A vision for ubiquitous sequencing

Yaniv Erlich1,2

1Department of Computer Science, Columbia University, New York, New York 10027, USA; 2New York Genome Center, New York, New York 10013, USA

Genomics has recently celebrated reaching the $1000 genome milestone, making affordable DNA sequencing a reality. With this goal successfully completed, the next goal of the sequencing revolution can be sequencing sensors—miniaturized sequencing devices that are manufactured for real-time applications and deployed in large quantities at low costs. The first part of this manuscript envisions applications that will benefit from moving the sequencers to the samples in a range of domains. In the second part, the manuscript outlines the critical barriers that need to be addressed in order to reach the goal of ubiquitous sequencing sensors.

The cost of DNA sequencing has plunged orders of magnitude in the last 25 years. Back in 1990, sequencing 1 million nucleotides cost the equivalent of 15 tons of gold (adjusted to 1990 price). At that time, this amount of material was equivalent to the output of all United States gold mines combined over two weeks. Fast-forwarding to the present, sequencing 1 million nucleotides is equivalent to the value of ~30 g of aluminum. This is approximately the amount of material needed to wrap five fast breakfast sandwiches at a New York City food cart. As a result of the breathtaking pace of advancements, DNA sequencing has become the ultimate back-end for a wide spectrum of biological assays (Shendure and Aiden 2012). A growing number of these assays simply take advantage of DNA or RNA for labeling, creating molecular post-its that can be read in a highly parallel and cost-effective manner (Zador et al. 2012; Levy et al. 2015). Paraphrasing the famous quote by James C. Maxwell (1864), the aim of many experimental techniques is to reduce the problems of nature to the determination of DNA sequences.

So, what is the next frontier of DNA sequencing? For decades, an affordable sequencing platform—the $1000 genome—has been the main focus (Green et al. 2011). While price was under strong selective pressure, the community has largely accepted sequencing devices in any shape and size. Examples of such devices include the now obsolete Heliscope by Helicos Biosciences, which contained a >100 kg granite slab to stabilize the sequencer (Davies 2010) and the 860 kg Pacific Biosciences’ RSII instrument, with its large footprint (Fig. 1A). In stark contrast, the last year has witnessed the emergence of small-footprint sequencers with the successful early access program of the handheld Oxford Nanopore MinION (Fig. 1B; Loman and Watson 2015) and an ongoing development of relatively small sequencers by other companies such as Genapsys (Fig. 1C).

With these exciting new developments, the next phase of the sequencing revolution is the emergence of DNA sequencing sensors. Different from massive sequencing platforms such as the Illumina X Ten and Pacific Biosciences RSII, sequencing sensors will be extremely miniaturized devices that include automatic sample preparation with the aim of real-time sequencing in the field (Table 1). Integrating these miniature sequencers in larger systems will add a DNA-awareness layer to various devices (Fig. 1D). That being said, sequencing sensors will not replace sequencing platforms in the foreseeable future; the latter will certainly be around for heavy-lifting tasks where scale matters, such as whole-genome sequencing. Rather, sequencing sensors will enable a new range of applications that can substantially benefit from moving the sequencers to the samples instead of the traditional way of moving the samples to the sequencers. With this goal in mind, the cost per base pair (bp), extremely high accuracy, and other traditional parameters of sequencing performance will be of secondary importance. More attention will be devoted to the cost of the device, seamless sample preparation, latency of sequencing results, and environmental robustness. Here, I envision potential applications of sequencing sensors in different domains, map chief engineering challenges, and briefly discuss potential regulatory issues of ubiquitous sequencing.

Potential applications of sequencing sensors

Sequencing at home

Sequencing sensors will enable the advent of DNA-aware home appliances, continuing the growing trend of smart devices that monitor the household environment such as Nest Protect that senses CO and smoke levels. Multiple appliances could benefit from integration with sequencing sensors, including air conditioning or the main water supply to monitor harmful pathogens. However, of all possible options, toilets may offer the best integration point. First, toilets regularly collect a large amount of biological material as part of daily life. There is no need for new routines or special procedures, solving a major usability barrier. In addition, it is also possible to monitor pathogens in the water supply. Second, toilet integration can solve certain engineering complications, such as access to fresh water and a place to drain sequencing waste (although access to electricity may be complicated). Most toilets are designed to have space below the water tank, which is rarely occupied. This could be a convenient area to place a small system without substantial miniaturization burden. Third, as opposed to most appliances such as refrigerators or dishwashers, many houses have more than one toilet, increasing the potential market size for such a device. A smart toilet system can convey extremely rich information about household members for medical and dietary purposes.
It will allow monitoring of fluctuations in the host microbiome and other potential indicators of the well-being of the gastrointestinal and renal systems. Since waste also contains host DNA, it will be possible to seamlessly assign the sequencing results to a specific individual without additional user involvement. In addition, it might be possible to track food composition from waste in order to automatically log dietary habits.

Another exciting opportunity is bringing sequencing sensors to the hands of the general public as a stand-alone device. With the growing interest in self-tracking, an affordable sequencing sensor with automatic sample preparation has the potential to be a trendy—and admittedly quite geeky—technological accessory.

Just a year ago, ~13,000 Kickstarter backers pledged over $2 million to develop SCiO, a low-cost handheld spectrometer that is intended to sense the chemical makeup of food and drugs, highlighting the potential market for similar DNA-aware devices. Beyond the educational advantages, bringing sequencing sensors to the masses can also assist in crowd-sourcing DNA or RNA signatures from various sources. This can facilitate the development of interpretation apps in domains that have received relatively little attention from traditional scientific endeavors such as collecting microbiome signatures at various spoiling stages of food or detecting mislabeled food products.

Health-related applications

Real-time DNA sequencing has remarkable potential for diagnosing infectious diseases by bringing the sequencers to at-risk individuals. For example, rapid sequencing at airport checkpoints might be useful to control pathogen outbreaks and offer medical assistance to affected passengers. Similarly, a portable sequencer will enable physicians to provide more accurate diagnoses in the field during humanitarian crises or in the clinic without the need to waste time by sending samples to a lab.

A more bold approach is to leverage the at-home sequencing devices presented above. By careful development of at-home tests, sequencing sensors will allow patients to obtain data on infectious disease in the comfort of their own homes, reducing both the misery of getting out of bed while sick and the risk of infecting others. Specialized apps will integrate the sequence reads with other physiological measures and patient complaints. A virtual physician visit (or an FDA-approved fully automated program) will conclude the diagnosis and decide on the course of treatment specific to the pathogen.

Sequencing sensors can also allow pathogen surveillance by constant monitoring of key junctions for disease-spreading routes such as hospital laundry (Fijan and Turk 2012), hotel water systems (Mouchtouri et al. 2007), central air conditioning systems (Tringe et al. 2008), and municipal sewage (King 2014). Multiple studies have highlighted the utility of DNA sequencing for disease surveillance including the 2011 outbreak of E. coli O104:H4 in Germany (Rohde et al. 2011) and the 2014 Ebola pandemic originating in West Africa (Gire et al. 2014). These efforts played a key role in identifying the pathogenic strain, tracing its evolution, and reconstructing the spread of disease. However, it should be noted that pathogen surveillance from environmental samples before an outbreak will be far more challenging than these retrospective analyses that had the advantage of diagnosed patients. Recent studies have shown that, under certain conditions, species annotation pipelines can report questionable results such as the presence of endemic species in samples that were collected far away from their natural habitats or the detection of nearly extinct pathogens (Gonzalez et al. 2014; Afshinnekoo et al. 2015; Mason 2015). These errors stem from reads with local sequence similarity to the genomes of unrelated organisms or reflect horizontal gene

---

**Table 1. Sequencing platforms vs. sequencing sensors**

| Feature       | Sequencing platform | Sequencing sensor |
|---------------|---------------------|-------------------|
| Operation     | Static              | Mobile            |
| I/O           | Fancy touch screens | API               |
| Cost of sequencing | Less important | Important         |
| Cost per bp   | Important           | Less important    |
| Capacity      | Important           | Less important    |
| Speed         | Less important      | Important         |
| Accuracy      | Important           | Less important    |
| Bandwidth     | High                | Low               |
| Latency       | High                | Low               |
| Main application | Finding genetic variants | Quantifying DNA molecules |
| Made for      | Human               | Machines          |

---

Figure 1. Sizes of sequencing platforms vs. sequencing sensors. For other differences, see Table 1. (A) Three men haul an 860-kg Pacific Biosciences RSII, a sequencing platform, to the University of Exeter (photo courtesy of @PsyEpigenetics). (B) MinION sequencing sensor. (C) An early prototype of a Genapsys flowcell. The company develops an iPad-size sequencer. (D) A commodity digital camera chip ready for cell phone integration. Can DNA sequencers be that small?
transferred between species. In addition, the field has yet to establish a comprehensive knowledge of the genomic sequences of many microorganisms, increasing the likelihood of species misidentification. Addressing these challenges will necessitate long sequence reads for increased specificity, further maturation of analytical frameworks, and expanding databases of annotated species.

Additional potential usage for DNA sequencing sensors includes integration with medical devices for real-time monitoring of cell-free DNA and RNA in patients. These molecules are released into the plasma due to various biological processes, including cancer, conferring a diagnostic vantage point without invasive biopsies (Van Der Vaart and Pretorius 2008). Multiple lines of evidence have shown that these nucleic acids can be used as biomarkers for a series of acute conditions, such as liver response to drug overdose (Wang et al. 2009), heart attack (Creemers et al. 2012), sepsis (Dwivedi et al. 2012), and brain trauma (Ohayon et al. 2012). Real-time bedside sequencing at relatively short intervals can serve as the basis for a universal apparatus to capture these biomarkers, adding a new layer of information to patient status.

Forensic and security applications

DNA forensics for counter-terrorism and law enforcement applications can highly benefit from rapid sequencing sensors. Currently, the DNA evidence processing chain includes evidence collection, shipment of evidence to a local crime lab, isolation of DNA, generating a short tandem repeat profile using capillary electrophoresis, and a database search—a process that usually takes days to complete (Kayser and de Knijff 2012). Quick sequencing of DNA evidence at the crime scene can reveal the suspect’s identity in the critical window after the event to reduce the chance that the suspect will successfully flee, destroy evidence, or mount an additional attack. First responders of serious crimes will be able to operate the system to automatically notify other forces or border control of the suspect’s identity to better focus efforts to capture the suspect. Such a technology might also have military uses. Recent reports revealed that, at least in one instance, U.S. Special Forces used a DNA test for positive identification of a target of high interest (Whitlock and Gellman 2013). However, despite the importance of the target, it took >8 h before the assay could provide a final confirmation of the person’s identity.

Another application for rapid DNA-based sequencing is human identification at security check-points such as airports. While other biometric signatures such as fingerprints and physiometric properties are accurate and easy to obtain, DNA has the advantage of positive identification in the absence of a person’s inclusion in a database (Erlich and Narayanan 2014). Technically, this can be done by familial searches that rely on finding autosomal matches with first-degree relatives (Bieber et al. 2006) or by surname inference techniques that leverage Y Chromosome matches of distant relatives (Gymrek et al. 2013). With careful implementation that is sensitive to genetic privacy and cultural issues (Kim and Katsanis 2013), such technology at checkpoints could play a role in fighting human trafficking. This horrific worldwide problem affects millions of victims, most of whom are children or young females and takes a substantial societal and economical toll on certain regions (U.S. Department of State 2014). Family members of the abducted victim will contribute their own DNA to a specialized database that will only be used to locate relatives, similar to a missing person database. Security forces at airports or borders can check individuals suspected of being at risk. To prevent misuse, samples will be immediately destroyed and results will not be saved upon a negative match.

From a technical perspective, the identification of human DNA requires very low sequencing throughput. Currently, forensic DNA profiles are developed by genotyping 13–20 autosomal STR markers, which could be achieved with <4000 bp of total sequencing throughput (Ge et al. 2012). However, to avoid the lengthy target enrichment, it might be more advantageous to utilize an extremely low-throughput shotgun sequencing of the sample. Previous work has shown that any 200 common SNPs in linkage equilibrium create a unique profile for any person on earth (with the exception of monozygotic twins) (Lin et al. 2004). At least theoretically, a similar number of long (>1000 bp) reads could ascertain this number of SNPs. Practically, identification will probably require a few thousand reads to collect additional SNPs in order to obtain higher levels of confidence. The variant calling can be done together with aggressive genetic imputation which was shown to substantially improve the calling of common variants despite extremely low sequencing coverage (Pasaniuc et al. 2012). Finally, sample identification will rely on querying a forensic database of whole-genome sequencing or genome-wide array data.

Food industry

Rather than having consumers sequence food products at home, quality control can be integrated at various stages of the entire supply chain. Preliminary studies have shown that DNA sequencing can answer various food-related applications including distinguishing morphologically similar poisonous and edible mushrooms, detecting the accumulation of pathogenic microbes in meat, tracing the fermentation of cheeses, identifying hidden traces of allergens, and even predicting the ripening time of avocados (Dopico et al. 1993; Galimberti et al. 2013; Dugat-Bony et al. 2015; Wolfe and Dutton 2015). As such, sequencing sensors can be a highly flexible backend to monitor various issues in the food supply chain. The real challenge, however, will be to perform these assays outside of a laboratory setting.

Instead of analyzing naturally occurring sequences, ubiquitous sequencing in the food industry can utilize barcoding of food products with artificial DNA tags. Recent work has tested the usability of such tags in the food supply chain by delivering exogenous DNA fragments within a coat of silica, a common food additive, at an extremely low cost. It has been shown that these tags could be used to trace the source of milk in dairy products such as yogurt (Bloch et al. 2014) and serve as a long-term authenticity mark of premium olive oil (Puddu et al. 2014). Moving forward, artificial DNA labels can create an edible data structure that includes the type of food, producer, lot number, nutrient information, and presence of known allergens or simply encode a URL to a webpage that contains this information. The length of the DNA label can be relatively short, as recent DNA-specific encoding schemes have proposed methods to robustly code 1.6 bits of information in each DNA base pair (Goldman et al. 2013). DNA-labeled products can establish an inexpensive method to identify ingredients and sources of processed food, tracking authenticity, and revealing hidden traces of allergens. In addition, if it is possible to precisely quantify DNA labels from human waste, this scheme coupled with smart toilet systems (discussed above) will enable automatic logging of ingredient consumption. This can revolutionize the tedious manual process of documenting food intake, which is a key component of various diet programs and can help researchers gain substantial amounts of data.
correlating ingredient consumption and illness. These DNA tags could also label oral medications. Sequencing sensors at toilets could trace the presence of the drug in waste, creating a technology to monitor patient adherence regardless of the chemistry of the drug, addressing a major public health issue.

**Technological challenges**

Full realization of DNA sensors will require overcoming a multitude of barriers. Below is an outline of the main obstacles.

**Sample preparation**

Mainstream protocols for shotgun sequencing are not compatible with the aim of real-time miniaturized DNA sensors. They involve multiple chemical reactions and physical manipulations to purify DNA and tether the molecules to the sequencer apparatus, inducing a substantial latency and complicating miniaturization. Multiplexed target enrichment protocols, such as in-solution capture, are even more involved. They consist of more complex steps, including condensation of intermediate reactions, rapid cooling and heating, and a long (>24 h) hybridization step.

Recent advancements have the potential to greatly facilitate the advent of rapid and miniature library preparation methods. A new single-cell sequencing method has proposed a microfluidic-based technique to perform most of the RNA library preparation within droplets (Klein et al. 2015; Macosko et al. 2015). This procedure compresses a large number of processing stages into one step and creates a scalable method for in situ library preparation. GnuBIO, a Bio-Rad company, develops a microfluidic-based system for genomic library preparation that is completely integrated within their benchtop sequencer. If successful, this system will allow an end-to-end automated sequencing pipeline that starts with genomic DNA samples. Another alternative is to simplify library preparation by avoiding multiplexed target enrichment. Most of the applications mentioned above could either utilize shotgun sequencing or focus on a single region using a miniaturized PCR machine.

**Supply of reagents**

Low maintenance and environmental robustness are key issues for sequencing sensors. Current sequencing technologies require multiple types of reagents such as DNA polymerase (Illumina, Ion Torrent, and Pacific Biosciences, Genia), biological pores coupled to a motor enzyme (Oxford Nanopore), modified nucleotides (Illumina and Pacific Biosciences), and exogenous DNA primers (all technologies). These have relatively low durability and some also need to be stored at 4°C or −20°C. One possible solution to reduce the burden of loading reagents onto sequencing sensors is a user-friendly cartridge on which sensitive reagents can be lyophilized to allow storage at room temperature. A nonmutually exclusive effort is to minimize the types of reagents and prolong their durability. Previous studies have suggested that solid-state nanopores can offer increased robustness and durability over protein-based nanopores in lipid bilayers (Dekker 2007; Venkatesan and Bashir 2011). These solid-state nanopores might have the ability to sense naturally occurring DNA without any modification, which can greatly reduce the need for a reagent supply.

**Analytics**

An open question is how to perform the data analytics for sequencing sensors. Current sequencing platforms push low-level data from the machine and use external computers for almost all of the downstream data processing, from registering the sequencing signals via base-calling to alignment. Mobile sequencing sensors can wirelessly push raw data to the cloud or a local server as they sequence the samples. However, this will require data streams that are sufficiently compact to work with the bandwidth of mobile communication.

Latency is another factor of data analytics. Most sequence analysis algorithms are devised as offline programs that assume that all input data are available at the beginning of the run and that files are read and written at each stage (but see Chiang et al. 2014). Latency time could benefit from online bioinformatics pipelines that will align sequence reads as the flow of bases continues to arrive from the basecaller and make on-the-fly decisions before the full data are analyzed, such as pathogen detection from sufficient evidence. Unlike most data analytics today, sequencing sensor analytics should provide succinct and concrete information. The typical user will not care about quality recalibration or alignment of indels and will have zero knowledge or patience to deal with bioinformatics issues. All he wants are short answers such as “Meat is spoiled: Don’t eat” or “Suspect identified as Mr. X.” Simple and reliable reports will be key.

**ELSI aspects of ubiquitous sequencing**

The advent of ubiquitous sequencing can trigger a wide range of ethical, legal, and social implications (ELSI). The general public often considers DNA information as more sensitive compared to other types of personal data (Lunshof et al. 2008; Presidential Commission for the Study of Bioethical Issues 2012; Rodriguez et al. 2013). This fact is also reflected in certain laws that provide extra privacy protections for DNA information.

To highlight the complexities, consider the recent case of the “devious defecator” (Gilbert 2015). In this case, a grocery store company, Atlas, found that one or more of its employees defecated in one of its warehouses, ruining products. The company hired a forensic lab to obtain DNA from the feces and requested a DNA test from two employees thought to be responsible. The employees both sued Atlas since, according to the US Genetic Information Nondiscrimination Act (GINA), it is unlawful for employers “to request, require, or purchase genetic information with respect to an employee.” The court ruled in favor of the plaintiffs and awarded $2.25 million in damages although the genetic test was done on CODIS STR markers that do not reveal any health status. Most people are not aware of the challenges associated with analyzing human DNA samples, and the democratization of sequencing can expose them to legal liabilities.

Other laws also prohibit the collection of human DNA without consent. For example, in the UK, the Human Tissue Act requires consent for any collection of human DNA if there is intent to perform DNA analysis. The Israeli Genetic Information Nondiscrimination Act (GINA) is unlawful for employers “to request, require, or purchase genetic information with respect to an employee.” The court ruled in favor of the plaintiffs and awarded $2.25 million in damages although the genetic test was done on CODIS STR markers that do not reveal any health status. Most people are not aware of the challenges associated with analyzing human DNA samples, and the democratization of sequencing can expose them to legal liabilities.

Other laws also prohibit the collection of human DNA without consent. For example, in the UK, the Human Tissue Act requires consent for any collection of human DNA if there is intent to perform DNA analysis. The Israeli Genetic Information Nondiscrimination Act only authorizes genetic labs to perform tests on human DNA, with the exceptions of research and law enforcement purposes. In the US, abandoned DNA is not protected by the federal fourth amendment, and it is legally possible to collect and analyze samples from public places (Joh 2006), as was well demonstrated by a recent art project (Gambino 2013). However, certain states have imposed further regulation. For example, according to Section 79-L of the New York State Civil Rights Law, it is unlawful to “diagnose the presence” of genetic variations that are linked to human diseases without consent. A California senator recently proposed a more strict law similar to the UK law that will prohibit...
any collection of human DNA without consent with the exception of law enforcement purposes.

These laws can complicate even benign applications such as environmental tracking of bacterial communities in the sewage system. Previous shotgun sequence-based studies showed that such samples could contain small levels of human contamination. While human identification is probably not easy from these contaminants, their actual analysis could be interpreted as unlawful.

Since it is nearly impossible to avoid collecting human DNA in some settings, it will be beneficial to devise mitigation methods. These can include a warning message to the user that human DNA might be collected as part of the sequencing operation, which might violate local laws. Additional mitigation techniques could rely on a built-in suppression of reads that align to the human genome before analysis or storage. While previous work showed that residual human sequences can pass this procedure (Ame et al. 2015), implementing it will show that a reasonable effort was made to address privacy concerns. These mitigation techniques should be incorporated as part of the sequencer API by setting standard commands such as “get consent” or “suppress human DNA.” Such API commands will reduce the burden on developers and will standardize the mitigation steps. Besides technical mitigation, device manufacturers can facilitate a public discussion about social norms and etiquette on using such sequencing sensors by the general public. Notably, privacy concerns were one of the factors that hampered the adoption of Google Glass, emphasizing the importance of addressing these ELSI issues early.

Summary and the path forward

With the advent of miniature sequencing devices such as Oxford Nanopore’s MinION, we are on the cusp of truly democratizing DNA information by placing sequencers in the hands of the general public. Ubiquitous sequencing will create an incredibly powerful vantage point for observing massive amounts of DNA and RNA information within its natural context. This will open the possibility of integrating DNA data with other types of sensor information and obtaining a more comprehensive picture of the world around us.

This paper presented a range of potential applications that could benefit from sequencing sensors. But what could be the path forward to map which of these applications would be of interest to the general public? Importantly, most of the applications are in domains that are quite conservative and highly regulated, such as DNA forensics or medical devices. These might be less amenable testing grounds for new technologies. An interesting alternative is to consider citizen developers as an initial focus of sequencing sensors. Such hacker communities have played a pivotal role in the emergence of multiple technologies, from personal computers in the 70s, Linux in the 90s, and 3D printing and Raspberry Pi more recently. Similar communities exist in genetic genealogy, where nonscientists have created an impressive set of applications to analyze genetic information (e.g., GedMatch.com). By utilizing the power of citizen developers, it will be possible to test applications and accelerate R&D to mature sensors before testing them in one of the more conservative fields.

In any case, like any powerful technology, sequencing sensors will create a range of societal questions necessitating an ongoing discussion among all stakeholders about the benefits and risks and placing safeguards that will increase public trust in the concept.

Competing interest statement

Y.E. serves on the Scientific Advisory Board of Kailos Genetics and BigDataBio.

Acknowledgments

I thank Dina Zielinski, Brian Houck-Loomis, Itsko Pe’er, and Shree Nayar for useful discussion. Y.E. holds a Career Award at the Scientific Interface from the Burroughs Wellcome Fund. This study was partially supported by a National Institute of Justice (NIJ) award 2014-DN-BX-K089.

References

Aashinneko E, Meydan C, Chowdhury S, Jaroudi D, Boyer C, Bernstein N, Martiz JM, Reeves D, Gandara J, Chihangawala S, et al. 2015. Geospatial resolution of human and bacterial diversity with city-scale metagenomics. Cell Syst 1: 6–7.

Ame SK, Gardner SN, Marti JM, Sleazak TR, Gokhale MB, Allen JE. 2015. Using populations of human and microbial genomes for organism detection in metagenomes. Genome Res 25: 1056–1067.

Bieber FR, Brenner CH, Lazer D. 2006. Human genetics. Finding criminals through DNA of their relatives. Science 312: 1315–1316.

Bloch MS, Paunescu D, Stoessel FR, Mora CA, Stark WJ, Grass RN. 2014. Labeling milk along its production chain with DNA encapsulated in silica. J Agric Food Chem 62: 10615–10620.

Chiang C, Layer RM, Faust GG, Lindberg MR, Rose DB, Garrison EP, Marth GT, Quinlan AR, Hall JM. 2014. Speedseq: ultra-fast personal genome analysis and interpretation. bioRxiv 012179.

Creemers EE, Tijsen AJ, Pinto YM. 2012. Circulating microRNAs: novel biomarkers and extracellular communicators in cardiovascular disease? Circ Res 110: 485–495.

Davies K. 2010. The $1,000 genome: the revolution in DNA sequencing and the new era of personalized medicine. Simon and Schuster, New York.

Dekker C. 2007. Solid-state nanopores. Nat Nanotechnol 2: 209–215.

Dopico B, Lowe AL, Wilson JD, Menidio C, Grierson D. 1993. Cloning and characterization of avocado fruit mRNAs and their expression during ripening and low-temperature storage. Plant Mol Biol 21: 437–449.

Dugat-Bony E, Straub C, Teissandier A, Onésime D, Loux V, Monnet C, Irlinger F, Landaud S, Loëtzer-Perlat MN, Bento P, et al. 2015. Overview of a surface-ripened cheese community functioning by metacosymes analyses. PLoS One 10: e0124360.

Dwivedi DJ, Toftl JJ, Swystun LL, Pogue J, Liaw KL, Weitzt JJ, Cook DJ, Fox-Robichaud AE, Liaw PC. Canadian Critical Care Translational Biology Group. 2012. Prognostic utility and characterization of cell-free DNA in patients with severe sepsis. Crit Care 16: R151.

Erlich I, Narayanan A. 2014. Routes for breaching and protecting genetic privacy. Nat Rev Genet 15: 409–421.

Fijan S, Turk SS. 2012. Hospital textiles, are they a possible vehicle for healthcare-associated infections? Int J Environ Res Public Health 9: 3330–3334.

Gambino M. 2013. Creepy or cool? Portraits derived from the DNA in hair and gum found in public places. http://www.smithsonianmag.com/science-nature/creepy-or-cool-portraits-derived-from-the-dna-in-hair-and-gum-found-in-public-places-50266864/?no-ist.

Ge J, Eisenberg A, Budowle B. 2012. Developing criteria and data to determine best options for expanding the core CODIS loci. Investig Genet 3: 11.

Gilbert N. 2015. Why the ‘devious defector’ case is a landmark for US genetic-privacy law. Nature doi: 10.1038/nature.2015.17857.

Gine SK, Goba A, Andersen KG, Sealfon RS, Park DJ, Kanneh L, Jalloh S, Momoh M, Fullah M, et al. 2014. Genomic surveillance elucidates Ebola virus origin and transmission during the 2014 outbreak. Science 345: 1369–1372.

Goldman N, Bertone P, Chen S, Dessimoz C, LeProust EM, Sipos B, Birney E. 2013. Towards practical, high-capacity, low-maintenance information storage in synthesized DNA. Nature 494: 77–80.

Gonzalez A, Pettengill J, Vaughz-Baeza Y, Ottesen A, Knight R. 2014. Accurate detection of pathogens in microbial samples (and avoiding the conclusion that the pathovirus rules the earth). http://linguistics.berkeley.edu/ Up4mGEE.png; https://github.com/biocore/Platypus-Conquistador.

Green ED, Guyer MS, National Human Genome Research Institute. 2011. Charting a course for genomic medicine from base pairs to bedside. Nature 470: 204–213.
