We evaluated three popular germline variant callers that support polyploid genotypes: Octopus¹⁷; GATK4¹⁸; and FreeBayes¹⁹, on all synthetic polyploid Illumina datasets, and in the diploid HG002 sample to get performance baselines. Other notable germline callers, such as DeepVariant²¹, Strelka2²⁰, and Platypus²², were not included because they do not support polyploid calling. We also ignored methods that call polyploid SNVs but not indels, such as polyRAD²⁴. Other than specifying the ploidy and requesting genotype qualities from FreeBayes, we used default setting for all callers (Methods). Octopus calls were random forest filtered, GATK4 and FreeBayes calls were hard-filtered using recommended thresholds (Methods). Variants were compared using RTG Tools³¹ vcfeval (Methods).

Genotyping accuracy was considerably worse for polyploids compared with diploids (Fig. 1a and Supplementary Table 1). For $30\mathrm{x}$ sequencing depth, on average 1/200 diploid genotype calls were incorrect, in contrast with 1/11 for tetraploid and 1/6 for hexaploid. Sensitivity was similarly affected; there were $8\mathrm{x}$ and $16\mathrm{x}$ more false negatives on average for tetraploid

and hexaploid, respectively, compared with diploid, for 30x sequencing. There were also more substantial differences in accuracy between callers for polyploids compared with diploids. Notably, Octopus made 25% less errors than GATK4 and FreeBayes overall, and half the errors on some depths; the largest F-measure difference between callers occurred at moderate sequencing depth: 30x for tetraploid, 50x for hexaploid. F-measure showed a typical logarithmic relationship with sequencing depth for both tetraploid and hexaploid samples, but also suboptimal response considering ploidy; the F-measure lost from doubling the ploidy was not recovered by doubling the depth, and the differential increased with depth.

The majority of false positives resulted from incorrect genotype copy-numbers: 87% of false positive biallelic genotype calls (96% of all false positives) were due to incorrect copy number. The most common false positive for all depths were the balanced heterozygotes: AAaa and AAAaaa (Fig. 1b), 92% of which were due to incorrect copy number. A larger fraction of these were made when the true genotype had -1 alternative allele copy rather than +1 copy (65% vs 34%; Supplementary Fig. 1). The most common biallelic false negatives in tetraploids were heterozygotes with a single variant copy (simplex), while for hexaploids it was heterozygotes with two

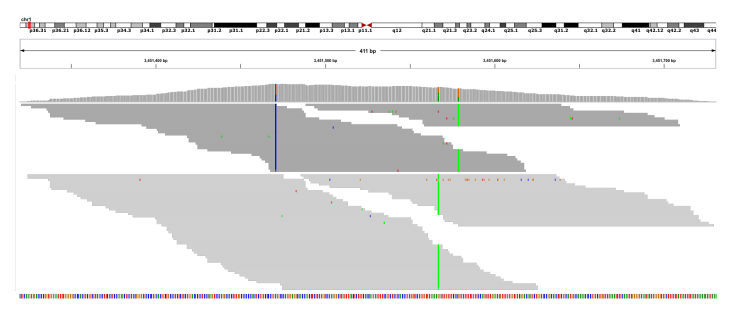


Fig. 2 | Read pileup of HG003-HG004 tetraploid colored and grouped by supported haplotype. There are two distinct haplotypes (light and dark grey). The true genotypes for the three SNVs (T>C, G>A, G>A) are AAAa, Aaaa, AAAa. The alternative allele read depths are 30/78 (38%), 38/64 (59%), and 20/58 (34%) respectively. GATK4 and FreeBayes both miscall the first two SNVs as AAaa – the most likely genotypes assuming binomially distributed allele observations. Octopus makes the correct calls because it phases all three SNVs, and the first haplotype (including the first and third SNVs) is supported by 74/114 (65%) of reads.

variant copies (Supplementary Fig. 1). However, normalising by the true prevalence shows that the most frequent false negative for depths $\geq 30 \mathrm{x}$ is the balanced heterozygote for both tetraploid and hexaploid; for depths $\leq 20 \mathrm{x}$ the most frequent false negative was simplex (Fig. 1b).

Genotype quality scores were generally well calibrated for all callers (Fig. 1c and Supplementary Fig. 2). The average F-measure percentage change for filtered verses unfiltered calls on all tests was $-0.1\%,\ -0.2\%,\ {\rm and}\ +3.5\%,\ {\rm for}\ {\rm Octopus},\ {\rm GATK4},\ {\rm and}\ {\rm FreeBayes},\ {\rm respectively}.\ {\rm Notably},\ {\rm filtering}\ {\rm did}\ {\rm not}\ {\rm always}\ {\rm improve}\ {\rm F\text{-}measure};\ {\rm only}\ {\rm Octopus}\ {\rm improved}\ {\rm F\text{-}measure}\ {\rm on}\ {\rm all}\ {\rm tests}\ {\rm with}\ {\rm filtering},\ {\rm highlighting}\ {\rm the}\ {\rm advantage}\ {\rm of}\ {\rm machine-learning}\ {\rm based}\ {\rm filters}\ {\rm compared}\ {\rm to}\ {\rm hard}\ {\rm filters}.\ {\rm However},\ {\rm performance}\ {\rm differentials}\ {\rm between}\ {\rm callers}\ {\rm were}\ {\rm similar}\ {\rm for}\ {\rm unfiltered}\ {\rm calls}\ ({\rm Supplementary}\ {\rm Table}\ 1),\ {\rm showing}\ {\rm that}\ {\rm most}\ {\rm of}\ {\rm Octopus}\ {\rm 's}\ {\rm performance}\ {\rm advantage}\ {\rm come}\ {\rm from}\ {\rm better}\ {\rm genotyping}\ {\rm rather}\ {\rm than}\ {\rm filtering}.$

Longer haplotypes improve genotyping accuracy. A plausible explanation for Octopus having better genotyping accuracy than GATK4 and FreeBayes is that Octopus considered longer haplotypes — on average — when calculating genotype likelihoods. If the true set of haplotypes including a subset of variants can be confidently determined, then the variance in the genotype posterior probability distribution is expected to decrease, in particularly with respect to copy number, with larger subsets (and therefore longer haplotypes) since the number of discriminating reads is expected to be proportional to the haplotype length (Fig. 2). To test this, we recalled genotypes in the 30x tetraploid sample using a parametrisation of Octopus designed to generate longer haplotypes than with default settings (Methods). The mean called haplotype length increased from x bases to y bases and the F-measure increased

by x% compared with default settings.

Banana genotyping. Dwarf Cavendish banana (*Musa acuminata*) is autotriploid consisting of 11 chromosomes with a haploid genome size of around $523 \mathrm{Mb}$, and is an important food source and export-product for many developing countries 7 . To support our previous results on real polyploid samples, we called variants (Methods) in a dwarf Cavendish banana specimen that was previously whole-genome sequenced with two Illumina technologies, NextSeq-500 and HiSeq-1500, to $65 \mathrm{x}$ and $55 \mathrm{x}$ coverage, respectively 32 . Both datasets were mapped to the DH Pahang v2 reference 7 , and genotypes were called with Octopus, GATK4, and FreeBayes.

Due to lack of truth data, we evaluated concordance on the two banana datasets using haplotype-aware intersections (Methods). Genotypes called by all callers in both datasets, while substantially the largest intersection set, only accounted for 40% of all distinct genotype calls; 20% of calls were unique to a single callset (Fig. 3). However, there were considerable differences in concordance between the two datasets for each caller: GATK4 had 37% more discordant calls compared with Octopus and 16% more than FreeBayes, despite making 0.5%less calls overall than FreeBayes and only 3% more than Octopus (Table 1). We also found high disconcordance when intersecting by called alleles (Supplementary Fig. 3 and Supplementary Table 2); 58% of distinct alleles were present in all callsets while 11% were unique to a single callset, indicating that that, in comparison to our results on synthetic data, a larger proportion of false calls arise from incorrect variant alleles rather than copy-number errors.

Manual review of discordant calls indicated that a large fraction were due to slightly different indels called in each dataset, suggesting failure to discover correct allele(s). To test this,

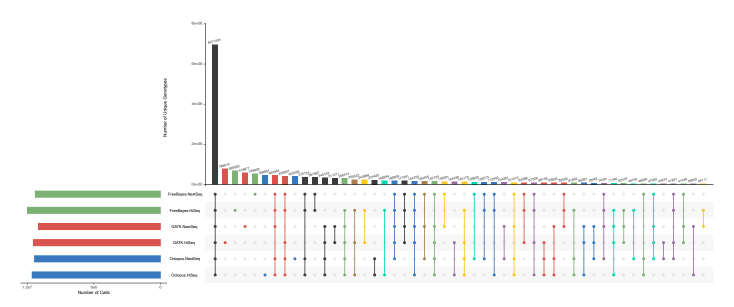


Fig. 3 | Comparison of genotypes called in two Illumina datasets (HiSeq and NextSeq) of banana specimen by Octopus, GATK4, and FreeBayes. 'UpSet' plot shows callset intersections for each caller-dataset pair. The largest 50/63 intersection sets are shown. Intersections are color coded by caller discordance between the two datasets: No discordances (black), Octopus (blue), GATK4 (red), FreeBayes (green), Octopus & GATK4 (purple), Octopus & FreeBayes (cyan), GATK4 & FreeBayes (yellow), All (brown). The total number of unique genotype calls was 17,277,217.

we recalled both datasets with Octopus using the union of the four callsets from Octopus and GATK4 as candidates (Online Methods). The number of called variants increased by x% and the fraction of concordant calls increased to x%, supporting this hypothesis. Further assessment of a selection of remaining discordant calls using haplotype-tagged and realigned evidence BAMs generated by Octopus (Methods) indicated that major sources of error were: i) lack of read depth or allele bias; ii) mismapped reads, possibly due to incomplete or divergent reference; iii) failure to discover a correct allele (in any callset); iv) probable structural variation.

Discussion

We have shown that genotyping is substantially more difficult in polyploids than in diploids using typical whole-genome sequencing depths, and that there are considerable differences in accuracy between callers. Notably, Octopus produced less than half the total errors of other methods. We believe this is due to Octopus modelling longer haplotypes during genotyping, as this increases statistical confidence in allele copy-number. Regenotyping using longer haplotypes increased genotyping accuracy, supporting this hypothesis.

Analysis of real autotriploid banana datasets revealed high discordance between callers, and more alarmingly, high discordance for callers on similar datasets of an identical specimen.

Table 1 | Concordance in two banana Illumina datasets Caller Concordant Discordant Total Concordant % FreeBayes 9.778.611 3,729,699 13,508,310 72% GATK4 9.025.920 4,421,440 13,447,360 67% Octopus 9,854,737 3,219,611 13,074,348

While these results were, at least, consistent with the relative accuracy of callers determined by our benchmarks using synthetic polyploid data (Octopus was the most concordant caller), absolute error rates were clearly higher in real polyploid data. Plausible reasons for this include: i) greater divergence from the reference genome; ii) higher levels of repetitive elements in the genome; iii) more structural variation; iv) less complete reference genome; v) higher rates of sequencing related errors, such as due to the use of PCR amplification. vi) callers optimised for human data.

We have only considered single sample polyploid calling in this work, however, multi-sample calling is important for studying population diversity. Population calling in humans is a difficult problem due to the computational complexities of joint calling and difficulties in merging independent callsets. Population calling in polyploids will likely be even more challenging, and would perhaps benefit from more sophisticated genotype priors as developed in other methods ²³.

Moving forward, there is clearly room for improvement in polyploid genotyping from sequencing. The creation of high quality validation sets with real polyploid samples would highly valuable in the development of polyploid calling algorithms, including Octopus. We hope that this work lays the groundwork for future developments.

Acknowledgements. This work was supported by The Wellcome Trust Genomic Medicine and Statistics PhD Program (grant nos. 203735/Z/16/Z to D.P.C). The computational aspects of this research were supported by the Wellcome Trust Core Award Grant Number 203141/Z/16/Z and the NIHR Oxford BRC. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health. We would like to thank Len Trigg at Real Time Genomics for kindly arranging an update to RTG Tools

Time Genomics for kindly arranging an update to RTG Tools for polyploid genotype support on our request.

Author contributions. D.P.C did the analysis and wrote the paper. D.W. and G.L critically reviewed the manuscript and supervised the project.

Ethics declaration. The authors declare no competing interests.

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Methods

Synthetic polyploids with real reads. Raw reads (FASTQ) generated for the PrecisionFDA Truth v2 challenge 33 were downloaded from the DNAnexus portal (https://precision.fda.gov/challenges/10). Each FASTQ was line counted to ensure realistic haplotype frequencies, before concatenating contributing samples to make the full data polyploid dataset. Downsampling was performed directly on the FASTQ files using seqtk with default seed. The sampling fraction was set using: $test\ depth$ / $full\ depth$, where full depth is $35 \times ploidy/2$. Illumina reads were mapped with BWA-mem using default alignment parameters.

Changes to Octopus for polyploid calling. While the models that we previously described for Octopus ¹⁷ are fully capable of polyploid calls, in practice we found some issues. Runtimes were prohibiting for high ploidies due to the model always considering every possible genotype for a given set of candidate haplotypes, which is reasonable for diploids but not polyploids. Moreover, sensitivity for low copynumber variants was less than ideal due to the variant discovery mechanisms not fully accounting for ploidy.

To resolve the runtime issue, we modified the genotype proposal algorithm so that an upper bound on the number of genotypes evaluated can be specified. The algorithm respects this limit by evaluating the full model on the maximum ploidy that results in less candidate genotypes than the limit for a given set of haplotypes, and then extends a subset of these with greatest posterior probability using each of the candidate haplotypes. The procedure is then applied iteratively, increasing the ploidy by one each iteration, until the desired ploidy is reached. We expect this procedure to work well when the number of unique haplotypes present in a region is not substantially greater than the first ploidy considered. We addressed the sensitivity issue by tweaking the pileup and local *de novo* reassembly candidate variant discovery algorithms to account for sample ploidy.

Variant calling synthetic polyploids. For GATK4, we called variants using BAMs with marked duplicates created by GATK4's *MarkDuplicates* tool. Raw BAMs were used for FreeBayes and Octopus. The sample ploidy was specified for all callers: --organism-ploidy (Octopus), --sample-ploidy (GATK4), and --ploidy (FreeBayes). For FreeBayes, we requested genotype qualities with the -= optiion.

Filtering variant calls. We used HiSeq-2500 data from GIAB for samples HG001, HG006, and HG007 to train Octopus's random forest for polyploids, using the same mixing procedure described or the evaluations. The forest was also trained on HG001. This forest was used to filter all polyploid calls, including the banana datasets. The v0.7.0 germline forest was used to filter diploid calls. For GATK4, we used filter expressions: "-filter 'QD < 2.0' -filter-name 'QD2' -filter 'QUAL < 50' -filter-name 'Q50' -filter 'GQ < 5' -filtername 'GQ5' -filter 'FS > 60.0' -filter-name 'FS60' -filter 'SOR > 3.0' -filter-name 'SOR3' -filter 'MQ < 40.0' -filter-name 'MQ40' -filter 'MQRankSum < -12.5' -filter-name 'MQRankSum-12.5' -filter 'ReadPosRankSum < -8.0' -filter-name 'ReadPosRankSum-8'". For FreeBayes, we used filter expression: "QUAL > 1 & GQ > 1 & SAF > 0 & SAR > 0".

Identifying copy-number errors. Biallelic sites were identified after intersecting false negative and false positive genotypes previously identified by RTG Tools vcfeval. The intersection was also performed using RTG Tools vcfeval with the *--squash-ploidy* and *--output-mode=annotate* options.

Long haplotypes with Octopus.

Variant calling banana. We tweaked caller parameters for banana calling to account for the high diversity present in banana species. For GATK4, we set *--heterozygosity* to 0.05, *--heterozygosity-stdev* to 0.05, and *--indel-heterozygosity* to 0.01. For FreeBayes, we set *--theta* to 0.05. For Octopus, we set emph--snp-heterozygosity to 0.05, *--snp-heterozygosity-stdev* to 0.05, and *--indel-heterozygosity* to 0.01, *--max-haplotypes* to 300, and *--max-genotypes* to 30000.

Banana concordance analysis. Callsets for the banana datasets were intersected using a custom script (https://github.com/dancooke/starfish) that invokes both RTG Tools vcfeval (that only supports 2-way comparisons) and bcftools to achieve multisample haplotype-aware comparisons.

Code availability. Octopus source code and documentation is freely available under the MIT licence from https://github.com/luntergroup/octopus. Custom Snakemake and Python code used for data analysis is available from https://github.com/luntergroup/polyploid.

Data availability. All primary data used for analysis was previously published in the Precision FDA Truth Challenge v2³³. Synthetic datasets can easily be reproduced using the online code.