Sequencing ancient RNA in bioarchaeology - Review

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Abstract

First-Generation Sequencing (FGS) revolutionized molecular biology in general, and bioarchaeology in particular. That allowed to study ancient DNA (aDNA) remains for the first time. Second-Generation Sequencing (SGS) further improved throughput, reducing time and cost per sequenced nucleotide base. Most importantly, it allowed to sequence full ancient-genomes, like the ones of Neanderthals. Third-Generation Sequencing (TGS) improved even more SGS throughput, also allowing to sequence single molecules, without requiring previous amplification or modification steps. That is revolutionary, overcoming putative biases associated to such steps. This opens the door for true-direct RNA sequencing [without retrotranscription into complementary DNA (cDNA)], including ancient RNA (aRNA). Different TGS platforms are arising, promising exciting new discoveries on bioarchaeology for both aDNA and aRNA.

Key words: functional genomics, epigenetics, messenger RNA, ancient transcriptomes, paleotranscriptomics.

Resumen

La secuenciación de primera generación (FGS; del inglés, “First-Generation Sequencing”) revolucionó la biología molecular en general, y la bioarqueología en particular. Ello permitió estudiar restos de ADN antiguo (ADNa) por primera vez. La secuenciación de segunda generación (SGS; del inglés, “Second-Generation Sequencing”) mejoró aún más el rendimiento, reduciendo el tiempo y coste por base nucleotídica secuenciada. Además, permitió secuenciar genomas antiguos completos, como los de neandertales. La secuenciación de tercera generación (TGS; del inglés, “Third-Generation Sequencing”) mejoró aún más el rendimiento de SGS, permitiendo además secuenciar moléculas individuales, sin requerir pasos previos de amplificación o modificación. Ello es revolucionario, evitando posibles sesgos asociados a tales pasos previos. Esto abre la puerta para la secuenciación directa verdadera del ARN [sin retrotranscripción en ADN complementario (ADNc)], incluyendo ARN antiguo (ARNa). Diferentes plataformas de TGS están surgiendo, prometiendo nuevos e interesantes descubrimientos en bioarqueología, tanto para ADNa como ARNa.

Palabras clave: genómica funcional, epigenética, ARN mensajero, transcriptomas antiguos, paleotranscriptómica.
Introduction

First-Generation Sequencing (FGS) revolutionized bioarchaeology, allowing to sequence ancient DNA (aDNA). The breakthrough was extended to ancient genomes with Second-Generation Sequencing (SGS), sporting higher throughput, faster turnaround and cheaper cost per read nucleotide base. Among others, that included sequencing of Neanderthal genome (Green et al. 2006; Noonan et al. 2006; Dorado et al. 2007-2015). Third-Generation Sequencing (TGS) further improves previous methodologies, allowing to sequence single-molecules, without previous modification or amplification steps, generating longer (kilobase; kb) reads.

Such advancements avoid biases associated to previous approaches, further increasing productivity, reducing time and cost (Bleidorn 2015). Therefore, SGS and TGS are considered High-Throughput Sequencing (HTS) technologies, being sometimes referred with the ambiguous “Next-Generation” Sequencing (NGS) terminology. Additionally, these methodologies can also be applied to study DNA epigenetics, including aDNA (Gokhman et al. 2014; Pedersen et al. 2014; Orlando et al. 2015; Seguin-Orlando et al. 2015; Smith et al. 2015).

Besides aDNA, it would be also interesting to sequence ancient RNA (aRNA), allowing to take functional snapshots of particular tissues at ancient times. Yet, RNA is more labile than DNA, and usually degrades after physiological activity supporting-life ceases. It is known that aDNA is usually scarce, physically broken (degraded) and may be also chemically modified. Indeed, aDNA may be hard to isolate and sequence, being prone to generate false-positive results due to modern DNA contamination. Thus, it is expected that aRNA should be harder to isolate and sequence, albeit messenger RNA (mRNA) is much shorter than genomic DNA, which is certainly an advantage for sequencing. But, is it possible to actually sequence ancient RNA? The answer is yes—to some extent, so far—, as described below (Fordyce et al. 2013; Guy 2013, 2014; Ng et al. 2014; Smith et al. 2014).

Sequencing ancient RNA

There are two basic approaches to sequence RNA: retrotranscription and direct sequencing. The first approach can be accomplished with FGS and SGS, to generate complementary DNA (cDNA) (Fig. 1) and further sequencing. Such methodology is sometimes known with the misleading name of RNA-seq, instead of the more suitable cDNA-seq terminology. Thus, different aRNA have been sequenced (Guy et al. 2014), including 50-year-old RNA peach latent-mosaic viroid (Guy 2013), white-clover cryptic virus-1 (Guy and Gerard 2016), 700-year-old viral genomes from caribou feces (Ng et al, 2014) and 750 year-old barley stripe-mosaic virus (Smith et al. 2014). Even the transcriptome of ancient (723 year-old) maize kernels has been sequenced (Fordyce et al. 2013). Curiously, since retroviruses insert into host genomes as Endogenous RetroViruses (ERV), their presence has been also detected sequencing modern genomes, as well as ancient ones. That is the case of 1,400 year-old
aDNA from coprolites (Rivera-Perez et al. 2015). Obviously, in this latter case, retrotranscription was carried out in living cells, being included here as a research curiosity.

Figure N° 1: RNA retrotranscription into cDNA. Example of retrotranscription, in which RNA is captured with paramagnetic beads (top-left black ball) and retrotranscribed into cDNA. Terminal-transferase activity of used retrotranscriptase allows to generate CC tails, to further hybridize single-stranded adapters, which become double-stranded after polymerization. RNA-cDNA hybrid is denatured, allowing to amplify the first-strand cDNA into double-stranded DNA, via Polymerase Chain Reaction (PCR). © 2016 Danandmike, Wikimedia Commons <http://commons.wikimedia.org> and Creative Commons <http://creativecommons.org>.

Fortunately, TGS platforms allow to directly sequence nucleic-acids, as indicated above, without previous retrotranscription or amplification steps, effectively opening to door to truly sequence RNA in general, as well as aRNA in particular. Among them are: i) Helicos BioSciences <http://www.helicosbio.com> true Single-Molecule Sequencing (tSMS), based on Sequencing-by-Synthesis (SbS). It was the first single-molecule sequencing technology available (Harris et al., 2008), being later on offered as a service by “Sequence the Lower Limit” (SeqLL) <http://seqll.com>; ii) Pacific Biosciences
(PacBio) <http://www.pacb.com> Single-Molecule Real-Time (SMRT) sequencing, derived from former Nanofluidics <http://nanofluidics.com>, via SbS; iii) Complete Genomics <http://www.completegenomics.com> combinatorial Probe-Anchor Ligation (cPAL) sequencing, based on Sequencing by Hybridization (SbH) and Sequencing by Ligation (SbL), which is offered as AllSeq <http://allseq.com> service (Fig 2); and iv) Oxford Nanopore Technologies <http://nanoporetech.com> sequencing, based on ionic (electrical)-current alterations, as nucleotides pass though biological or synthetic solid-state nanopores. It generates long reads of 230 to 300 kb, at a speed of 280 to 500 bases per second (bps). Interestingly, a miniature handheld MinION Mk 1B sequencer (87 g weight) has been successfully used by the National Aeronautics and Space Administration (NASA) <https://www.nasa.gov> for astrobiology. It has exciting potential for crew-disease diagnostics and nucleic-acid-based life research (Rezzonico 2014; John et al. 2016).

Figure N° 2: Single-molecule sequencing. Example of TGS (cPAL), in which an oligonucleotide anchor is hybridized to a known region of single-molecule to be sequenced. Further, a fluorescent interrogating-probe (containing A, C, G or T nitrogenous base at the interrogating position) is hybridized and ligated to the anchor. Non-hybridized probes are removed, and bound fluorescence is detected. The process is repeated for the rest of interrogating nucleotides. Then, the probe-anchor is removed and the process is repeated for other probes-anchors, until the target nucleic-acid is sequenced. © 2016 Suspencw, Wikimedia Commons <http://commons.wikimedia.org> and Creative Commons <http://creativecommons.org>.
TGS has been used to sequence not only DNA, but also RNA by Direct RNA Sequencing (DRS) (Ozsolak et al. 2009). It has also been used to sequence ancient genomes, including Pleistocene horse (Orlando et al. 2011; Ginolhac et al. 2012; Schubert et al. 2012).

Future prospects and concluding remarks


Feasibility of direct RNA sequencing has been demonstrated (Ozsolak et al. 2009), albeit the age limit of aRNA suitable for sequencing remains to be determined, as previously done for aDNA (Allentoft et al, 2012; Millar and Lambert 2013). The possibility to sequence ancient transcriptomes (paleotranscriptomics) is certainly exciting, mostly when using TGS. Indeed, the direct and longer reads of new TGS approaches (like those using nanopores) are quite convenient for such a goal. Yet, special assembly algorithms may be also needed for aRNA (as in the case of aDNA), since such material is usually degraded, as previously pointed out. Besides, increasing computing power will be required to analyze and compare the exponentially-growing data generated from genomics, transcriptomics and proteomics studies.

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