# Mapping of RNA modifications by direct Nanopore sequencing and JACUSA2

<sup>3</sup> Christoph Dieterich\*<sup>1,2,3</sup>, Amina Lemsara<sup>1,2</sup>, and Isabel Naarmann-de Vries<sup>1,2,3</sup>

<sup>1</sup>Klaus Tschira Institute for Integrative Computational Cardiology, University Heidelberg, 69120 Heidelberg, Germany

<sup>2</sup>Department of Internal Medicine III (Cardiology, Angiology, and Pneumology),

University Hospital Heidelberg, 69120 Heidelberg, Germany

<sup>3</sup>German Centre for Cardiovascular Research (DZHK)-Partner Site

Heidelberg/Mannheim, 69120 Heidelberg, Germany

10 Abstract

to be written

2

6

Keywords: Bayesian, 10X Genomics, Cell barcode assignment, Nonsensemediated mRNA decay (NMD)

## INTRODUCTION

Chemical modifications on DNA and histones, also known as epigenetics 15 marks, strongly impact gene expression during cell differentiation and in 16 several other biological programs. In the 1970s, it was recognized that RNA is also subjected to extensive covalent modification, and studies in the late 1980s revealed the widespread deamination of bases (termed RNA editing), 19 which can lead to recoding if it occurs within coding sequences. Impressive 20 development in the RNA modification field occurred during the past eight 21 years, with the discovery of an extensive layer of base modifications in mR-NAs. These can influence gene expression and have been already shown to 23 be involved in primary cellular programs such as stem cell differentiation, response to stress, and the circadian clock. The study of RNA modifications 25 and their effects is now referred to as epitranscriptomics, and it reveals strik-26 ing similarities to what is known for epigenomics. To date thirteen distinct 27 modifications have been identified on mRNA transcripts [?]. These modifications are catalyzed by a variety of dedicated enzymes and can be divided into two classes: modifications of cap-adjacent nucleotides and internal mod-30 ifications.

 $<sup>{\</sup>rm *christoph.dieterich@uni-heidelberg.de}$ 

In contrast to the m7G cap, the impact of internal modifications on gene 32 regulation has been less studied apart from RNA editing, which is mediated 33 by RNA deaminases (e.g. the ADAR family). The most widespread internal 34 mRNA modification is N6-methyladenosine (m6A). By modulating the pro-35 cessing of mRNA, m6A can regulate a wide range of physiological processes 36 and its alteration has been linked to several diseases???. The modifica-37 tion is catalyzed co-transcriptionally by a Mega-Dalton methyltransferase 38 complex, which includes the heterodimer METTL3-METTL14 and other 39 associated subunits?. This modification is reversible since two proteins of the AlkB-family demethylases can remove m6A from mRNA transcripts [??]. 41 In mammals, m6A preferentially localizes within long internal exons and at 42 the beginning of terminal exons at so-called DRACH motif (D = A/G/U, 43 R = A/G, H = A/C/U) sites [???]. Once deposited, m6A is recognized 44 by several reader proteins that can affect the fate of mRNA transcripts in nearly every step of the mRNA life cycle, which includes alternative splicing [??]. The best-described readers are the YTH domain family of proteins that 47 decode the signal and mediate m6A functions. By affecting RNA structure, 48 m6A can also indirectly influence the association of additional RNA-binding 49 proteins (RBPs) and the assembly of larger messenger ribonucleoprotein 50 particles (mRNPs). 51

Several approaches have been presented to map RNA modifications on 52 RNA. Herein, we focus on mRNA modification site detection in general and 53 on m6A in particular where antibody-based protocols (miCLIP), methylation-54 sensitive restriction enzyme assays (MazF) or transgenic approaches (TRIBE, 55 DART) have been presented. All of the aforementioned approaches rely on 56 high-throughput sequencing on the Illumina platform. This typically in-57 volves cDNA synthesis by reverse transcription and PCR-based library amplification. One recent addition to the tool is direct RNA single molecule 59 sequencing on the Oxford Nanopore Technology platform. While or software 60 workflow is able to deal with Illumina and Nanopore-based approaches, the 61 latter is the principal topic of our methods article. 62

## $_{53}$ MATERIALS

69

70

71

#### 64 ONT direct RNA sequencing

- 1. 500 ng polyA<sup>+</sup> RNA isolated from total RNA e.g. with Oligotex mRNA kit (Qiagen) or Dynabeads oligo dT<sub>25</sub> beads (Thermo Fisher Scientific). Store RNA at -80 °C and the mRNA purification kit as recommended by the manufacturer.
  - 2. Nuclease-free water. Store at room temperature.
  - 3. Direct RNA-sequencing kit (SQK-RNA002, Oxford Nanopore Technologies). Store at -20 °C.

- 4. NEBNext Quick Ligation Reaction Buffer (New England Biolabs). Store at -20 °C.
- 5. T4 DNA Ligase (New England Biolabs). Store at -20 °C.
- $^{75}$  6. dNTP Mix (10 mM each). Store at -20 °C.
- 76 7. SuperScript IV Reverse Transcriptase (Thermo Fisher Scientific). Store at -20 °C.
- 8. Agencourt RNAClean XP beads (Beckman Coulter). Store at 4 °C.
- 9. 70 % ethanol, freshly prepared.
- 10. Qubit dsDNA HS assay kit and Qubit Fluorometer (Thermo Fisher Scientific).
- 11. Flow cell priming kit (EXP-FLP002, Oxford Nanopore Technologies).
   Store at -20 °C.
- 12. Thermo cycler.
- 13. Gentle rotator mixer.
- 14. Magnetic stand for 1.5 ml tubes.
- 15. 1.5 ml DNA LoBind tubes (Eppendorf), 0.2 ml PCR tubes.
- 16. MinION or GridION sequencing device and MinION R9.4.1 Flow cells (FLO-MIN106D, Oxford Nanopore Technologies). Store Flow cells at 4 °C.

# Preparation of an in vitro transcriptome sample

#### Isabel

#### 93 Hardware requirements

- All analyses have been performed/tested on two alternative hardware sys-
- tems: a standard Linux desktop computer or an Apple iMac (Retina 5K,
- 96 ultimo 2014). The workflow requires a multi-core processor system with
- 97 minimal main memory of 16GB RAM and several GBs of free disk space
- 98 (depending on data set size).

## 99 Software dependencies and installation

Our analysis workflow has few requirements, which are detailed in Table 2. Specifically, to execute our workflow, the following prerequisites are neces-sary: a BASH shell, a JAVA runtime environment, a working PERL and R installation. Additional i.e. non-standard software to process and map Nanopore reads (bedtools, samtools and Minimap2) are obligatory, while the installation of a Nanopore read simulator (NanoSim) is optional and de-pends on your use case. Table ?? lists some additional R packages, which are required to run the R code. Detailed instructions on how to setup are found under https://github.com/dieterich-lab/MiMB\_JACUSA2\_chapter 

# 109 METHODS

Overview Figure 1

#### 111 Nanopore direct RNA sequencing

#### Isabel

- 1. Adjust 500 ng polyA<sup>+</sup> RNA to a total volume of 9 μl with nuclease-free water. Complete RT adapter ligation reaction (in 0.2 ml PCR tube) with 3 μl NEBNext Quick Ligation Reaction Buffer, 0.5 μl RNA CS (RCS, from SQK-RNA002), 1 μl RT-Adapter (RTA, from SQK-RNA002) and 1.5 μl T4 DNA Ligase. Incubate 10 min at room temperature.
- 2. Prepare reverse transcription master mix during ligation: 9  $\mu$ l nuclease-free water, 2  $\mu$ l 10 mM dNTPs, 8  $\mu$ l 5x SuperScript IV first strand buffer, 4  $\mu$ l 0.1 mM DTT.
- 3. Add the reverse transcription master mix to the ligation reaction and mix by pipetting. Add 2  $\mu$ l SuperScript IV reverse transcriptase and mix by pipetting. Incubate in a thermocycler with the following protocol: 50 min at 50 °C, 10 min at 70 °C, cool down to 4 °C.
- 4. Let the Agencourt RNAClean XP beads come to room temperature during incubation of reverse transcription. Carefully resuspend beads before use. Transfer reaction to a 1.5 ml DNA LoBind tube and mix with 72  $\mu$ l Agencourt RNAClean XP beads. Incubate 5 min at room temperature on a gentle rotator mixer.
- 5. Collect beads on a magnetic stand and remove supernatant. Wash pelleted beads two times (30 sec) with 200  $\mu$ l freshly prepared 70 % ethanol. Remove supernatant. Spin sample down and place on magnet again. Remove any residual ethanol.

- 6. Resuspend beads in 20  $\mu$ l nuclease-free water by gentle flicking and incubate 5 min at room temperature on a gentle rotator mixer. Collect beads on a magnetic stand and transfer 20  $\mu$ l eluate in a fresh 1.5 ml DNA LoBind tube.
- 7. For ligation of the RMX adapter, add the following to 20  $\mu$ l eluate: 8  $\mu$ l NEBNext Quick Ligation Reaction Buffer, 6  $\mu$ l RMX (from SQK-RNA002), 3  $\mu$ l nuclease-free water, 3  $\mu$ l T4 DNA Ligase. Mix by pipetting and incubate 10 min at room temperature.
- 8. Add 40  $\mu$ l carefully resuspended Agencourt RNAClean XP beads to the reaction and mix by pipetting. Incubate 5 min at room temperature on a gentle rotator mixer.
- 9. Collect beads on a magnetic stand and remove supernatant. Wash pelleted beads two times with 150  $\mu$ l wash buffer (WSB, from SQK-RNA002). Resuspend beads by flicking, spin down and return to magnetic stand. Remove supernatant from pelleted beads.
- 10. Resuspend beads in 21  $\mu$ l elution buffer (EB, from SQK-RNA002) by gentle flicking and incubate 5 min at room temperature on a gentle rotator mixer. Pellet beads on a magnetic stand and transfer 21  $\mu$ l eluate in a fresh 1.5 ml DNA LoBind tube.
- 11. Quantify 1  $\mu$ l of the library on a Qubit fluorometer with the Qubit dsDNA HS kit according to the manufacturerers protocol. Concentration should be usually in the range of 5 10 ng/ $\mu$ l.
- 12. Insert MinION R9.4.1 Flow cell in the MinION or GridION sequencing device and perform Flow cell check in the MinKNOW software.

  For successful sequencing of mammalian polyA<sup>+</sup> RNA at least 1,000 available pores are recommended.
- 13. Prepare Priming Mix by adding 30  $\mu$ l flush tether (FLT, from EXP-FLP002) to a vial of flush buffer (FB, from EXP-FLP002) and mix by pipetting. Open priming port. Remove air bubble from priming port by inserting the tip of a P1000 pipette into the priming port and slowly dialing up, until a small volume of storage buffer enters the pipette tip. Load 800  $\mu$ l Priming Mix via the priming port and carefully avoid introduction of air bubbles. Close the priming port and wait for 5 min.
- 14. Mix 20  $\mu$ l library with 17.5  $\mu$ l nuclease-free water and 37.5  $\mu$ l RNA running buffer (RRB, from SQK-RNA002) and mix by pipetting. Open the priming port and the sample port. Load 200  $\mu$ l Priming Mix via the priming port. Mix library by pipetting just before loading and load dropwise via the sample port. Carefully avoid introduction of air bubbles. Close the sample port and the priming port.

15. Start sequencing for 48 to 72 h in the MinKNOW software. Choose direct RNA-sequencing kit and high-accuracy basecalling as parameters. We recommend to adjust the output filter to a minimum Q score of 7 (instead of 9).

178 Preparation of an *in vitro* transcriptome sample

Isabel

- 180 Nanopore read processing
- 181 Minimap2 and samtools

Christoph

- <sup>183</sup> Use Case 1: Comparison of wildtype and knock-out samples
- 184 Xpore data

Christoph

186 Use Case 2: Comparison of wildtype and IVT samples

Christoph

187

188 Use Case 3: Comparison of wildtype to simulated IVT sample

Christoph

# 190 NOTES

191 Tips and Tricks

# 2 ACKNOWLEDGMENTS

The authors would like to thank Etienne Boileau, Thiago Britto Borges, Tobias Jakobi for proof-reading and comments. The authors are grateful

<sub>95</sub> to Marek Franitza for running the experiments on the 10x platform and to

196 Christian Becker for running ONT sequencing. This work was supported by

197 Informatics for Life funded by the Klaus Tschira Foundation.

# 98 FIGURE CAPTIONS

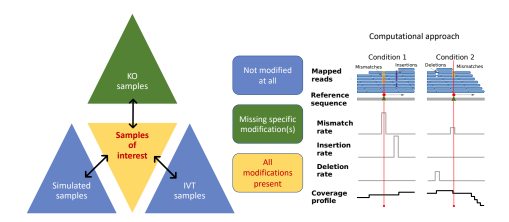


Figure 1: General outline of RNA modification detection by JA-CUSA2. A key feature of our approach is that multiple replicates can be compared as shown on the left. Samples of interests where all modifications are present could be compared with either KO samples where the modification of interest is missing or IVT/simulated samples where all modifications are absent. Read stacks (in blue) are compared head-to-head as shown on the right.

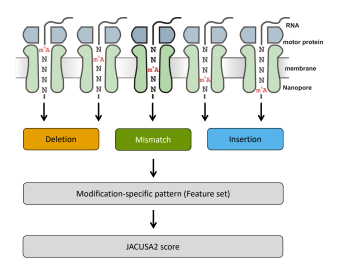


Figure 2: Motivation of 5mer context for RNA modification mapping. The nanopore covers 5 consecutive RNA residues. That is why we consider a 5mer context and derive 3 principal features for every position within a given 5 mer (15 features in total, with a central A residue in this example). We evaluate each feature set by previously learned patterns and compute a final score for modification site detection.

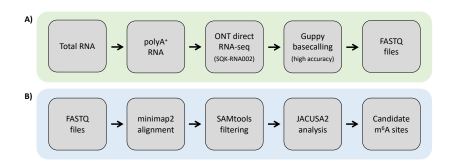


Figure 3: Experimental and computational workflow. tbd

Software	Version	Description
Minimap2	https://github.com/lh3/minimap2	https://lh3.github.io/minimap2/
	v2.22 or later	
samtools	https://github.com/samtools/	http://samtools.github.io/
	samtools v1.12 or later	
JAVA	openjdk 11.0.12 2021-07-20 - JAVA 11 or	OpenJDK Runtime Environment
	later	
R	https://www.r-project.org/ version	The R Project for Statistical Comput-
	3.5.1 or later	ing
PERL	https://www.perl.org/ version 5.28.1	Perl is a highly capable, feature-rich
	or later	programming language
BASH,	should be part of your Linux distribution	Misc.
sed,		
awk		
bedtools	https://github.com/arq5x/bedtools2	Perl is a highly capable, feature-rich
	version 2.29.2 or later	programming language
NanoSim	https://github.com/bcgsc/NanoSim	NanoSim is a fast and scalable read
	version 3.0.2 or later (optional)	simulator that captures the technology-
		specific features of ONT data

Table 1: Software dependencies blubba

# 199 TABLE CAPTIONS

# TABLES

R Pack-	Version	Description
ages		
ggplot2	https://cran.r-project.org/web/	ggplot2 is a system for declaratively
	packages/ggplot2/index.html - gg-	creating graphics, based on The Gram-
	plot2_3.3.0 or later	mar of Graphics.
NMF	https://cran.r-project.org/web/	Provides a framework to perform Non-
	packages/NMF/index.html-NMF_0.22.0	negative Matrix Factorization (NMF).
	or later	

Table 2: R Package dependencies blubba