Expanding the Conservation Genomics Toolbox: Incorporating Structural Variants to Enhance Genomic Studies for Species of Conservation Concern

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Publication Information

Wold, Jana; Koepfli, Klaus-Peter; Galla, Stephanie J.; Eccles, David; Hogg, Carolyn J.; Le Lec, Marissa F.; . . . and Steeves, Tammy E. (2021). "Expanding the Conservation Genomics Toolbox: Incorporating Structural Variants to Enhance Genomic Studies for Species of Conservation Concern". Molecular Ecology, 30(23), 5949-5965. https://doi.org/10.1111/mec.16141

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Expanding the conservation genomics toolbox: Incorporating structural variants to enhance genomic studies for species of conservation concern

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Abstract
Structural variants (SVs) are large rearrangements (>50 bp) within the genome that impact gene function and the content and structure of chromosomes. As a result, SVs are a significant source of functional genomic variation, that is, variation at genomic regions underpinning phenotype differences, that can have large effects on individual and population fitness. While there are increasing opportunities to investigate functional genomic variation in threatened species via single nucleotide polymorphism (SNP) data sets, SVs remain understudied despite their potential influence on fitness traits of conservation interest. In this future-focused Opinion, we contend that characterizing SVs offers the conservation genomics community an exciting opportunity to complement SNP-based approaches to enhance species recovery. We also leverage the existing literature—predominantly in human health, agriculture and ecoevolutionary biology—to identify approaches for readily characterizing SVs and consider how integrating these into the conservation genomics toolbox may transform the way we manage some of the world’s most threatened species.

KEYWORDS
conservation genetics, fitness traits, functional diversity, genome-wide diversity, pangenomes, structural variation
Over the past 10 years, the field of conservation genomics has largely focused on transitioning from a handful of microsatellite markers to thousands of genome-wide single nucleotide polymorphisms (SNPs) (Allendorf et al., 2010; Mable, 2019). Whether characterized using reduced-representation data or whole-genome sequence (WGS) methods, SNP-based estimates of genome-wide variation are being used to better inform conservation by delineating species, including cryptic species (e.g., Binks et al., 2021; Quattrini et al., 2019; but see Stanton et al., 2019), detecting hybridization and introgression (DuFresnes & Dubey, 2020; Peters et al., 2016; but see Forsdick et al., 2021 preprint; Hauser et al., 2021), identifying conservation units (Liddell et al., 2020), informing conservation translocations (Dresser et al., 2017; Glassock et al., 2021), guiding conservation breeding programmes (e.g., Gala et al.,; Wright et al., 2020) and identifying the genomic basis of adaptive traits (Dunt sch et al., 2020). However, SNPs constitute only one component of genome-wide variation (Ho et al., 2020). Structural variants (SVs) are genomic rearrangements such as insertions, deletions, duplications, inversions, and translocations, and are generally defined as ≥50 bp in length (Figure 1). Variation in chromosome structure was one of the earliest types of genetic variation studied (e.g., McClung, 1905; Sturtevant, 1921), and a recent focus on the impacts of these genomic rearrangements on sequence variation has found that SVs affect more overall genome content and intersect with genes more often than SNPs (Catanach et al., 2019; Chakraborty et al., 2019; Chiang et al., 2017; Frayling, 2014; Pang et al., 2010).

Structural variants can have significant evolutionary impacts on species. These impacts include genomic incompatibilities between ancestral and novel variants that may influence hybridization and introgression (Weissensteiner et al., 2020) and may even lead to speciation (Davey et al., 2016; Todesco et al., 2020). In addition, emerging evidence indicates that SVs can underlay fine-scale population structure, and may facilitate local adaptation (Cayuela et al., 2021; Dorant et al., 2020; Tigano et al., 2021). These findings—which are generally the result of integrating SNPs and SVs—highlight the opportunity for both variant types to inform conservation. Nevertheless, the broad application of SVs in conservation remains challenging in part due to the uncertainty in how best to detect and genotype all SV types at the population scale (Mérot et al., 2020). Further, although SVs are more likely to impact fitness traits than SNPs (Pang et al., 2010), the overall impacts of SVs on population persistence is relatively unexplored (Oomen et al., 2020).

Here, we leverage our cross-sector expertise in conservation (AWS, CJH, JG, JRW, KPK, MLL, SJG, TES), human health (DE), agriculture (JG), quantitative genetics (AWS) and bioinformatics (DE, JG, JRW, MLL) to consider the significance of SVs in population persistence and species extinction risk and discuss opportunities and challenges for relating SVs to fitness traits in small populations. We identify approaches for reliably characterizing SVs with an emphasis on leveraging new and existing genomic resources for species of conservation concern. We also describe how the incorporation of genome assemblies for multiple reference individuals can be used to improve SV discovery and genotyping, even for large and/or complex SVs (Alonge et al., 2020; Eizenga et al., 2020; Gao et al., 2019; Golicz et al., 2016; Tettelin et al., 2005). We contend that the study of SVs has the potential to improve conservation outcomes by enhancing our understanding of genome-wide variation and traits related to fitness.
be purged from the population, small populations may selectively accumulate slightly deleterious SVs. This enrichment indicates that SVs probably underpin some negative fitness traits associated with inbreeding depression. Although there are some clues as to the evolutionary dynamics of SVs in small populations, further study is critical for resolving the significance of SVs in population persistence and species extinction risk.

3 | RELATING STRUCTURAL VARIANTS TO FITNESS TRAITS

Advancements in WGS technologies coupled with increased computational capacities, have renewed interest in the role of SVs in determining trait differences (Chiang et al., 2017; Pang et al., 2010; Sadowski et al., 2019; Yi & Ju, 2018). For example, characterizing SVs in agricultural species has led to the identification of variants associated with economically significant traits in crops like grapevine (Zhou et al., 2019), maize (Yang et al., 2019), soybean (Liu et al., 2020) and tomato (Alonge et al., 2020), and identified specific genes and gene regions associated with domestication in vertebrates (Bertolotti et al., 2020; Cagan & Blass, 2016; vonHoldt et al., 2017).

The impact of these SVs on trait variation may be direct or indirect. For example, SVs can lead to direct changes to gene coding regions and gene regulation that change the function or expression of proteins (Bickhart & Liu, 2014; Collins et al., 2020). In contrast, in some cases where a particular SV is associated with a trait, the SV may not be the causal variant, but rather increase the likelihood of de novo causal mutations nearby. This is the case for a relatively common 1.3 Mb inversion on the human Y-chromosome where microdeletions accrue at inversion breakpoints, which can result in profound impacts on male fertility (Hallast et al., 2021). There is also growing evidence that SVs can further impact the gene regulatory landscape by altering the formation of topologically associating domains (TADs), genomic regions that physically interact with themselves more frequently than with regions elsewhere (Sadowski et al., 2019; Shanta et al., 2020).

There is growing evidence that SVs—in particular, the suppression of recombination and subsequent evolution of “supergenes”, or tightly linked coadapted alleles—can impact fitness traits in natural
populations (Huynh et al., 2011; Jay et al., 2018; Kim et al., 2017). For example, sperm swimming speed in zebra finch is determined by inversion haplotypes, with heterokaryotypic males producing faster sperm than homokaryotypic males (Kim et al., 2017; Knief et al., 2017). In addition, a supergene resulting from an inversion has been found to determine mating strategy and morphology in the Eurasian ruff (Küpper et al., 2016). Moreover, it is notable that this inversion is probably a lethal recessive variant (Lamichhaney et al., 2016). Similarly, a large inversion resulting in a supergene underlies significant morphological and behavioural differences among white-striped and tan-striped morphs in White-throated sparrows, with aggressiveness being monogenic in the white-striped morph (Merritt et al., 2020). Although many fitness traits are likely to be polygenic, including traits of conservation interest such as disease susceptibility, reduced fertility and developmental abnormalities (e.g., Moran et al., 2021; Murchison et al., 2012; Roelke et al., 1993; Savage et al., 2020 preprint), those impacted by supergenes are likely to have relatively simple inheritance patterns which will enable their characterization and management.

To date, there are generally two approaches to investigate the genomic basis of traits, both of which require clearly defined traits and well-curated data sets and high-quality, well annotated, reference genomes: (1) Comparative genomics, where well-characterized groups (individuals, populations or species) are used to identify highly differentiated genomic regions (Alonge et al., 2020; McHale et al., 2012; vonHoldt et al., 2017; Weissensteiner et al., 2020); and (2) Association studies, where associations between specific markers (i.e., SNPs and/or SVs) and traits are assessed (Chakraborty et al., 2019). For example, association studies and/or comparative approaches using SVs have revealed signatures of domestication (e.g., aquaculture salmon and dogs; Bertolotti et al., 2020; Cagan & Blass, 2016; vonHoldt et al., 2017), and the relative contribution of SVs to species evolution (e.g., Atlantic cod, Corvids, Heliconius butterflies, Sunflowers; Berg et al., 2017; Joron et al., 2006, 2011; Rieseberg et al., 1999; Weissensteiner et al., 2020).

Whole-genome SNP-based association studies have been extensively applied to model organisms, agriculturally significant species and humans, although they are increasingly applied to wild and non-model species (Santure & Garant, 2018). However, for many SNP-based association studies, significantly associated loci do not explain the vast majority of known trait heritability (i.e., the proportion of trait variation that is due to genetic rather than environmental differences between individuals; Clarke & Cooper, 2010; Eichler et al., 2010; Manolio et al., 2009). This "missing heritability" is often attributed to genetic variation not explained by genotyped loci, that is causal variants may remain uncharacterized (Manolio et al., 2009). Another hypothesis to explain this missing heritability is that SVs are a significant source of trait variation (Eichler et al., 2010; Frazer et al., 2009). Further, although a given SV is likely to be linked to a nearby SNP (Wildt et al., 2020), they may not be in strong linkage disequilibrium with each other, so that the SNP will not capture the impact of the SV on the trait (Pang et al., 2010). Structural variants may also lead to challenges in aligning reads and calling SNPs in a region, in turn preventing the identification of SVs from sequence data (Pang et al., 2010). The inclusion of SVs into association studies is therefore likely to lead to better power to detect causative variants, particularly given advances in GWAS methods, including machine-learning approaches, that can more effectively prioritize variants in a region and account for nonadditive interactions (Brieuc et al., 2015; Ramzan et al., 2020). This is particularly promising as evidence suggests that SVs have larger phenotypic effects and may be deleterious more often than SNPs (Chakraborty et al., 2019; Conrad et al., 2010; Cridland et al., 2013; Emerson et al., 2008; Rogers et al., 2015). For example, a quantitative trait locus (QTL) study for Drosophila melanogaster found that about half of all candidate genes underpinning mapped QTL were impacted by SVs, and that a large proportion of genes contained multiple rare SVs (Chakraborty et al., 2019).

## 4 | STRUCTURAL VARIANT DISCOVERY AND GENOTYPING IN THREATENED SPECIES

### 4.1 | A high-quality reference genome is an invaluable investment

While linkage mapping and cytological techniques can be used to characterize SVs, here we focus on the approaches most likely to be accessible to the conservation genetics community (Deakin et al., 2019). Partnerships with global genome consortia (e.g., Vertebrate Genomes Project, Rhie et al., 2021; Earth BioGenome Project, Lewin et al., 2018; Nature and Zoonomia Consortium, 2020) are providing increased accessibility to highly contiguous, well-annotated genome assemblies generated from both short and long read data and long-range scaffolding approaches for many species, including those of conservation concern and their close relatives (Whibley et al., 2020). As costs of generating and analysing WGS data continue to drop, a growing number of conservation genomicists working on species beyond global genome consortia are investing in the assembly and annotation of high-quality reference genomes. We readily recognize that not all conservation programs are able to access the resources or sample quality to generate a high-quality, well annotated reference genome, nor do we recommend this action for all threatened species. Where there is a clear hypothesis that SVs may be impacting a particular gene or region, comparative approaches (Alonge et al., 2020; Weissensteiner et al., 2020), and target capture methods (vonHoldt et al., 2017) may be the most cost-effective, especially if a reference genome from a closely related species can be leveraged instead. However, as described below, investment in a high-quality reference genome is a near necessity for accurate and precise SV discovery and genotyping, particularly if the goal is to characterize a broad range of SV types for downstream functional analyses (e.g., investigating the genomic basis of traits linked to fitness). Further, if the goal is to characterize a large proportion of SVs—including large and/or complex SVs—across the genome, integration of multiple
reference genomes provide a powerful tool for accurately characterizing genome-wide variation and may facilitate genotyping across multiple variant classes (i.e., both SNPs and SVs; Figure 2).

### 4.2 Alignment-based SV discovery methods

Once a high-quality reference genome has been assembled for a species of interest, SVs can be detected using alignment-based methods where short- and/or long read sequence data for multiple individuals are aligned to the reference without prior assembly. These methods rely on identifying patterns in read mapping between a sample and the reference genome. Common algorithms include read-pair, read depth, split-reads and de novo or local assembly (Hajirasouliha et al., 2010; Korbel et al., 2007; Yoon et al., 2009). No single algorithm is well suited to detecting all SV types. For example, read pair-based algorithms, where the orientation and distance between paired ends are assessed, are suitable for detecting deletions, duplications, and inversions. In the case of read depth approaches, deletions and duplications are identified through variation in mapping depth. Split-reads identify regions where alignments map across a breakpoint, which is best suited for discerning the ends of an inversion or translocation (Figure 3). Finally, de novo and local assembly algorithms are best suited to identifying variants not present in the reference (e.g., insertions) as reads aligned to contigs may be reassembled alongside unmapped reads for pairwise comparison to a reference (Mahmoud et al., 2019). Early implementation of SV discovery programs typically relied on a single approach (e.g., BreakDancer, Pindel; Chen et al., 2009; Ye et al., 2009). Although these programs are less computationally intensive, they are limited to calling only a few SV types and tend to underperform against generalist programs that incorporate at least three algorithms for SV detection (e.g., Delly, Lumpy, Manta, SvABA; Chen et al., 2016; Layer et al., 2014; Rausch et al., 2012; Wala et al., 2018).

Because of the challenges of identifying and classifying SVs, many SV discovery approaches have a high (and systematic) false positive rate (e.g., Cameron et al., 2019), especially when data is derived from short-read sequencing methods (Figure 3; see more below). In order to address this error, an ensemble approach for SV characterization has been extensively applied in human data and in domesticates (Du et al., 2021; Ho et al., 2020; Zhou et al., 2019). With an ensemble approach, multiple SV callers are integrated into a single pipeline as a means to create more certainty around SV discovery, with only variants that intersect multiple SV callers retained (Becker et al., 2018; Mohiyuddin et al., 2015; Zarate et al., 2018 pre-print). Further, because validated SV call sets have been developed for humans, it is straightforward to benchmark appropriate program combinations for SV discovery (Collins et al., 2020; Ho et al., 2020; Parikh et al., 2016; Zook et al., 2020).

When validated SV call sets are lacking, we recommend using a generalist SV discovery program that combines multiple SV discovery algorithms to target a range of SVs at once. These generalist programs enable a combination of assembly-based methods alongside read depth, read pairs and/or split-read approaches (i.e., GRIDSS, Cameron et al., 2017; Manta, Chen et al., 2016). These programs not only perform well on their own but have the added benefit of performing well over a range of SV types (Cameron et al., 2019; Kosugi et al., 2019). Using a generalist program has an additional benefit that it overcomes the challenge faced by many ensemble methods, where distinguishing between true variants and false positives is difficult.
due to the significant overlap in false positives across methods (Cameron et al., 2019). However, conservation geneticists should be aware that the computational resources required to characterize SVs at the population scale using generalist SV discovery programs are substantial (e.g., for a diploid 1.15 Gb genome with 170 individuals sequenced to ≥25× coverage, 72 physical cores, 460 Gb RAM, >3 Tb storage was required to implement paired-, split-read and assembly-based SV discovery algorithms; JRW personal observation).

4.3 Multiple reference genomes improve genome-wide structural variant discovery and genotyping

A pan-genome is the aggregate characterization of genomic variation present in a group of interest, including species and populations (e.g., variation between strains of tomato, Alonge et al., 2020). Pan-genomes offer a straightforward solution to address the challenges associated with SV discovery and genotyping with short-read sequence data.
Although originally developed to characterize variation in bacteria (Tettelin et al., 2005), they are commonly used in studies of trait diversity in humans (Pang et al., 2010) and agriculturally significant species (e.g., cattle, goats, soybean, and maize; Bickhart et al., 2020; Della Coletta et al., 2021; Golicz et al., 2016; Liu et al., 2020; Low et al., 2020; McHale et al., 2012; Yang et al., 2019). There are two components of a pangenome: the "core" genomic regions that do not vary among individuals, and "accessory" genomic regions that vary among individuals (Bayer et al., 2020; Golicz et al., 2016; Hurgobin & Edwards, 2017; Figure 4). In a pangenomic approach, genomes of multiple individuals are assembled de novo using multiple platforms (e.g., long reads, Hi-C, Optical mapping; Song et al., 2020; Soto et al., 2020; Weissensteiner et al., 2020; Zhou et al., 2019), followed by pairwise comparisons of whole-genome alignments for SNP and SV discovery (e.g., Cortex, MUMmer, Minimap2; Delcher et al., 1999; Iqbal et al., 2012; Li, 2018). Once variant discovery is complete, genome graphs representing the variation in the pangenome may be constructed to efficiently represent "core" and "accessory" regions (Eizenga et al., 2020; Li, 2018; Rakocevic et al., 2019; Tettelin et al., 2005) that are encompassed in analyses of copy number variation (CNV) and presence-absence variation (PAV) (Della Coletta et al., 2021). Genome graphs are a powerful method for population-level genotyping and consistently outperform alignment-based genotyping (e.g., Ebler et al., 2020 preprint; Eggertsson, 2017; Iqbal et al., 2012; Kim et al., 2019; Li, 2018). As a result, pangenomic approaches can resolve complex variants (i.e., multiple overlapping events) that may otherwise go undetected in alignment-based approaches, hampering the discovery of causal variants (Alonge et al., 2020; McHale et al., 2012).

For species of conservation concern, individuals selected for reference genomes may not be the most representative in that some are chosen to showcase individuals known to the public (e.g., animal ambassadors) or are the result of opportunistic sampling. Regardless, in most cases, it is unlikely that a single individual accurately represents the genomic variation within the species as a whole, even in highly inbred species (Gao et al., 2019; McHale et al., 2012). The primary advantage of a pangenomic approach is that multiple alternative alleles are assessed while read mapping, effectively removing reference bias (Paten et al., 2017). In other words, sequence reads are not precluded from mapping if they are not represented in the primary reference (e.g., insertions, Figure 3). The promise of pangenomes is exemplified in crop species such as soy, wheat and Brassica spp. where many more SVs are resolved, leading to a paradigm shift in crop improvement (Della Coletta et al., 2021; Golicz et al., 2016; Liu et al., 2020; Montenegro et al., 2017; Song et al., 2020). However, the ability of conservation programs to establish pangenomes largely depends on the availability of existing genomic resources and the ability to secure substantive funding.

**Figure 4** Characterization of SVs using assembled whole-genomes and genome graphs using multiple reference genomes: (a) Schematic of structural variant (SV) discovery using alignment of assembled whole genomes. (b) Schematic of genome graphs, which can then be used to characterize a pangenome and facilitate rapid genotyping of individuals, as "core" regions (genomic regions that do not vary among individuals) are readily distinguished from "accessory" regions (genomic regions that do vary among individuals). Genome graphs can also facilitate accurate genotyping as more than one allele may be considered at once. The use of multiple reference genomes can reduce reference bias and are better able to capture insertions, inversion haplotypes and complex SVs (i.e., genomic regions where multiple SVs occur in close proximity) [Colour figure can be viewed at wileyonlinelibrary.com]
4.4 | SV discovery and genotyping with short-read sequence data

Moderate coverage (e.g., ~10x) short-read sequencing is becoming the predominant approach in conservation genomics (Cam et al., 2020; Galla et al., 2018; Lado et al., 2020; Lew et al., 2015; Lujan et al., 2020; Oyler-McCance et al., 2015; Robinson et al., 2016). Moderate coverage data sets are generally cost effective and appropriate for characterizing SNPs using a single reference genome, whereas a minimum of 30x coverage is often recommended for de novo SV discovery and genotyping (Ahn et al., 2009; Kosugi et al., 2019; Sims et al., 2014; Wang et al., 2008). This is because at low coverage it is challenging to determine whether a detected variant is an artifact of sequencing/mapping error or a “true” variant (Figure 3). For example, mapping errors may occur when the relative size of a SV spans a large portion or the entire length of a read, or in the case of a complex rearrangement, prevents mapping altogether (Sedlazeck et al., 2018; Yi & Ju, 2018). In addition, read insert size (400–600 bp) may bias the discovery of insertions and deletions in some pipelines (Kosugi et al., 2019). By increasing average read depth to 30x, the likelihood of distinguishing between sequencing error, read mapping errors and true genomic variation increases as well. However, due to the increased costs, such high coverage short-read data sets may be challenging to generate for many species of conservation concern. The question then becomes how best to fully utilize moderate coverage short-read data sets to investigate SVs. Recent studies demonstrate that an average read depth of 10x is sufficient for population-scale comparisons, provided sampling is representative and a high-quality reference genome is available (Collins et al., 2020; Du et al., 2021; Zhou et al., 2019).

Regardless of coverage, it is important to note that genotypes called in short-read SV discovery programs are prone to high error rates (Chander et al., 2019). One strategy to alleviate this is to conduct SV discovery across all individuals to establish a SV call set and then use dedicated programs for SV genotyping (e.g., BayesTyper, SVTyper, Paragraph) (Chander et al., 2019). In addition, SV genotyping programs have variable performance across a range of SV sizes and types. As such, many programs are unable to genotype across the whole range of SVs that may have been called during discovery. A final consideration is that genotyping programs are still an area of active development. Although they may alleviate some of the false discovery rates prevalent in SV discovery programs, they do not provide a definitive solution (Chander et al., 2019).

As a result, putative SVs identified using short-read data alone should be treated as preliminary. One method that may aid in removing false variants is a trio-binning approach, whereby SVs that fail to abide by Mendelian inheritance patterns (either due to challenges with accurate genotyping or incorrect variant calls) are removed from the SV call set (e.g., Patel et al., 2014; Pilipenko et al., 2014). This is particularly relevant for intensively managed threatened species, as pedigree relationships are often known, and suitable trios can be identified for sequencing. Alternatively, if sufficient resources are available, the sensitivity of long-read sequence data may be leveraged across multiple reference genomes approach for SV discovery and may facilitate the use of lower coverage short-read resequencing data for SV genotyping, but this largely remains untested. Target-capture methods for genes or regulatory regions of interest can also provide an affordable alternative to WGS for species with large genomes and projects with tight budgets (Andermann et al., 2020). For example, vonHoldt et al. (2017) used a targeted-sequencing approach combined with PCR validation to characterize a region on chromosome 6 under positive selection in domestic dog breeds, which significantly reduced sequence coverage requirements.

4.5 | SV discovery and genotyping beyond short-read sequence data

When characterizing genomic features, especially SVs, there are many sequencing platforms and approaches to choose from, and although they may perform well when addressing specific challenges, each has its own caveats (Table 1). Two providers prominently feature in long-read sequencing: Pacific Biosciences (PacBio) and Oxford Nanopore Technologies (ONT). Since the launch of these technologies in 2011 and 2014, the long-read sequencing space has been characterized by fast-paced progress and innovation as demonstrated by the first telomere to telomere assembly of the human X chromosome achieved with ultra-long-read sequencing (Miga et al., 2020). The precise error rates between these two technologies remain somewhat contentious (Dohm et al., 2020; Lang et al., 2020 preprint), but as a general rule, ONT currently provides longer average read lengths than PacBio overall (Logsdon et al., 2020) at the cost of higher sequence error rates. Further, when PacBio sequencing providers are overseas, ONT offers the advantage that data can be generated in-house, which may, for example, be more responsive to the needs and aspirations of Indigenous Peoples (Collier-Robinson et al., 2019; Galla et al., 2016) and/or alleviate the need for export permits. Despite the challenges, the power of long-read sequencing technologies to span a significant portion, if not the entire length, of complex regions of the genome in a single read provides a powerful tool for SV discovery and population-level genotyping. When used in conjunction with a high-quality, well annotated reference genome, this improves confidence in read mapping across the genome (Amarasinghe et al., 2020), and substantially increases precision (the proportion of variant calls that are “true”) and recall (the proportion of “true” SVs detected) rates for both SNPs and SVs (Wenger et al., 2019). In addition, platforms that directly sequence native DNA remove the amplification bias common
| Feature                                      | Short-reads                                                                 | Long-reads                                                                 | Multiple reference genomes                                                                 | Optical mapping                                                                                     |
|----------------------------------------------|-----------------------------------------------------------------------------|-----------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Type of SV that can be assessed              | Deletions, Insertions, Inversions, Copy Number Variants, and some Translocations | Deletions, insertions, inversions, translocations, duplications, complex SVs | Deletions, insertions, inversions, translocations, duplications, complex SVs                     | Deletions, Insertions, Inversions, Translocations, Duplications                                      |
| High-quality reference genome                | Required                                                                    | Required                                                                    | Required (multiple)                                                                            | Optional                                                                                           |
| Technology used to generate data for SV discovery and genotyping | Illumina HiSeq, NovaSeq, NextSeq                                              | Pacific Biosciences (PacBio) Sequel; Oxford Nanopore Technologies (ONT)     | 1. Multiple reference genomes generated using a combination of short-, long-, linked-reads (e.g., Hi-C, Pore-C) for representative individuals 2. Short- and/or long-reads generated for population-level genotyping | Bionano Genomics Saphyr                                                                               |
| DNA quality                                  | Moderate to high molecular weight                                           | High to ultra-high molecular weight<sup>a</sup>                            | High to ultra-high molecular weight<sup>a</sup>                                                | Ultra-high molecular weight<sup>a</sup>                                                            |
| Length of SVs detected                       | 50 bp to <1 Mb                                                             | 50 bp to >1 Mb                                                             | 50 bp to >1 Mb                                                                                 | 500 bp to >1 Mb                                                                                   |
| Sequence coverage required                   | ≥10x                                                                        | ≥10x                                                                        | 1. ≥10x to generate multiple reference genomes 2. ≥10x for population-level genotyping         | >30x                                                                                              |
| Method of SV discovery                       | Alignment-based, including read-pair, read depth, split-reads and local assembly | Alignment-based with local assembly                                        | 1. Alignment of multiple de novo assembled whole reference genomes; 2. Genome graphs               | Alignment-based, either to reference genome or to optical maps from different samples             |
| SV genotype evidence                         | Multiple sources of evidence required depending on SV type (read pair, read depth, split read) | Single source of evidence (long continuous reads) usually sufficient        | Two sources of evidence required (multiple references and population-level sampling)             | Order, position, and orientation of fluorescently-labelled sequence motifs                          |
| Example algorithms/programs                  | BreakDancer; LUMPY; Manta                                                  | SMRT-SV (PacBio); NanoVar (ONT)                                            | Minimap2; MuMmer; Graphtyper2                                                                  | Bionano SVCaller                                                                                  |
| Proportion of SVs discovered across the genome | Moderate                                                                   | High                                                                        | High                                                                                            | High                                                                                             |
| Recall rate (proportion of “true” variants that are detected) | Low                                                                        | High                                                                        | High                                                                                             | High                                                                                             |
| False positive rate for detection            | High                                                                        | Low                                                                         | Low to moderate                                                                                  | Low                                                                                               |

<sup>a</sup> Depending on the technology used.
in many short-read sequencing approaches (Depledge et al., 2019). Furthermore, there are emerging “adaptive” sequencing approaches that have the potential to selectively sequence specific regions of the genome (Payne et al., 2020 preprint).

Structural variants significantly alter genome topology and impact the gene regulatory landscape (Sadowski et al., 2019; Shanta et al., 2020). In light of these impacts, the hierarchical organization of DNA within the nucleus is of particular interest when investigating the relationship of transcriptional regulation mechanisms.

Chromatin conformation capture (3C) based sequencing approaches enable the investigation of the organization of chromatin across genomes (Kong & Zhang, 2019) and have identified the chromatin signature in gene expression (Lieberman-Aiden et al., 2009; Lupiáñez et al., 2015; Shanta et al., 2020). In addition, there are emerging advancements in Nanopore sequencing methods to integrate chromatin conformation capture with long-read sequencing (i.e., Pore-C; Ulahannan et al., 2019 preprint). Rather than the amplification bias introduced by preparing a short read library, long-read sequencing provides data on chromatin at a range of distances along the linear genome and enables contacts to be sequenced without amplification. This information is particularly beneficial for non-model species where gene annotations, regulatory element annotations and gene regulatory networks are un- or poorly-characterized, leading to challenges in predicting the impacts of SVs on gene expression. For example, 3C-based approaches combined with SV detection and analysis have revealed the molecular basis for human developmental diseases previously diagnosed through karyotypes (Melo et al., 2020) and resolved the mechanisms underlying reproductive disorders in goats (Guang-Xin et al., 2021).

Optical mapping approaches are a useful complement to long-read sequencing approaches, and have enhanced genome assembly outcomes by providing insights into the “big picture” of large-scale genomic variants (as per Weissensteiner et al., 2020). Optical mapping utilizes a technique based on light-microscopy to identify specific sequence motifs (such as restriction enzyme cut sites), which are then used to generate images of fluorescently-labelled DNA molecules (Schwartz et al., 1993), enabling the characterization of large, complex rearrangements missed by long-reads alone (Yuan et al., 2020). On average, optical maps span 225 kb, providing information on the physical distance and relationship among genomic features. Besides being used to improve the scaffolding of genome assemblies (Howe & Wood, 2015; Zhang, 2015), including those of endangered species (Rhie et al., 2021), optical mapping methods directly enable the identification of both intra- and interspecific SVs (Levy-Sakin et al., 2019; Zhiihai et al., 2016). The primary current commercial provider of optical mapping technology is Bionano Genomics and their Saphyr instrument, which uses a nano-channel microfluidic chip to linearize and capture images of fluorescently-labelled ultra-long DNA fragments to generate optical maps at a resolution of 500 bp (Yuan et al., 2020). While optical maps provide information on the physical topology of chromosomes, they do not provide sequence information on an allele. Because long-reads and optical maps complement each other, the ideal data set for SV

| Feature | Short-reads | Long-reads | Optical mapping |
|---------|-------------|------------|------------------|
| Choice of method influences number and type of SVs detected | High | Moderate, depending on sample number | Moderate, depending on sample number |
| Benchmark data must be verified for non-model species | High | Moderate, depending on sample number | Moderate to high |
| Low precision for detecting complex SVs, near SNPs, and small indels | Moderate | High | High |
| Low performance in low complexity regions | Moderate | High | High |
| Low molecular weight DNA ≤20 Kb fragments; ultra-high molecular weight DNA ≥200 Kb fragments | High | Moderate, depending on sample number | Moderate, depending on sample number |
discovery would include both data types (e.g., Soto et al., 2020; Weissensteiner et al., 2020).

5 | CONCLUDING REMARKS

Structural variants provide an exciting opportunity to complement SNP-based approaches and expand our understanding of genome-wide variation. However, there remains much to learn about the integration of SVs in the small population paradigm. In the meantime, mounting evidence indicates that SVs play an important role in determining fitness traits for both model and non-model species. Whereas SNPs impact single nucleotides, SVs can capture multiple genes and gene regions simultaneously and are therefore more likely to impact fitness. Emerging approaches for SV discovery and genotyping, and relating SVs to fitness traits, are now in reach for threatened species, and we are confident that discoveries related to the role of SVs in determining fitness traits of conservation interest are imminent. Moreover, the development of SV-specific tools for analyses of within and between population variation will further enhance the utility of SVs in species of conservation concern (e.g., SV-Pop; Ravenhall et al., 2019).

However, developing strategies to manage fitness traits will be challenging, as selection for or against them can lead to the reduction of genome-wide diversity (Kardos & Shafer, 2018; Kardos & Shafer, 2021 preprint). Despite this, current management practice can include such actions. For example, individuals that carry, or express, negative fitness traits can be assessed in conservation breeding programs, followed by appropriate actions or recommendations (e.g., Attwater’s prairie chicken; Hammerly et al., 2013). Alternatively, when inheritance of these traits is known, individuals may be screened prior to selective pairing (e.g., California Condor; Moran et al., 2021). Regardless, as we learn more about the mechanisms underlying fitness traits, especially traits associated with negative fitness, there is a pressing need to establish comprehensive evidence-based management strategies that mitigate the impact of these traits while minimizing the loss of genome-wide diversity. Where we see the most promise for this is the emerging practice of integrating WGS data into individual-based models, which will enable different management actions to be modelled and compared (Seaborn et al., 2021). Such models will allow researchers to forecast the consequences of selection-based management of fitness traits underpinned by adaptive or deleterious SVs on genome-wide diversity, thus better informing conservation management actions. To achieve this, strong collaboration between conservation practitioners, population ecologists (e.g., modelers), quantitative geneticists, and conservation genomicists will be required to establish the most effective models possible (Hohenlohe et al., 2020).

ACKNOWLEDGEMENTS

We wish to thank the Guest Editors of the “Whole-genome sequences in Molecular Ecology” Special Issue for the invitation to contribute this Perspective and three anonymous reviewers for their insightful comments and constructive feedback throughout the peer review process. We are grateful to the Kākāpō125+ Consortium led by Genomics Aotearoa for providing the opportunity to explore structural variants in a conservation context. We extend our thanks to the University of Canterbury’s Conservation, Systematics and Evolution Research Team (ConSERT), including Roger Moraga at Tea Break Bioinformatics for robust dialogue—and Kylie Price and Joanna Roberts for their support and guidance—on this topic. We acknowledge the University of Canterbury, New Zealand Department of Conservation and Genomics Aotearoa for their support of JRW and NSF Track 2 EPSCoR Program (award number OIA-1826801) for their support of SJG.

AUTHOR CONTRIBUTIONS

J.R.W., K.P.K., A.W.S. and T.E.S. were the lead investigators on this perspective. J.R.W. designed all figures and K.P.K. contributed Table 1. J.R.W. led the writing of the manuscript. All authors contributed to manuscript drafts, provided constructive responses to reviewer feedback and gave final approval for publication.

DATA AVAILABILITY STATEMENT

Not applicable.

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