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Enhancing the accuracy of next-generation sequencing for detecting rare and subclonal mutations

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Abstract | Mutations, the fuel of evolution, are first manifested as rare DNA changes within a population of cells. Although next-generation sequencing (NGS) technologies have revolutionized the study of genomic variation between species and individual organisms, most have limited ability to accurately detect and quantify rare variants among the different genome copies in heterogeneous mixtures of cells or molecules. We describe the technical challenges in characterizing subclonal variants using conventional NGS protocols and the recent development of error correction strategies, both computational and experimental, including consensus sequencing of single DNA molecules. We also highlight major applications for low-frequency mutation detection in science and medicine, describe emerging methodologies and provide our vision for the future of DNA sequencing.

Clonal

When referring to a genetic variant or mutation, it is one that is present in all or most molecules in a population being sequenced. The term typically implies that it arose from a common ancestor, such as a fertilized egg in the case of germline variation, or the earliest founder cell of a tumour.

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Genetic heterogeneity underlies the evolution and adaptation of all life on Earth. This is equally true of stochastically generated variants in germ cells as it is of somatic cells within tumours and ageing tissues. Rare variants can selectively proliferate upon exposure to new environments through natural selection^{1–3}. Intercellular genetic diversity underlies many elements of human disease, including the emergence of therapeutic resistance to antimicrobial and anticancer therapies^{4,5}, the development of certain inherited genetic diseases⁶ and ageing and age-associated pathologies^{7,8}. Yet despite its importance, until the past decade, our tools for quantifying and studying genetic variation in heterogeneous cell populations have been limited.

Beginning in 2005, a new generation of tools, referred to by the now somewhat anachronistic moniker of next-generation sequencing (NGS), emerged and wholly reshaped genetics⁹. NGS technologies have reduced the cost and increased the scale of genomic investigations by many orders of magnitude. They have transformed the study of genetic variation in humans and model organisms, elucidated the genetic basis of some diseases and advanced our understanding of the organization, regulation and function of genomes with unprecedented granularity¹⁰. Multiple distinct NGS platforms now exist, but all share the same fundamental feature of parallel interrogation of millions of individual DNA templates. The digital nature of the approach stands in contrast to the prior gold-standard Sanger method of sequencing aggregate populations of molecules¹¹. Although

this makes NGS methods potentially more sensitive for quantifying genetic heterogeneity, the vastly higher throughput means that, in practice, the absolute number of errors to contend with is greater than ever before.

Nearly all measurements in science are limited by the signal-to-noise ratio of the assay (FIG. 1), and genetic heterogeneity is no exception. The rarer the variant, the more sensitive a technique must be to find it. Historically, scientific aspirations in this field exceeded the capabilities of available tools. For germline sequencing where variants are clonal, high-confidence genotypes can still be obtained, despite the modest accuracy of standard NGS, by redundantly sequencing identical genomic copies from multiple cells of an individual and ignoring erroneous read-to-read variation. By contrast, shortcomings in accuracy fundamentally limit the sensitivity of routine NGS for detection of low-level genetic variation in subclonal populations encompassing fewer than ~1% of the DNA molecules in a sample^{12,13}. This has been a particular challenge for the reliable identification of somatically acquired mutations in multicellular organisms and for disentangling mixed microbial populations. Several recent advances have now markedly improved NGS accuracy and therefore our capability to detect rare variants; the deeper we have been able to look, the more reasons we are now discovering to look even deeper.

In this Review, we summarize the transformative impact of NGS on resolving molecular heterogeneity and survey the technical evolution of computational,

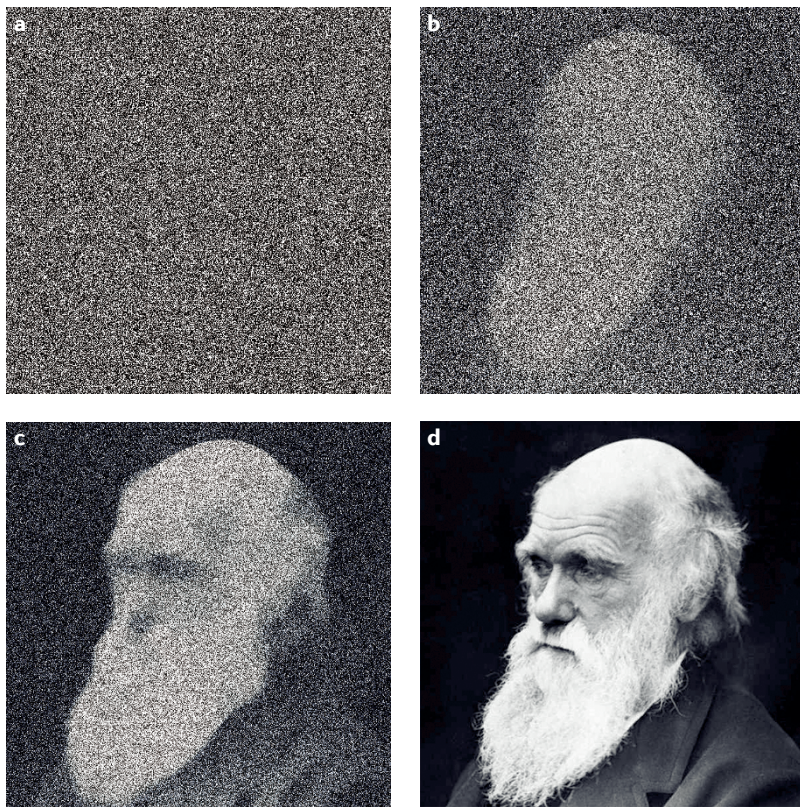


Figure 1 | The signal-to-noise problem. The accuracy of all analytical measurements, DNA sequencing included, depends on the ratio between true value and the precision of the detection method. This is analogous to the noisiness of a digital camera image: at a low signal-to-noise ratio, an image is indecipherable (part **a**); with increasing sensor quality an oval shape appears (part **b**), becomes apparent as a face (part **c**) and then as a specific individual (part **d**). Figure is adapted from REF. 205, CC-BY-4.0 (<https://creativecommons.org/licenses/by/4.0/>).

biochemical and recently developed single-molecule consensus methods for error correction that help mitigate the inherently high error rates of NGS platforms. We discuss emerging technologies that have the potential to further enhance our understanding of the role of rare genetic variants and highlight major fields of research that either have or will soon benefit from advances in sequencing accuracy for the purpose of obtaining higher sequencing sensitivity. We close with a discussion of future opportunities and the next wave of technical sequencing challenges we see on the horizon.

Subclonal mutation detection

For many areas of medicine and biology, genetic heterogeneity is the rule rather than the exception. Although several sensitive technologies for low variant allele frequency (VAF) mutation detection predated the advent of NGS, these were limited to interrogation of very small genomic regions and not easily transferred between loci^{14,15}. Various methods for selective amplification of low-frequency variants facilitate detection but lack the ability to precisely quantify their relative abundance¹⁶. Digital PCR is a powerful technique that can be used for both precise molecular counting¹⁷ and, in allele-specific forms, robust low-frequency mutation detection¹⁸.

Subclonal

When referring to a genetic variant or mutation, it is one that is present in only a subset of molecules being sequenced. This may refer to either a variant carried by a subpopulation that arose and expanded within a larger population or through mixing of two or more distinct populations.

Sequencing accuracy

The number of errors made per base pair sequenced. It may be stratified by subtype of error, such as a specific type of base substitution.

In recent years, the method has become more widespread with the advent of convenient high-throughput emulsion-based platforms, referred to as digital droplet PCR¹⁹; however, variants being sought typically must be known a priori. NGS is indisputably the most generalizable method of mutation detection, but it has been only recently that technical advances have allowed it to achieve a comparable accuracy.

At the outset, we acknowledge that comparing the accuracy of different NGS protocols is challenging. The error rate of conventional NGS is approximately 1% and can be as low as 0.1% in optimal scenarios²⁰. However, the precise value varies by specific platform, chemistry version, sequence context, filtering stringency and various other factors that make for lively discussions but few hard-and-fast agreements among researchers. The accuracy of many error-correction methodologies is similarly affected by variables that are difficult to normalize between studies, such as degree of DNA damage and whether the DNA standard being sequenced is, itself, truly free of mutations. Diluted oligonucleotide templates with defined sequences are often used to represent rare variants in published mixing experiments, but the frequency of errors during oligonucleotide synthesis may be as high as 1 in 100. Similarly, although standardized cell lines are an attractive source of DNA that can be benchmarked against by different researchers, it has been shown that, in at least some situations, the mutations that accumulate during a few months in culture can exceed those accrued over an entire human lifetime²¹. For all these reasons, we have opted to limit our comparisons of method accuracy to orders of magnitude.

Improving the accuracy of NGS

Most attempts to lower the limit of detection for commercially available NGS platforms can be grouped into one of three broad categories. The first category consists of purely computational and statistical strategies to exclude sequences of low confidence from conventionally generated NGS data. The second strategy consists of library preparation protocols that either biochemically remove or limit the formation of mutagenically damaged nucleotides in templates before sequences are generated. We review both of these categories in this section. The third and most successful type of strategy is single-molecule consensus sequencing, which we describe in subsequent sections. This more contemporary latter approach involves an intertwined combination of chemical labelling before sequencing and informatic deconvolution thereafter, which allows for the identification and exclusion of errors that invariably occur despite all other measures.

Computational reduction of sequencing artefacts.

Initial efforts to reduce the background error rate of NGS focused on data filtering schemes to discount low-confidence sequences caused by technical artefacts. Phred quality scores, originally developed for Sanger sequencing electropherograms to estimate the probability of an error at each sequenced base²², were adapted for the image-based output of NGS platforms²³.

Quality score filtering does not improve accuracy if the errors are introduced before the sequencing, for example, during PCR amplification. A variety of bioinformatics tools, such as MuTect²⁴ and VarScan2 (REF. 25), use additional filters, such as whether variants are biased towards the beginning or ends of reads (reflecting erroneous end-repair of fragments or mapping errors), or require that true variants be seen in multiple independent sequencing reads or both read orientations²⁶. Newer alignment algorithms and read-trimming tools are better able to avoid artefacts from off-target mapping or inadvertent sequencing into artificial adapter or primer sequences used in library preparation²⁷. Increasingly sophisticated software packages that are specifically designed for the detection of low-frequency variants in traditional NGS data are able to eliminate many false calls through rigorous statistical approaches that involve modelling the error profile of specific sequencing applications, or even individual sequencer runs, and using these to appropriately assign confidence to particular conclusions^{28,29}.

Some sequencing errors can be identified and removed empirically by simultaneously sequencing a test sample that is thought to have subclonal mutations (for example, a tumour) alongside an apparently homogeneous control sample³⁰. Low-level variants in the control sample are assumed to reflect sites with a higher propensity for technical errors and are discounted from all samples. As increasingly large sequencing data sets are generated, databases of specific sequence contexts that recurrently yield artefactual errors can be identified and viewed more sceptically. Particular mutation patterns, such as those resulting from oxidative DNA damage^{31,32}, or engineered polymerases and nucleotides used during sequencing³³ can be partially remedied informatically. Nevertheless, even the most careful computational data scrubbing cannot universally produce high-confidence calls of subclonal mutations much below 1% abundance^{13,34–36}, and the more aggressive the approach, the greater the risk of excluding true rare variants.

Biochemical reduction of sequencing artefacts. Errors arising during the generation of NGS data can occur at many stages. Mistakes can occur on the sequencer itself as a result of optical imperfections, enzymatic errors during cluster formation or overlapping or polyclonal clusters, among other issues^{33,37}. Substantial errors can also arise during pre-sequencing library preparation as a result of PCR misincorporations, chimeric PCR products, template switching or hairpin formation^{37–40}. Although high-fidelity proofreading polymerases are typically used for amplification steps, lower-fidelity polymerases may be used for preceding repair and A-tailing of library fragments. Furthermore, all polymerases are considerably more error prone when copying across damaged nucleotide templates. Such damage may be present at the time of sampling from normal cell processes or environmental exposures but can also occur from extrinsic manipulations, for example chemical extraction, heating or clinically used stabilization methods such as formalin fixation^{41,42}.

A common step in library preparation is ultrasonic shearing of DNA into short fragments. This produces sufficient energy to break the phosphodiester backbone, which can also oxidize bases and lead to artefactual C:G to A:T transversion mutations. This can be somewhat reduced by pH buffering and cation chelation^{31,32}. Furthermore, nicks and non-blunt-ended breaks produce regions of single-stranded DNA, which are both biochemically more susceptible to damage⁴³ and subjected to copying by lower-accuracy polymerases during end-repair steps in adapter-ligation-based library preparation methods. Enzymatic DNA fragmentation methods avoid some of these issues, but at the same time may produce other low-level artefacts as a result of nicks, abasic sites or other incomplete cleavage products that vary by the enzymatic mechanism⁴⁴.

Although some DNA damage may be prevented, other sources are unavoidable. Both formalin-fixation and heating accelerate the rate of spontaneous deamination, particularly of cytosine to uracil, to produce C:G to T:A substitutions upon PCR amplification^{42,45}. Uracil bases in DNA can be excised by treatment with uracil-DNA glycosylase (UDG), which yields abasic sites that are resistant to amplification^{46,47}. However, abasic sites are, themselves, somewhat mutagenic if copied: polymerases typically mis-insert an A when encountered^{48,49}, which also generates a single nucleotide C:G to T:A substitution. Destruction of the remnant phosphodiester backbone at abasic sites with a DNA lyase, commonly endonuclease VIII, may somewhat abrogate this⁴².

Another common mutagenic base modification, 8-oxo-dG, is formed by oxygen free radicals and readily mis-pairs with adenine to lead to C:G to A:T mutations⁵⁰. 8-Oxo-dG can be biochemically excised by incubating with formamidopyrimidine [fapy]-DNA glycosylase (FPG), which has both glycosylase and lyase activity^{42,46}. As with UDG, treatment can reduce artefacts but may also render some highly damaged DNA unamplifiable. This can be a particular problem with some low-input applications described later. Combinations of glycosylases with other repair enzymes to replace the damaged base may improve amplifiability³², yet this treatment itself may introduce errors at low levels.

Nevertheless, although biochemical approaches, in concert with computational strategies, have a positive effect on improving NGS accuracy, it is relatively modest. Not all mutagenic damage can be prevented, and not all damage that occurs can be easily corrected. For example, spontaneous deamination of 5-methylcytosine generates a canonical thymine base. To achieve error rates below 1 in 1,000, other techniques are required.

Molecular consensus sequencing strategies

Following the release of the first commercial NGS platform in 2005, efforts to improve accuracy initially focused on refining the core elements of sequencers themselves. During the next several years, iterative improvements to optics, polony formation methods, sequencing-by-synthesis chemistry and on-machine data filtering boosted the accuracy of raw outputted data

Sequencing sensitivity

The ability to detect a variant at a particular variant allele frequency. This depends on both the sequencing accuracy and the number of independent DNA molecules successfully sequenced that include the genomic position (or positions) of interest.

Variant allele frequency

(VAF). The fraction of all molecules being sequenced that carry a specific genetic change or mutation at a particular genomic position.

Digital PCR

DNA amplification carried out in single-molecule reaction chambers. Recently, this has most often entailed microscopic aqueous droplets immersed in oil. When DNA input is sufficiently low, only one molecule will seed each reaction. When allele-specific amplification conditions are used, the number of droplets that successfully amplify can be digitally tabulated to determine the variant allele frequency.

Polony

A population of identical amplification copies that originated from a single founder molecule and are spatially colocalized, such as on the surface of a microbead or as a spot on a surface. It is the biochemical analogue of a bacterial colony on a Petri dish.

Tag-based error correction

Also known as consensus sequencing, an approach for error correction whereby individual DNA molecules are uniquely labelled before amplification and sequencing, and the sequences of the related derivative copies are then compared with each other to exclude errors.

Short-read platforms

Next-generation sequencing systems that generate reads that are dozens to several hundreds of nucleotides in length, for example, the current Illumina and Thermo Fisher Scientific Ion Torrent platforms and previously manufactured Roche 454 and ABI SOLiD platforms. Current versions sequence amplified colonies, not single molecules.

Long-read platforms

Next-generation sequencing systems that generate reads that are thousands to tens of thousands of nucleotides in length. These currently include Pacific Biosciences (PacBio) and Oxford Nanopore Technologies, which sequence single molecules, not colonies, and therefore have a higher error rate than short-read platforms.

Molecular barcode

Also known as a unique molecular identifier (UMI). A set of DNA nucleotide codes where each is affixed to only one or a subset of individual DNA molecules within a sample. The purpose is to uniquely label single molecules for consensus-based error correction or molecular counting. These may be informatively combined with molecule fragmentation points for greater label diversity.

Index sequence

A particular DNA nucleotide code affixed to all molecules within a given DNA sample that is used for multiplexing samples on a single sequencer run.

Sequencing depth

The number of sequencing reads that include a particular genomic position in their sequence. Some may be simply PCR copies of the same molecule.

by about an order of magnitude. Paired-end sequencing allowed additional confirmation of the identity of bases sequenced from both ends of a molecule. During the past several years, although NGS throughput has increased dramatically, improvements to raw accuracy have largely plateaued; on some high-output platforms, it has even decreased⁵¹. Recognizing that some biochemical mistakes are unavoidable, in approximately 2009, an innovative solution for improving accuracy was developed that focused on identifying and ignoring errors, rather than preventing them entirely⁵². The approach, eventually becoming known as single-molecule consensus sequencing, tag-based error correction or molecular barcoding, rapidly emerged through the work of multiple investigators to become a new standard for high-accuracy NGS applications^{53–55}. We begin with reviewing different embodiments of this concept that apply to the short-read platforms, which currently comprise most of the NGS market. In the subsequent section, we discuss consensus sequencing approaches that apply to the commercially available long-read platforms that rely on direct sequencing of single DNA molecules.

Barcoding of individual DNA molecules. During conventional short-read platform NGS, a DNA library is typically PCR amplified before sequencing. It is often impossible to definitively know whether two identical sequence reads arose from copies of the same starting molecule or from two independent molecules. However, if a unique tag (that is, a molecular barcode) is applied to each molecule before amplification, this label will be propagated to all derivative copies and independent sequence reads can thus be recognized as having arisen from a common founder. It is worth noting that the concept of a molecular barcode (also known as a unique molecular identifier (UMI), a single-molecule identifier (SMI) or simply a tag) is different from that of an index sequence. Molecular barcodes serve to uniquely label individual molecules within a sample, whereas index sequences are identical DNA labels that are affixed to all molecules in a given sample for the purpose of sample multiplexing.

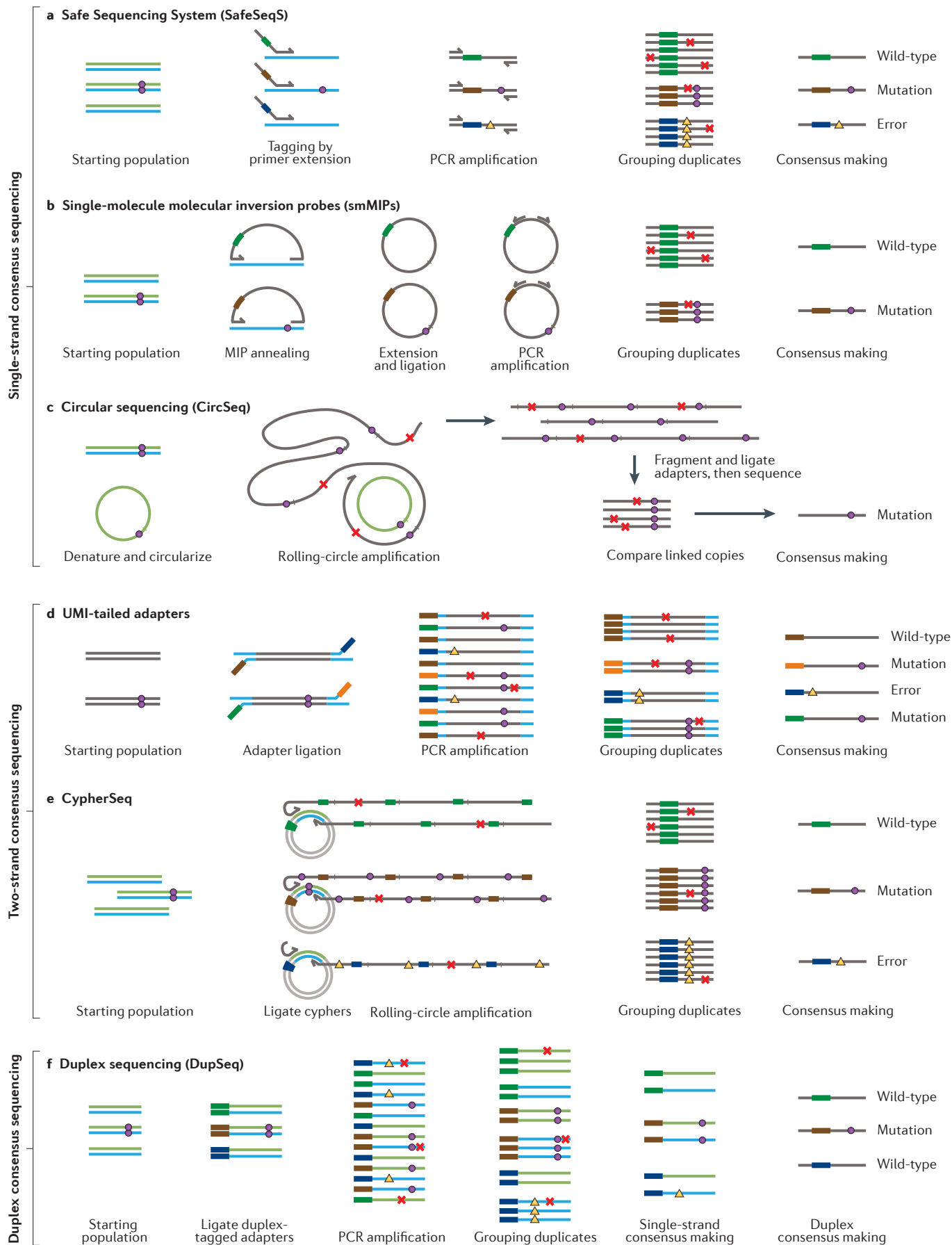
Molecular barcodes can be used to improve the accuracy of counting DNA or RNA molecules in mixtures by eliminating biases from variable amplification^{54,56–59}. More importantly, because all identically tagged reads will have derived from a common founder (provided that barcodes are designed carefully), any variation between their actual sequences must necessarily reflect technical errors^{53–55}. Tag-based error correction relies on this principle: independent reads sharing a common tag are recognized and grouped as amplicon copies of the same starting molecule; any sites of sequence differences among the reads are discounted as errors when forming a consensus sequence (FIG. 2). A fundamental element of the approach is the need to intentionally produce and sequence redundant molecular copies, which requires relatively higher raw sequencing depth than conventional NGS and, thus, additional costs.

Molecular barcodes come in two forms: exogenous and endogenous^{53,55}. Exogenous barcodes entail random

Figure 2 | Methods of consensus-based error correction on short-read platforms. a

The Safe Sequencing System (SafeSeqS) uses randomly generated molecular barcodes carried by PCR primers (coloured thick bars) to reduce errors by independently labelling each single-stranded DNA molecule, thus allowing identification of derivative copies. True mutations (circles) can be discerned from sequencing errors or late PCR errors (crosses) because the latter occur only in a subset of identically labelled duplicate reads. PCR errors that occur during the first cycle of amplification (triangles) can be propagated to all duplicates and escape error correction. **b** | Single-molecule molecular inversion probes (smMIPs) entail two targeting arms joined by a linker that contains a molecular barcode. The molecules are hybridized with single-stranded DNA and then extended and ligated to form closed loops that are amplified and sequenced. Consensus-based error correction is similar to SafeSeqS and similarly susceptible to first-cycle amplification artefacts. **c** | Circular sequencing (CircSeq) entails circularization of single-stranded DNA fragments without any molecular barcodes followed by rolling-circle amplification, fragmentation and sequencing of short stretches of concatemered fragments. The molecular fragmentation points of the starting molecules serve as unique molecular identifiers (UMIs) for consensus-based error correction. As with other single-stranded consensus methods, recurrent amplification errors may fail to be identified and corrected. **d** | UMI-tailed adapters can be ligated to a library to uniquely mark each single strand. Despite both strands in a complex being tagged, no means are provided to relate the consensus of one strand to that of its mate for comparison, and early PCR errors (triangles) may go unrecognized. **e** | CypherSeq circularizes double-stranded DNA molecules using a single adapter-molecule-containing double-stranded molecular barcode. Targeted enrichment is achieved with rolling-circle amplification using primers directed to each DNA strand. Although information from both strands may contribute to consensus making, lack of asymmetry between the two strands makes it impossible to discern whether one or both strands is successfully amplified. Recurrent early amplification errors (triangles) can escape error correction when only one strand worth of data is successfully recovered because this cannot be recognized. **f** | Duplex sequencing (DupSeq) allows true duplex error correction on high-throughput short-read sequencing platforms by applying molecular barcodes to each double-stranded DNA molecule in such a way that amplification products of the two strands can be informatively related to each other (thick coloured bars) but also distinguished (blue versus green strands). After tagging, derivative PCR products are grouped by molecular barcode and by strand. Consensuses are made for each strand group and then compared to that of the complementary strand. True mutations (circles) can be confidently distinguished from both sequencing errors and late PCR errors (crosses) as well as first-round PCR errors (triangles) because complementary errors are extremely unlikely to occur by chance at the same position on both DNA strands. See the main text for a detailed description of each method. MIP, molecular inversion probe.

or semi-random artificial sequences that are incorporated into either sequencing adapters or PCR primers. Endogenous barcodes describe the randomly or semi-randomly generated fragmentation points at the



ends of DNA molecules in ligation-based library preparation methods. The two approaches can be used either alone or in combination⁶⁰.

With either approach, it is important that a sufficient variety of possible tag sequences exist such that the probability of two independent molecules being tagged the same way is low. With sequencing at low molecular depth, the chance of two independent DNA fragments having the same shear points by chance is small, and these endogenous sequences alone suffice as tags^{60,61}. At the other extreme is deep sequencing following an amplification-based library preparation. In this case, molecular ends are defined by invariant primer sites, not random fragmentation, so all tag information must come from exogenous tags⁵⁵. A similar problem arises with targeted enzymatic fragmentation⁶². If barcode diversity is inadequate, tag clashes can occur, whereby independent molecules are identically labelled. In this scenario, true low-frequency variants can be erroneously discarded as errors. If barcodes are too complex, they may develop errors themselves and artificially create false families that incorrectly appear as arising from distinct molecules. Both problems can be mitigated with careful design and strategies for tolerating errors in barcodes^{63–65}.

Over the past 5 years, molecular consensus sequencing has proved itself as the most impactful means for reducing NGS errors. Different implementations variably reduce sequencing error rates from $\sim 10^{-2}$ to 10^{-4} – 10^{-7} or lower. The variety of approaches developed to date can be grouped into three basic categories: single-strand consensus sequencing; two-strand consensus sequencing; and duplex consensus sequencing (FIG. 2).

Single-strand consensus sequencing. One of the most widely cited early implementations of tag-based NGS error correction is the safe sequencing system (SafeSeqS) technique, which applies tags via PCR primers carrying a degenerate sequence tail⁵⁵. In this method, after a small number of PCR cycles with the barcoded primers, additional amplification with a second set of universal primers is carried out to generate multiple copies of each tag-defined founding molecule, which are then sequenced and grouped into families for consensus-based error correction (FIG. 2a). A substantial challenge to widespread implementation was the difficulty of sequencing large numbers of targets at the same time because of PCR multiplexing problems caused by the random tags. Newer variations on the technique that protect the tags from nonspecific binding through the use of a hairpin design have somewhat reduced this problem^{66,67}.

Primer-based tagging is convenient in so far as it facilitates sequence targeting at the same time as tagging for error correction, but because several PCR tagging cycles are needed, the same DNA molecule will occasionally be labelled by more than one tag and create false families. Another tagging method that circumvents this issue is the use of single-molecule molecular inversion probes (smMIPs)^{68,69}. Instead of a pair of primers, a single oligonucleotide with two targeting arms connected by a linker region with a molecular barcode is

hybridized to a DNA sample and then extended and ligated to form tagged, closed-loop products that can then be enriched, amplified and sequenced (FIG. 2b). With smMIPs, many targets can be easily multiplexed together, and there is little risk of double-tagging the same molecule. Design constraints around the narrow proximity window for the targeting arms add a challenge, but improved software algorithms have recently made the method more tractable⁷⁰.

Another consensus sequencing method that takes a very different approach is circular sequencing (CircSeq)^{46,71}. In this technique, DNA is fragmented and melted into very short single-stranded pieces that are then circularized and copied into concatemers via rolling-circle amplification. The concatemers are further amplified and then sequenced (FIG. 2c). Instead of exogenously applied barcodes, the unique genomic coordinates of fragmentation points serve as molecular identifiers to define which sequence reads derived from a given starting molecule. The fact that tandem copies of the sequence are physically joined means that each sequencing read contains the necessary information for an initial level of consensus calling. In contrast to tag-based barcoding of unlinked copies, which may have either too few or an excess of copies present, this linkage improves cost-efficiency by keeping the duplicate rate more uniform. The two major challenges to the approach are the very limited length of sequences that can be genotyped as tandem copies on short-read platforms (some solutions have subsequently been developed⁷²) and the risk of tag clashes that stems from the use of the inherently limited number of possible shear points as identifiers⁷³. Nevertheless, the concept of sequencing tandem linked copies is powerful and will undoubtedly become more relevant as the performance of long-read platforms improves.

Most of the applications for NGS do not require an entire genome to be sequenced, and targeting of specific regions is important to reduce costs. NGS library preparation workflows involving DNA fragmentation, adapter ligation and then hybrid capture enrichment of loci of interest are slightly more time consuming than amplicon-based methods but are generally easier to design and, thus, are more widely used. One of the most easily implemented and popular consensus sequencing approaches that has made its way into many commercial products in the past 2 years incorporates a degenerate UMI sequence into one adapter strand⁷⁴. Depending on the implementation, either one or both (the version shown in FIG. 2d) of the library fragment strands are thus labelled. The combination of variable shear points and high-diversity exogenous UMIs substantially reduces the risk of tag clashes.

All single-strand consensus techniques reduce errors by two to three orders of magnitude, which is far greater than any prior computational or biochemical approach, and make it possible to accurately identify rare variants below 0.1% abundance. However, certain errors persist. All four methods rely on consensus sequencing of tagged copies derived from just one strand of what are natively double-stranded DNA molecules. Mistakes that occur

Molecular depth

The number of collapsed consensus reads derived from an independent DNA molecule that include a particular genomic position.

Tag clashes

The occurrence of two independent molecules being identically labelled by random chance. This may happen if the diversity of the applied molecular barcodes is too low for the number of DNA molecules sequenced. True mutations may erroneously be excluded.

False families

Sets of related molecules where an error has occurred during amplification that mutates the common tag sequence to erroneously make it appear that two independent molecules gave rise to these molecules.

during the first round of amplification can be propagated to all other copies as jackpot errors that escape correction (that is, the yellow triangles in FIG. 2a). This is particularly true of mis-incorporation errors at sites of mutagenic DNA damage, especially 8-oxo-guanine adducts and deaminated cytosine bases. This is clearly apparent in the spectrum of background errors from any single-stranded tagging method: G>T and C>T mutations stemming from oxidation and deamination, respectively, are far more frequent than the reciprocal mutations C>A and G>A. True mutations should be present in equal proportions of complements⁶⁰. The developers of CircSeq noted that this mutational bias could be partially abrogated by treatment with damage-removing glycosylases⁴⁶.

Two-strand consensus sequencing. DNA is a double-stranded molecule for a reason. The ability to unwind and independently copy each half facilitates cell division. The biochemically enforced rule-based pairing of nucleotides is relied upon by cellular machinery to ensure high fidelity of strand replication: any mismatches are quickly identified and repaired. Conceptually, a tag-based sequencing approach that takes into account the genotype of both DNA strands should achieve a higher degree of error correction for analogous reasons. One effort to improve upon the SafeSeqS method used bar-coded PCR primers targeted against both the reference and anti-reference strands of regions of interest⁷⁵. In concept, if a mutation is seen in both PCR products, it can be viewed with greater confidence. Despite being theoretically higher in accuracy than SafeSeqS, because the two PCR products that derive from the individual strands of any particular molecule will carry different random barcodes, the resulting sequences cannot be directly related to each other. Although both the reference and anti-reference strands of molecules in a population can be genotyped, there is no way to compare the sequence of one strand of a particular double-stranded molecule with that of the other, so true double-stranded error correction is impossible when concurrently sequencing multiple molecules. The same is true of the adapter-based UMI approach shown in FIG. 2d.

The first reported tag-based NGS error-correction method relied on a ligation-based approach where both strands of double-stranded molecules were labelled with identical molecular tags followed by PCR amplification, then concatemerization of amplicons and sequencing⁵³. After amplification, PCR products derived from both the reference and anti-reference strands carry the same tags and can be grouped to produce an error-corrected consensus. A more recent variant upon this method, known as CypherSeq⁷⁶, incorporates rolling-circle amplification from primers targeting both strands after ligation into a circularized adapter sequence to achieve a degree of target enrichment before PCR amplification (FIG. 2e). With both methods, PCR products derived from each strand of individual molecules can be used to form a consensus sequence; however, as the amplicons of the two strands are indistinguishable, it is impossible to tell whether the resulting consensus is based on single-strand or

double-strand data. Because one strand often fails to amplify, either owing to DNA damage or stochastic factors, or is simply not sampled⁶⁰, jackpot amplification errors can still escape detection (that is, the yellow triangles in FIG. 2e). A recent preliminary technique known as proximity sequencing (Pro-Seq) involves concurrent amplification of both the reference and anti-reference strands of individual DNA duplexes with physically linked primers in emulsion droplets⁷⁷. This enables genotyping of both strands within the same flow-cell cluster for cost savings but has an identical limitation in that amplification of both strands cannot be ensured.

Duplex consensus sequencing. In 2012, our group described duplex sequencing (DupSeq), a technique that uses a special form of molecular tagging to independently barcode each strand of individual DNA duplexes in such a way that sequence reads derived from one strand can be related to but also distinguished from that of the other⁶⁰. Exogenous tags within each strand of the sequencing adapters and/or DNA fragment shear points serve as UMIs that informatically relate reads from the two strands. A non-complementary portion of the adapters introduces strand asymmetry that allows the products to be distinguished from each other and, importantly, allows confirmation that both strands have been sequenced (FIG. 2f). To achieve true duplex error correction when the strands are separately amplified, the adapted molecule must contain both a UMI and an asymmetric strand-defining element (SDE). Together, these pieces allow a separate consensus sequence to be produced for each strand for comparison to that of its mate⁷⁸. In this way, early PCR jackpot errors can be confidently recognized and discounted (that is, the yellow triangles in FIG. 2f).

The theoretical DupSeq error rate of $<10^{-9}$ reflects the low probability of a complementary jackpot error occurring at the same position on both strands⁶⁰. In fact, one challenge to experimentally verifying this error rate is identifying a gold-standard source of DNA that is truly devoid of mutations: the lowest frequency of single-base substitutions we have measured in the DNA of newborn infants is between 10^{-7} and 10^{-8} , which is consistent with mutation frequencies extrapolated from differences between human generations⁷⁹. We and others have applied DupSeq to measure variations in the spontaneous mutation frequency of microbial populations⁶⁰ and in mammalian tissues in the setting of ageing^{61,80,81}, neurodegeneration^{82,83}, inherited DNA repair defects⁸⁴ and genotoxic exposures^{61,85}. In all cases, the mutagenic effect of near or below one mutation per million base pairs could only be recognized because of the extremely low error rate.

In theory, a variety of other molecular tools could serve as DupSeq UMIs and SDEs. Other than shear points and DNA-based tags, single-molecule compartmentalization methods that keep paired strands in physical proximity⁷⁷ or other non-nucleic acid tagging methods could serve the strand-relating function. Similarly, asymmetrical chemical labelling of the adapter strands in a way that they can be physically separated

can serve an SDE role. A recently described variation of DupSeq uses bisulfite conversion to transform naturally occurring strand asymmetries in the form of cytosine methylation into sequence differences that distinguish the two strands^{86,87}. Although this implementation limits the types of mutations that can be detected, the concept of capitalizing on native asymmetry is noteworthy in the context of emerging sequencing technologies that can directly detect modified nucleotides⁸⁸.

Having distinct elements that both relate and distinguish strands in DupSeq reflects the need to add molecular identity information about an original molecule that is lost when the paired strands are physically uncoupled and copies are made. Whereas this is currently the most practical approach with short-read platforms, for newer single-molecule long-read sequencing technologies, strand uncoupling and/or DNA amplification are often not required.

Direct single-molecule consensus sequencing

At the present time, only two types of single-molecule sequencers are commercially available: those manufactured by Pacific Biosciences (PacBio), which rely on detection of fluorescent nucleotide incorporation by single, immobilized polymerases within zero-mode wave guides⁸⁹, and those of Oxford Nanopore Technologies, which capitalize on the differential current changes caused by nucleotides as a single-stranded DNA molecule traverses a molecular nanopore between two chambers to record the sequence⁹⁰. These platforms currently make up a relatively small portion of all sequencing, largely because of lower raw accuracy and throughput than prevailing short-read technologies, but both have gained an increased following as the technologies have matured.

Because of their long reads, the most common use for single-molecule sequencers is *de novo* genome assembly and sequencing of complex repetitive regions or structural rearrangements^{90,91}. However, long-read capability can be repurposed to improve genotyping accuracy of shorter regions through sequencing of tandem copies. One example is the intramolecular-ligated nanopore consensus sequencing (INC-Seq) method whereby the terminal 5' and 3' ends of single-stranded molecules are intramolecularly ligated to form closed loops that can be subjected to rolling-circle amplification followed by nanopore sequencing. The resulting reads comprise a long string of linked sense or antisense strands akin to the results of the CircSeq technique but with many more copies of much longer fragments⁹² (FIG. 3a). Although better than simple nanopore sequencing, the very low raw nanopore accuracy coupled with amplification errors leads to a final consensus error rate that is still inferior to the best short-read platforms.

A unique feature of single-molecule sequencers is that consensus sequencing can be achieved without any amplification. For the Oxford Nanopore platform, use of a hairpin adapter to link the two strands of an individual DNA duplex is the simplest form of amplification-free consensus sequencing, which requires neither an SDE nor a UMI⁹³ (FIG. 3b). The incremental error correction

achieved by sequencing both strands of a duplex (2D) as opposed to only one strand (1D) is still dwarfed by the substantial baseline nanopore error rate. In the future, it should be possible for the same linked pair of molecules to be repeatedly passed back and forth through a nanopore for more rigorous consensus building⁸⁸.

The first consensus sequencing method to verifiably correct errors using both strands of an individual DNA molecule was the circular consensus sequencing (CCS) SMRTbell technique on the PacBio platform⁹⁴. This approach relies on ligation of hairpin-shaped adapters to either end of a double-stranded template to form a closed loop, which is directly sequenced and produces multiple passes of data from each strand without prior amplification (FIG. 3c). CCS has been applied to high-accuracy sequencing of error-prone repetitive sequences⁹⁵, non-invasive detection of low-frequency cancer-derived mutations⁹⁶, metagenomic deconvolution⁹⁷ and direct measurement of DNA polymerase error rates^{38,98}. Under optimal conditions, the method can achieve accuracies of the order of 10^{-7} because jackpot artefacts from one strand are unlikely to occur as complementary changes on the other. However, because of lower raw accuracy, a dozen or more sequencer passes across the tandem copies, as well as treatment with DNA repair enzymes, are needed to achieve the maximum resolution³⁸. The lower nucleotide output of the PacBio system as compared with more standard short-read platforms means that ultra-rare variant detection is more challenging from a cost and time perspective.

Other emerging single-molecule technologies involving direct electrical base detection on microchips⁹⁹ may eventually allow similar opportunities. A recently announced single-molecule sequencing-by-hybridization platform repeatedly interrogates individual bases by iterative hybridization with different overlapping probes, which is effectively another embodiment of consensus sequencing¹⁰⁰; the system should allow complete duplex error correction using UMIs or physical linkage of complementary DNA strands. With further developed consensus-based error-correction approaches, the unique benefits of all these newest-generation platforms, namely, very long reads, rapid library preparations and, in some cases, easy portability¹⁰¹, can be meaningfully realized in applications to detect low-frequency variants.

Considerations in choice of consensus method

Trade-offs between accuracy, cost, recovery, speed and read length mean that no single consensus-based error-correction method is optimal for every application. Because most sequencers in use are short-read systems, techniques designed for these platforms have generally been favoured. Duplex approaches are by far the most accurate but also generally the most expensive per error-corrected base because of the need to sequence more duplicates to have a reasonable probability of recovering copies of both strands. Put another way, the consensus-making efficiency is lower because a greater raw sequencing depth is required to obtain a similar molecular (consensus) depth than with single-stranded methods. Because the sensitivity for detecting a rare variant

Consensus-making efficiency

The number of raw sequencing reads that are required to form a consensus read. This typically refers to an average: total raw reads divided by total consensus reads.

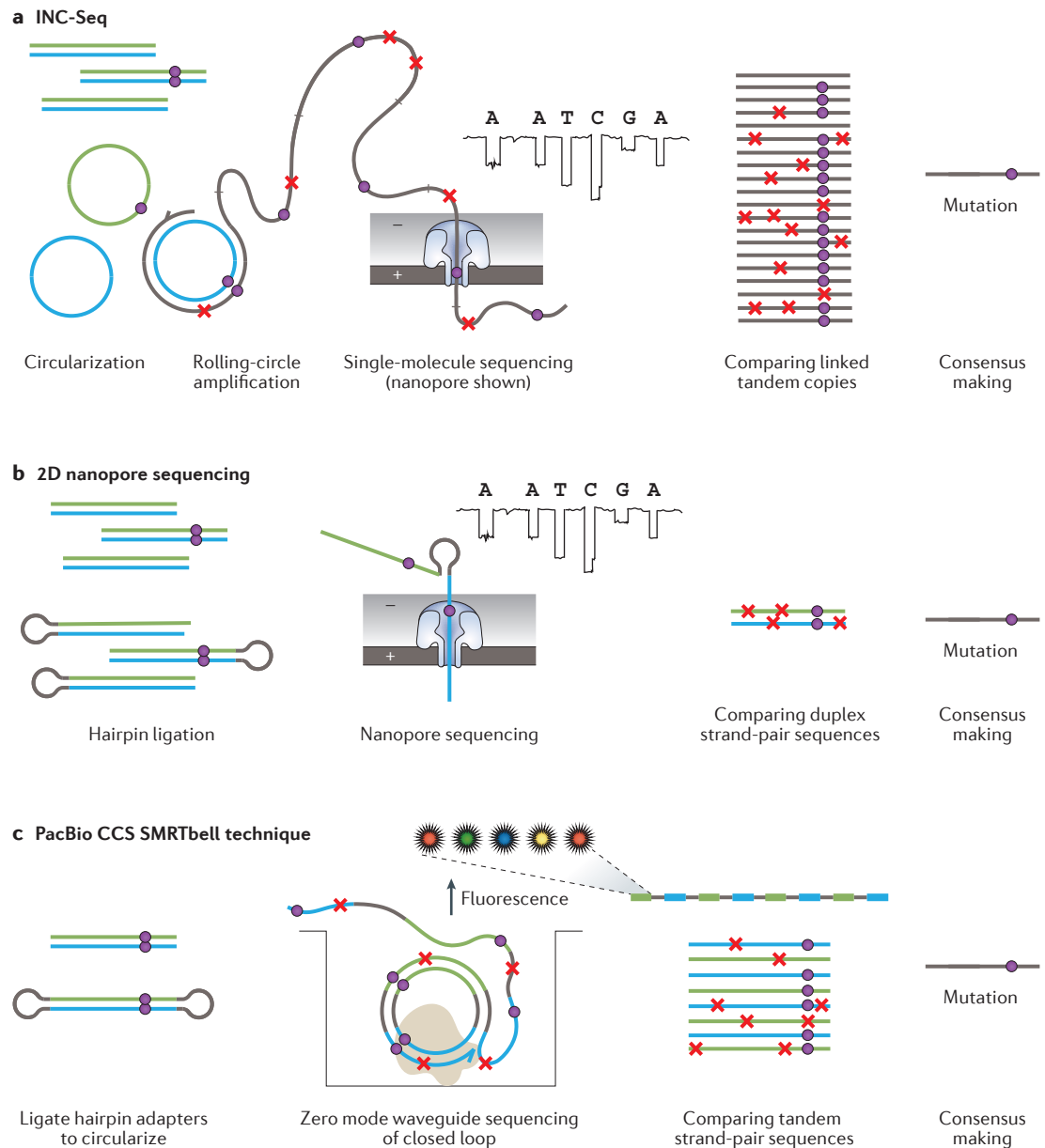


Figure 3 | Methods of single-molecule sequencing consensus-based error correction. a | The intramolecular-ligated nanopore consensus sequencing (INC-Seq) method begins by circularizing single-stranded DNA fragments followed by rolling-circle amplification of the loop; each product is a long DNA strand comprising concatenated copies of one of the strands of the starting molecule. This is sequenced on a long-read platform. True mutations (circles) can be discerned from most sequencing or amplification errors (crosses). **b** | 2D nanopore sequencing involves ligation of a hairpin adapter to one end of a duplex DNA molecule followed by tandem nanopore sequencing of the linked original strands. **c** | The circular consensus sequencing (CCS) SMRTbell technique entails ligation of hairpin adapters to each end of a molecule, followed by direct sequencing of the closed loop on the long-read Pacific Biosciences (PacBio) platform. Both strands are sequenced together in multiple passes. In the latter two cases, consensus sequences incorporate data from both DNA strands.

is influenced by both error rate and the number of consensus families generated at a given locus, what duplex methods gain in error rates comes at a greater cost for attaining sufficient molecular depth. For small genomic targets, this trade-off is acceptable but may become an important consideration with large multi-gene panels.

One substantial source of consensus-making inefficiency is non-uniformity of the amplification that is used

to generate molecular copies. With duplex techniques, if one strand replicates better than the other, this may prevent a consensus from forming or require an inordinately large number of raw reads to achieve a consensus. With single-stranded methods, if one locus amplifies better than another, a similar problem arises. Careful attention to factors that bias PCR efficiency, such as fragment length, help to abrogate the issue. For example, in

a preliminary report, we recently described the use of targeted CRISPR–Cas9 digestion to generate DupSeq libraries of uniform length and further capitalized on this size specificity to enrich for genomic loci of interest, thus further reducing competition during PCR⁶². Together, these features can improve consensus-making efficiency by an order of magnitude. A clever approach to eliminate amplicon competition and normalize yield per founding molecule using digital emulsion amplification has been demonstrated with both PCR and rolling-circle amplification^{71,77,102}. Ultimately, the most efficient consensus sequencing approaches are those that do not require any amplification, but single-molecule technologies will need to improve considerably before this becomes a major consideration.

Another important factor when selecting a method is the molecular conversion efficiency — that is, the fraction of input DNA molecules that are recovered as consensus sequences. This is typically lower with duplex than with single-strand approaches because both halves of a molecule must successfully amplify, which may be impossible if one is damaged or missing. Amplicon-based library preparation methods offer higher recovery and more rapid workflows than ligation followed by hybridization capture methods but do not retain complete duplex information. Simplicity and speed are important in clinical and some commercial settings. In situations where available DNA is limited, for example, in forensics or liquid biopsy applications, maximum recovery is important to detect low-frequency variants. However, greater recovery does not necessarily portend higher sensitivity if substantially more errors are introduced. FIGURE 4 illustrates the relationship between sequencing accuracy and positive predictive value for standard NGS, single-strand consensus sequencing and DupSeq. Although single-strand consensus methods have an absolute accuracy of the order of 10^{-5} , when attempting to detect variants that are present at a frequency of 1 in 100,000, approximately 80% of called mutations will be errors. Higher-accuracy duplex methods are necessary to reliably call mutations at this level. Increasingly sophisticated computational techniques are being developed to statistically integrate information from different types of consensus sequences and proportionally weight the certainty of called variants as a way to maximize data recovery while also retaining accuracy^{103,104}. In addition, emerging hybrid biochemical methods that combine the benefits of PCR-based targeting on library preparation speed and conversion efficiency with the advantages of adapter-based molecular tagging are likely to further narrow current performance trade-offs^{105,106}.

An important caveat is that with extreme accuracy comes new challenges from non-sequencing sources of errors. Certain artefacts that may be negligibly uncommon with standard NGS become significant when the sequencing background is reduced. For example, rare mapping errors of closely related pseudogenes or low-level cross contamination between samples during tissue processing can artificially appear as low-frequency variants. Slight carry-over between sequencer runs, amplification chimaeras or clustering errors that leads to index

shuffling^{37,51} can falsely make a clonal variant from one sample appear as a low-frequency mutation in another. Furthermore, when the technical background is reduced, newly apparent rare variants resulting from one process can easily be mis-assumed to be the result of another. For example, mutations commonly found in cancers can be detected at very low levels as part of normal ageing in healthy individuals and be a source of false positives in sensitive diagnostic tests¹⁰⁷. High-accuracy sequencing is a powerful tool, but it necessitates particularly thorough controls and well-thought-out experimental designs and interpretations.

As a final note, we wish to emphasize that the above descriptions and the illustrations in FIGS. 2,3 are highly simplified representations of complex techniques and do not fully convey many of the informatic subtleties of analysis. For a better appreciation, we refer interested readers to a select list of relevant software packages at the end of this article and those included as supplementary information in cited papers.

Applications of rare variant detection

The history of innovations in NGS and technologies for improving its accuracy are best understood in the

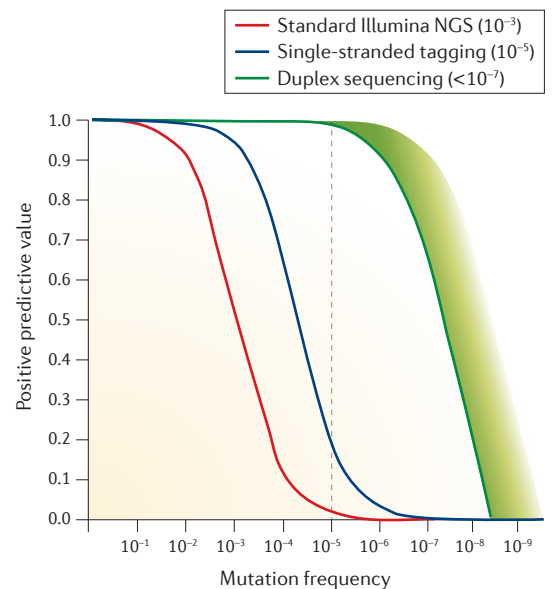


Figure 4 | Impact of error correction technology on detection sensitivity. The positive predicted value (the expected number of correct positive calls divided by the total number of positive calls) is plotted as a function of the variant allele frequency in a molecular population for each sequencing method of a specified error rate. As seen by curve overlap, nearly all mutant calls will be correct using any method if the frequency of detected variants is greater than 1 per 10. However, the error rates of standard Illumina sequencing and single-stranded tag-based error correction result in critical losses in positive predictive value at variant frequencies of ~1 per 100 and 1 per 1,000, respectively. The extremely low error rate conferred by duplex sequencing enables confident identification of variants below 1 per 100,000 (dotted line). NGS, next-generation sequencing.

Molecular conversion efficiency

The fraction of inputted DNA molecules of interest that are recovered as consensus sequences. This is often described in terms of genome-equivalents.

context of scientific questions they have been developed to answer. In this section, we highlight major fields in which the detection of minority genetic variants is important for medicine, biology and industry and provide examples of where the different approaches described above have been (or could be) applied (FIG. 5).

Cancer. Cancer is the ultimate disease of genetic heterogeneity^{3,108}. In the past decade, how it arises, progresses and spreads in the context of somatic evolution has been increasingly recognized^{109–113}. The cells in a tumour are not uniform. Mutations arising during cell division under the influence of selective pressures, as well as random drift, can lead to the outgrowth of genetically divergent clones in spatially distinct areas of a primary tumour^{114–117} and derivative metastases^{118,119}. Minority clones, which may be present at frequencies below the detection limits of conventional NGS techniques, can both drive tumour growth¹²⁰ and be an important source of resistance to therapy and basis for relapse^{5,121–123}; in fact, a higher degree of subclonal heterogeneity portends a worse prognosis in multiple tumour types^{124–126}.

A particular challenge has been developing technologies that are robust enough for resolving heterogeneity in the clinical setting. Pre-existing subclonal drug-resistance mutations in blood cancers have implications for the choice of initial therapy^{127,128}. After treatment, detection of rare cells with leukaemia-associated mutations using consensus sequencing indicates minimal residual disease (MRD) and the need for further treatment¹²⁹. In solid tumours, subclonal drug-resistance mutations are similarly relevant but may be missed by physical biopsy¹³⁰. An intense area of research is the use of liquid biopsies to non-invasively genotype cell-free DNA (cfDNA) shed by tumours into plasma, which allows genetic sampling of more than one region of a cancer in a way that can be readily repeated as a tumour evolves^{104,131–133}. Liquid biopsies are being used to detect both drug-sensitizing and drug-resistance mutations¹³⁴, low-level MRD after surgery^{135,136} and to follow the response to treatment¹³⁷. Tumour DNA can be found at low abundance in many body fluids^{138–142}. In each case, consensus sequencing-based error-correction approaches (single-strand, duplex or both) have been demonstrated to improve the detection of rare subclonal mutants. The detection of mutations in a variety of body fluids has been used to screen for specific cancer types in asymptomatic individuals for some time^{140,141,143}, but an even more ambitious prospect is the possibility of a universal pan-cancer-screening blood test¹⁴⁴. To achieve the exceptionally low false-positive rate needed for use in healthy populations, especially given the breadth of the genome that must be examined at high depth to identify very early tumours of many types, near-perfect technical accuracy is required using the most robust error-corrected sequencing methods possible.

Ageing. Mutations occur with each cell division; thus, it is no surprise that they increase in number with age⁸. A long-standing and unanswered question is whether these mutations reflect a cause or effect of ageing;

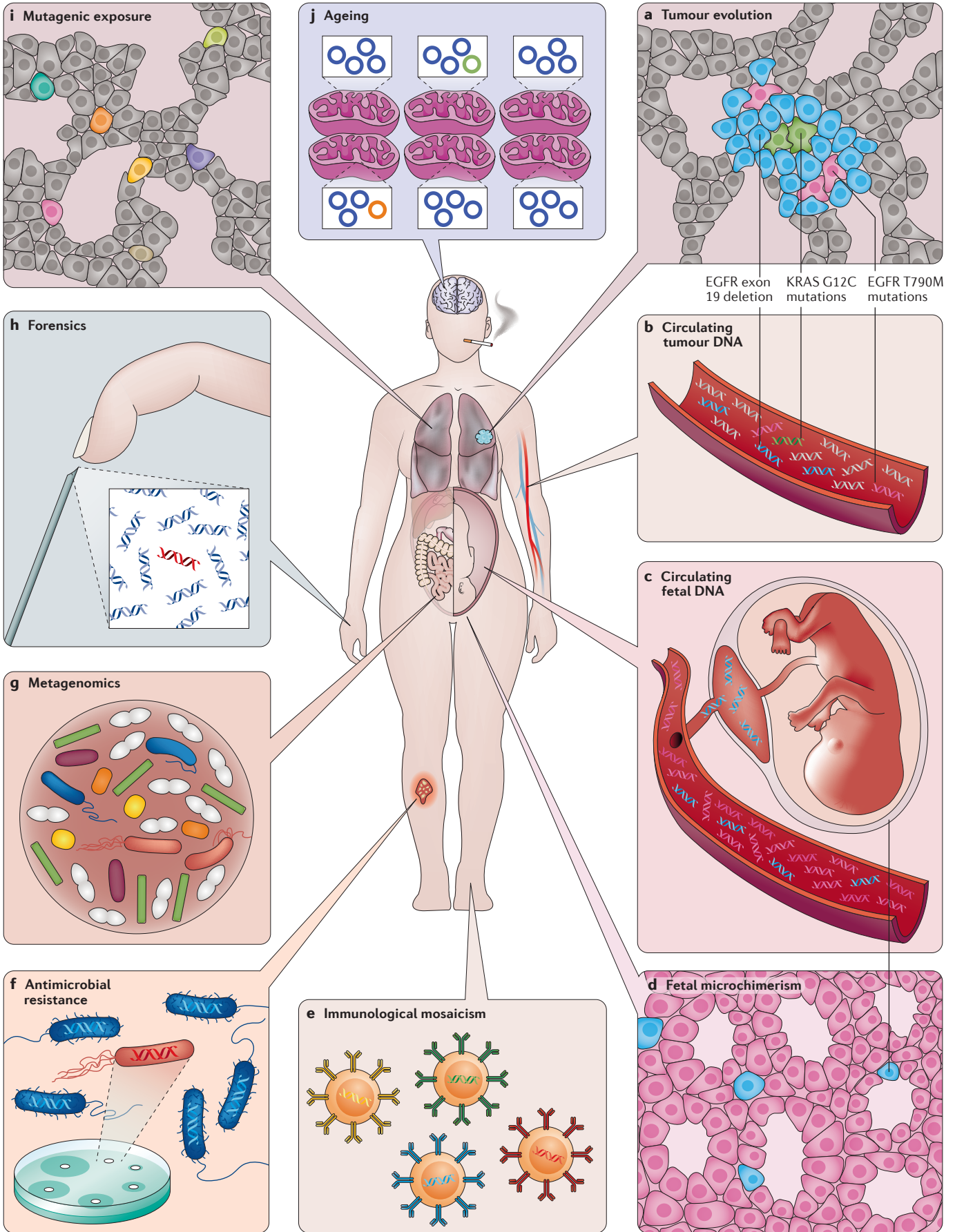
regardless, the association that has been found using duplex sequencing approaches is strong^{61,80}. Subclonal mutations have been associated with the onset of age-associated pathologies, such as neurodegeneration^{7,82}. The greatest risk factor for cancer, a disease caused by mutations, is ageing¹⁴⁵. Studies using conventional NGS have found clones bearing leukaemia-associated mutations in the blood of a subset of healthy individuals at a size and frequency that increases with age^{146,147}. Higher-accuracy single-strand consensus sequencing methods have identified these mutations at lower frequencies in nearly all adults¹⁴⁸. Using ultra-high-accuracy duplex consensus sequencing approaches, low-frequency age-associated mutations have been directly measured in multiple human tissues⁶¹, many of which are common to cancers^{149,150}. This highlights both the novel discoveries that come with greater accuracy and the new challenges the knowledge brings, for example, in mutation-based cancer screening¹⁰⁷. Whether age-associated subclonal mutation patterns will be able to predict future cancer risk or longevity remains to be explored.

Mutagenesis. Our bodies are exposed to endogenous and exogenous mutagens throughout life. It has long been known, using artificial selection-based assays, that mutations can be induced by genotoxic chemicals but only recently has it been possible to detect these directly^{60,61}. Instead of being limited to reporter genes, the highest accuracy duplex sequencing approaches can evaluate mutagenesis at any genomic site of any organism and have the potential to become a new standard of genotoxicity evaluation^{85,102,151–153}. New methods of therapeutic genome modification can lead to rare off-target mutagenesis through mechanisms that are not easily recapitulated in model organisms¹⁵⁴. The same holds true for treatments that are not intended to modify the genome but with the theoretical potential for mutagenesis¹⁵⁵. Emerging cellular therapies where stem cells are harvested and propagated ex vivo can introduce substantial numbers of mutations²¹. Given the rapid emergence of new medical technologies, it is important that we have equally powerful tools to carefully monitor their genetic consequences.

Maternal–fetal biology. Pregnancy is a state of chimerism: two genomes inhabit the same body and intermix. Even after birth, a mother and child remain genetically intertwined. Extremely rare populations of fetal cells can persist in a woman for decades¹⁵⁶. This interesting state of prolonged immunotolerance may play a contributory role in autoimmune disease, miscarriages and pre-eclampsia but also appears to confer certain benefits, particularly cancer protection. Until recently, genetic techniques for studying these rare populations of cells have been inadequate¹⁵⁷. During pregnancy itself, apoptosis of placental cells releases fetal DNA into the maternal circulation where it can be collected for non-invasive prenatal testing (NIPT). Fetal aneuploidies can be clinically detected by NGS with relative technical ease by simply counting the relative proportions of different chromosomes, even though fetal DNA is in the minority^{158,159}. In fact, one

Aneuploidies

Abnormal numbers of chromosomes in a cell. This may be inherited, such as trisomy 21, the basis of Down syndrome, or somatically acquired, such as in cancer.



◀ **Figure 5 | Applications of rare variant detection.** **a** | Cancer evolution. Genetic heterogeneity within tumours is thought to be responsible for the emergence of therapeutic resistance. In lung adenocarcinomas with certain epidermal growth factor receptor (EGFR) mutations under treatment with targeted inhibitors, drug-resistance mutations arise at low levels then clonally expand. **b** | Cell-free tumour DNA. Tumour cells release fragments of DNA into plasma and other body fluids that can be sampled via liquid biopsy. This serves as a non-invasive means of determining the genetic makeup of a tumour without a physical biopsy and is a sensitive way to detect minimal residual disease and early relapse. **c** | Circulating fetal DNA. Placental-derived DNA in the maternal circulation can be used to non-invasively detect fetal genetic traits or abnormalities. **d** | Fetal microchimerism. Fetal cells that engraft into a mother may persist many years after birth. These have important immunological consequences. **e** | Immunological mosaicism. Somatic V(D)J recombination and hypermutation in B cells and T cells create heterogeneity that helps the body adapt defences to new infectious and neoplastic threats. **f** | Antimicrobial resistance. Low-frequency variants in single-cell organism populations can be responsible for drug-resistance outbreaks. **g** | Metagenomics. Complex mixtures of microorganisms exist throughout the living world. The human body is colonized with symbiotic microorganisms and, in some diseases, health problems can arise from disrupted microbial diversity. **h** | Forensics. Mixtures of human tissues are routinely recovered at crime scenes or natural disasters. In some scenarios, the abundance of DNA from one individual may be much greater than that of another. **i** | Mutagenic exposure. DNA damage can be caused by normal ageing and by carcinogens. Very-low-frequency mutation load may be proportional to future cancer risk. **j** | Ageing. DNA damage occurs throughout life from exogenous and endogenous processes. Low-frequency mutations in both the nuclear and mitochondrial genome (the latter is shown here) may play a role in certain age-related pathologies in addition to cancer, such as neurodegeneration and autoimmunity. Subclonal mutations might serve as a biomarker of disease risk or even longevity.

early indication of the feasibility of DNA-based cancer screening was the incidental detection of maternal cancers from non-fetal chromosomal imbalances found during NIPT¹⁶⁰. Complete fetal genomes have been assembled from sequencing of cfDNA in maternal blood based on paternally inherited single nucleotide polymorphisms (SNPs), albeit using low-accuracy conventional NGS. To become viable as a clinical means of prenatally detecting de novo point mutations, ultra-accurate methods will need to be employed. This is particularly true of disease-causing mutations that arise mosaically during embryogenesis^{161,162}. A distinct but related application of identifying foreign SNPs in cfDNA is the early detection of organ transplant rejection¹⁶³.

The immune system. An individual's immune system is finely tuned for generating genetic heterogeneity through sanctioned V(D)J recombination and somatic hypermutation; a single blood sample may contain hundreds of thousands of different T cell receptor and immunoglobulin sequences¹⁶⁴. This genetic profile changes in response to infections, immunization and age, among many other states^{165,166}. The adaptive immune system plays a role in defence against neoplasia, and the genetic pattern of tumour-infiltrating lymphocytes (TILs) can reflect both prognosis and the likelihood of response to therapy^{166,167}. On the other end of the immune spectrum is overactivity. It has been theorized that some autoimmune disease may develop in response to subclonal somatic mutations in non-immune tissues¹⁶⁸. Chronic stimulation of the immune system also causes low-frequency off-target somatic hypermutation in lymphocytes themselves, leading to lymphoma¹⁶⁹. In each category, high-accuracy consensus sequencing methods

of multiple varieties are proving critical for disentangling the immunological heterogeneity of both physiological and pathological states.

Microbial populations. Our bodies contain at least as many microorganisms as human cells. Although some can cause disease, a greater number serve symbiotic functions, such as building or metabolizing molecules, outcompeting pathogenic organisms and training our immune systems^{170,171}. Colonization begins even before birth, and the stability of the ecosystem that develops influences health processes, including allergy development and autoimmune reactions¹⁷², changes in body weight¹⁷³ and responses to medications¹⁷⁴, among others. Early successes with therapeutic microbiota transplantation further indicates the importance of these populations^{175,176}.

The study of the composition of metagenomics ensembles with traditional techniques is not trivial. Some organisms grow poorly in culture or not at all. Viruses, fungi, protozoa and highly divergent phyla of bacteria that coexist in vivo often require unique growth conditions in vitro. In the case of a suspected infection, being able to quickly identify the causative agent or agents and predict which therapies will be most efficacious is clinically important but not always possible. Whereas clinical decisions about antimicrobials often must be made in minutes or hours, traditional culture takes hours or days and, in some cases, even weeks¹⁷⁷. NGS has emerged as a powerful and universal means of characterizing pan-kingdom microbial populations¹⁷⁸. High-speed NGS workflows can now identify organisms causing sepsis and predict their drug sensitivity in a matter of hours^{179,180}.

However, speed is not the only challenge in clinical microbial sequencing. When the genomes of organisms in a polymicrobial sample are very distinct, standard NGS can readily identify minority populations. When a population's constituents are genetically similar, however, sequencing errors prevent variant detection below approximately 1%. Because the evolutionary success of many pathogens is predicated on rapid evolvability to circumvent host immune responses, many mutate readily to form heterogeneous populations of closely related, but genetically distinct members, collectively known as a quasispecies^{181,182}. Deep sequencing has shown that the emergence of low-level drug-resistance mutations in multiple types of infections including HIV^{4,13,183}, viral hepatitis^{184,185} and tuberculosis¹⁸⁶ can predict therapeutic failure. As in oncology, consensus-based NGS error-correction approaches enable detection of rarer drug-resistance mutations⁵⁶, potentially affording the opportunity for earlier interventions if found. Perhaps just as importantly, the ability to confidently recognize the absence of such mutations might allow one to avoid unnecessary use of broad-spectrum agents.

Other applications. The utility of deconvolving heterogeneous, closely related mixtures of nucleic acids extends beyond medicine. Of the innumerable free-living microorganisms in the natural world, the majority have never been isolated or cultured¹⁸⁷. Direct high-accuracy NGS

Metagenomics

The study of complex microbial populations encompassing many co-mingling species that form an ecosystem, for example, an individual's gut microbiota.

Box 1 | The future of DNA sequencing: complete genomics and multi-omics

Being able to resolve heterogeneity among an aggregate collection of molecules is, itself, not sufficient to deeply understand evolving populations. Mutations do not act in isolation: the phenotype they confer often depends on the sequence of other genes and regulatory elements sharing the same cell. Methods for high-throughput identical barcoding of all the molecules from single cells using flow sorting or droplet compartmentalization are becoming widespread for both DNA and RNA^{199,200}. Use of endogenous cell membranes as individual barcoding reaction chambers is another high-throughput means of single-cell molecular labelling²⁰¹. We anticipate that the combination of high-throughput compartmentalization methods with ultra-long-read single-molecule sequencing technologies that incorporate duplex consensus-based error correction will enable truly whole-genome sequencing of large populations of single cells in the foreseeable future.

Single-cell transcriptomics^{200,201} and proteomics²⁰² can be achieved using high-throughput sequencing technologies. DNA carries important epigenetic information beyond its primary sequence in the form of methylation, other non-canonical nucleotides, chromatin structure and nuclear colocalization patterns²⁰³. In RNA, more than 100 types of naturally occurring chemical modifications have been described²⁰⁴. All these plus other emerging ‘omic’ technologies have been applied to single cells but have yet to be meaningfully combined. Amassing this parallel data from individual cells while retaining information about their relative spatial relationships in their native 3D tissue context will be a Herculean undertaking. Perhaps the greatest challenge to consider is simultaneously capturing the fourth dimension: nearly all current genomic methods represent finite snapshots in time, rather than a temporally dynamic measurement. Whether we will ever be able to non-disruptively determine the sequence of the complete in vivo genomes and epigenomes of billions of intact cells in a living organism is probably a question for our children or perhaps our children’s children.

Phasing

The proper assignment of two or more variants at spatially distant genomic locations to the derivative nucleic acid molecule, for example, the maternal or paternal allele.

of environmental samples is an important tool for identifying novel protein variants with potential industrial applications¹⁸⁸, for monitoring ecological health¹⁸⁹ and for facilitating food safety testing, including the detection of low-frequency drug-resistant microbial strains¹⁹⁰. In forensics, the ability to confidently distinguish mixtures of DNA contributed from different individuals is critical, given the consequence of mistakes^{106,191}. High-accuracy sequencing methods are especially important when heterogeneous mixed samples are degraded and therefore more error prone, as may occur at both modern crime scenes and with ancient DNA from archaeological sites¹⁹².

Conclusions

Within the span of a single human lifetime, we have gone from recognizing DNA as genetic material¹⁹³ to sequencing the human genome¹⁹⁴. Despite this remarkable progress, many opportunities remain. For example, the structure of certain highly repetitive regions in many organisms is still unknown. In fact, for this reason, we have yet to assemble even one genuinely complete human genome. Both emerging long-read technologies and techniques for phasing and accurately assembling short-read data using fragment barcoding methods^{72,195} or DNA crosslinking techniques¹⁹⁶ are bringing us closer, but they are not yet accurate enough to reliably detect very-low-frequency subclonal heterogeneity of structural variation despite the clinical importance in both the somatic and neoplastic settings^{197,198}. Another substantial challenge on the horizon is combining ultra-accurate single-cell genomic information with other types of ‘omic’ technologies and contextualizing with higher-order topological relationships between individual cells within a tissue (BOX 1).

We began this Review by enumerating NGS technologies that have had a major impact on the ability to detect subclonal variants and discussed the many fields in which NGS has benefitted or will soon benefit from this high accuracy. Biological and clinical questions that were previously intractable can now be approached. A meaningful vision of the future should not only focus on improvements to established areas but also address the role of high-accuracy DNA sequencing in delineating new and important hypotheses. Sequencing technology is advancing more quickly than ever before. Just as evolution is the driving force behind the progression of all life on Earth, so too is it with technology: the popularity of new sequencing methods will continue to rise and fall, but the discoveries that come from each will remain immortal. It is important to remember that methods are merely tools, not themselves answers; it is on the creative and responsible applications of these new technologies where we should focus our greatest attention.

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Author contributions

J.J.S and L.A.L. contributed to discussion of content and reviewing and editing the manuscript before submission. J.J.S. was primarily responsible for researching data and writing the manuscript. M.W.S. researched the literature and contributed to writing parts of the initial drafts of the article.

Competing interests

J.J.S., M.W.S. and L.A.L. are equity holders in TwinStrand Biosciences, Inc.

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