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- The effect of reverse transcription enzymes and conditions on high throughput
- amplicon sequencing of the 16S rRNA
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12 ABSTRACT

It is assumed that the sequencing of ribosomes better reflects the active microbial community than the 13 sequencing of the ribosomal RNA encoding genes. Yet, many studies exploring microbial communities 14 in various environments, ranging from the human gut to deep oceans, questioned the validity of this 15 paradigm due to the discrepancies between the DNA and RNA based communities. Here we focus on 16 an often neglected key step in the analysis, the reverse transcription (RT) reaction. Previous studies 17 showed that RT may introduce biases when expressed genes and ribosmal rRNA are quantified, yet its 18 effect on microbial diversity and community composition was never tested. High throughput sequencing 19 of ribosomal RNA is a valuable tool to understand microbial communities as it better describes the active 20 population than DNA analysis. However, the necessary step of RT may introduce biases that have so 21 far been poorly described. In this manuscript, we compare three RT enzymes, commonly used in soil 22 microbiology, in two temperature modes to determine a potential source of bias due to non-standardized 23 RT conditions. In our comparisons, we have observed up to 6 fold differences in bacterial class abundance. 24 A temperature induced bias can be partially explained by G-C content of the affected bacterial groups. 25 thus pointing towards a need for higher reaction temperatures. However, another source of bias was due 26 to enzyme processivity differences. This bias is potentially hard to overcome and thus mitigating it might 27 require the use of one enzyme for the sake of cross-study comparison. 28

29 INTRODUCTION

Massively parallel amplicon sequencing revolutionized our view of microbial world: by sequencing 30 a taxonomic tag such as 16S rRNA encoding gene, it allows taxonomic description of the microbial 31 communities (Quammen, 2018). However, the existing approaches introduce caveats: the DNA amplicon 32 sequencing may capture 'relic DNA', which is a recalcitrant genetic material from dead cells or naked 33 DNA (Carini et al., 2016) in addition, amplicon sequencing carries technical biases due to sample 34 preparation, DNA extraction methods (Pan et al., 2010), amplification reaction (Pfeiffer et al., 2014) and 35 analysis (Pollock et al., 2018). Moreover, DNA-based microbiome information can describe the total 36 community, but it cannot report which members are metabolically active (Blazewicz et al., 2013). In 37 contrast to DNA based tools, analysis of ribosomes can describe the metabolically active members of a 38 given community. The combination of data generated from rRNA encoding genes and ribosomes led to a 39 wide range of ecological insights, including the response to climatic changes (Angel et al., 2013), pH and 40 water availability (Romanowicz et al., 2016), and biogeochemical processes (Freedman et al., 2015). 41 Ribosomal analysis studies are based on an assumption that ribosomes are more abundant in active cells 42 compared to dormant ones (Blazewicz et al., 2013; Lennon and Jones, 2011). However, this assumption 43

⁴⁴ may not always be correct. Dormant bacteria may be misclassified as active, when ribosomes are present

in cells and spores that are inactive (Segev et al., 2013; Blagodatskaya and Kuzyakov, 2013). In contrast,

active bacteria with low metabolic turnover and low ribosomal count could be labeled as dormant when 46 sequencing depth is insufficient (Steven et al., 2017; Joergensen and Wichern, 2018). In spite of various 47 biases that introduce discrepancies in the community structure (Forney et al., 2004), ribosomal analysis 48 can capture the biological variability highlighting large differences between samples. However, if more 49 50 subtle differences are of interest, technical biases could confound biological interpretations (Lever et al., 2015; McCarthy et al., 2015). This is due to specific challenges introduced RNA based analysis (Bustin 51 and Nolan, 2004, 2017). Therefore, to confidently compare results across ribosome-based amplicon 52 sequencing studies, we must determine which component of the analysis: RNA extraction, processing or 53 data analysis may influence the outcome and introduce biases. 54 Prior studies focused on biases in the steps of RNA extraction, amplification and sequencing, but 55 disregard any biases that may occur during RT (Creer et al., 2016). At the crucial step of RT, most 56 researches simply 'follow the manufacture instructions' (Table 1). However, RT kits typically detail a 57 wide range of temperatures, primer, template and reaction options, which may lead to different results. 58 The reverse transcriptase (RT) enzyme requires sequence priming to initiate a reaction. Primers could be 59 poly-A complementary, random or sequence specific. Poly-A priming is limited to eukaryotic mRNA 60 which makes it unsuitable for use with ribosomal taxonomic tags. Opinions vary about the usefulness of 61 random and sequence-specific priming for the analysis of microbiomes: Random priming may produce 62 higher yield of cDNA and improve the detection limit (Zhang and Byrne, 2015; Ståhlberg et al., 2004a), 63 but may decrease the reproducibility and introduce bias (Bustin and Nolan, 2004; Hansen et al., 2010). 64 Sequence specific primes require fine tuning of the reaction conditions and higher template concentration 65 than random priming (Ståhlberg et al., 2004b). Moreover, the results of the RT reaction is determined not 66 only by the type of the RT enzyme used, but also by the reaction conditions (Curry et al., 2002; Ståhlberg 67 et al., 2004a,b; Bustin and Nolan, 2004; Sieber et al., 2010). 68

Ideally RT efficiency is near 100 %, but in practice it varies dramatically: 90 % efficiency was reported
 for SuperScript III (mutated MMLV RT) (Ståhlberg et al., 2004a), 20 % for Murine Leukemia Virus
 (M-MLV) RT (Curry et al., 2002), and as low as 2 % for Avian Myeloblastosis Virus (AMV) RT (Ståhlberg
 et al., 2004a). Moreover almost two order of magnitude difference were reported between mutated and
 wild-type AMV RT (Ståhlberg et al., 2004a).

To the best of our knowledge no study has yet compared the RT reaction conditions for environmental microbiome profile. We hypothesize that during RT reactions, varying RT enzyme types and temperature conditions will yield different results in microbial diversity and community composition. We further predict that variations in communities will be G-C dependent. To test our prediction, we present a comparative study of commonly used RT enzymes in the field of environmental microbiology as well as a comparison of two different reaction temperatures.

Manufacturer	RT Enzyme	RT origin	Temperature Suggested	e [°C] Used	RNA type	Primer type	Ref
Promega	MMLV ImProm-II	MMLV AMV	37–42 37–55	NA NA 42 NA	rRNA rRNA & mRNA rRNA & mRNA rRNA	926R Random hexamers Random hexamers Random hexamers	Carson et al. (2010) Pratscher et al. (2011) Angel et al. (2013) Ke et al. (2015)
Qiagen	QuantiTect Omniscript	Quantiscript Quantiscript	42 <i>-</i> 50 37	NA 37 NA NA	rRNA rRNA mRNA mRNA	Unique RT Primer Mix Random hexamers Random hexamers invA-R	Barnard et al. (2015) Placella et al. (2012) Paulin et al. (2013) García et al. (2010)
Takara	PrimeScript II	AMV	42-50	NA NA	mRNA rRNA	Random hexamers Random hexamers	Huang et al. (2016) Che et al. (2018)
Roche	Roche reverse transcription kit	AMV	42-60	42 & 50 42 & 50	rRNA rRNA	Random hexamers Random hexamers	Nunes et al. (2018) Jurburg et al. (2017)
Thermo Fisher	MMLV SuperScript-II	MMLV MMLV	37-42 42-55	45 NA NA NA	rRNA rRNA rRMA mRNA	900R Random hexamers 1492R Random hexamers	Lillis et al. (2009) Baldrian et al. (2012) Degelmann et al. (2009) Nacke et al. (2014)
	SuperScript-III	MMLV	42-55	NA NA NA	rRNA rRNA	Random hexamers 27F & LR3	Angel and Conrad (2013) Romanowicz et al. (2016)

Table 1. Literature overview of RT conditions applied in soil microbiological studies.

MATERIALS AND METHODS

81 Study site and sample collection scheme

- ⁸² Soil samples were collected at the central Negev Desert highlands, Israel (Zin Plateau, 30°86'N, 34°80'E)
- at an established ecological research site. The mean annual precipitation at the sampling site is 90 mm and

the mean annual temperature is 30 $^{\circ}$ C (LTER data). Samples were collected under the canopy of perennial

shrub *Hammada scoparia* in October 2015 at the end of the dry season as previously described (Baubin

- et al., 2019). Briefly, sampling was conducted in seven random blocks, each providing two technical
- ⁸⁷ replicates resulting in 14 samples. Samples were collected from the top 5 cm of the soil, following the
- removal of crust and debris. The soil samples were processed within 24 h of collection. Samples were homogenized using 2 mm sieve and the duplicates from each block were composited. This resulted in
- ⁸⁹ homogenized using 2 mm sieve and the duplicates from each block were co
 ⁹⁰ seven final replicates.

RNA preparation

- ⁹² Total RNA was extracted from each of the seven samples using a phenol-chlorophorm extraction pre-
- viously described by Angel (2012). The reaction buffer pH was adjusted to 5. The total RNA was
- ⁹⁴ subsequently purified with the MagListoTMTotal RNA Extraction Kit (Bioneer, Daejeon, Republic of
- ⁹⁵ Korea). Contaminant DNA was removed using a DNAse I from the MasterPure RNA Purification Kit
- ⁹⁶ (Epicenter, Madison, WI, USA) with two successive treatments of 30 min according to manufacturer's
- ⁹⁷ instructions. The reaction mixture was purified using the MagListoTMTotal RNA Extraction Kit (Bioneer).
- ⁹⁸ The absence of contaminant DNA was verified using total bacterial primers 341F (5' CCTACGGGAG-
- ⁹⁹ GCAGCAG 3') and 515R (5' TTACCGCGGCTGCTGGCAC 3') (Klindworth et al., 2013) and DreamTaq ¹⁰⁰ DNA polymerase (Thermo Scientific, Waltham, MA, USA) under the following conditions: 95 °C for
- 5 min, followed by 26 cycles of 95 °C for 15 s, 60 °C for 30 s and 72 °C for 30 s for extension followed by
- 102 72 °C for 5 min for final extension. If amplification was detected the sample was discarded, re-extracted,
- ¹⁰³ purified and tested. Only DNA-free samples were used in this study.

104 Reverse transcription reaction conditions

RT kits used in this experiment were chosen to represent the most commonly used enzymes in the 105 field (Figure 1). All seven samples were reverse transcribed by the same kit to reduce batch effects. 106 Each RT kit used in this study originated from a different source: (I.) ImProm-II Reverse Transcription 107 System enzyme (Promega, Madison, WI, USA) originates from AMV RT, (II.) SuperScript IV Reverse 108 Transcriptase Kit enzyme (ThermoFisher Scientific, Waltham, MA, USA) originates from MMLV RT 109 and (III.) TGIRT originates from the mobile group II introns reverse transcriptase (Mohr et al., 2013) 110 TGIRTTM-III Enzyme (InGex,St. Louis, MO, USA). Each reaction consisted of 50 ng of total RNA 111 template, measured by Ouanti-iTTMRNA Assay Kit (ThermoFisher), and random hexamer primers 112 (0.5 µg/reaction). Template and primer mix were heated to 70 °C (ImProm-II) or 65 °C (SuperScript 113 IV). Each reaction was subsequently cooled to 4 °C for 5 min and incubated at 42 °C (ImProm-II), 55 °C 114 (ImProm-II and Superscript) or 57 °C (TGIRT) for 60 min (ImProm-II), 120 min (TGIRT) or 10 min 115 (SuperScript IV). All reactions were terminated and DNA was removed by alkaline lysis using 2 µl of 116 1 M NaOH, incubating for 12 min at 70 °C. After which the reaction was neutralized using 4 µl of 0.5 M 117

acetic acid (Table 2).

119 Illumina sequence preparation

The V3 and V4 regions of the resulting cDNA were amplified using 341F (5' CCTACGGGAGGCAGCAG 120 3') and 806R (5' GGTCTGGACTACHVGGGTWTCTAAT 3') (Klindworth et al., 2013) primers. Each 121 reaction was performed in triplicate and consisted of 1 mM bovine serum albumin (Takara, Kusatsu, 122 Japan), 2.5 µl of 10x standard buffer, 5 µM primers, 0.8 mM dNTPs, 0.4 µl DreamTaq DNA polymerase 123 (Thermo Scientific, Waltham, MA, USA), and 4 µl of template cDNA. The reaction mixtures were 124 subsequently amplified using the following PCR conditions: 95 $^{\circ}$ C for 30 s, 27 cycles of 95 $^{\circ}$ C for 15 s, 125 $50 \,^{\circ}$ C for 30 s, 68 $^{\circ}$ C for 30 s and 68 $^{\circ}$ C for 5 min. Resulting amplicon presence was verified using 1.5 % 126 agarose gel electrophoresis. Resulting technical triplicates were combined, and the sequencing libraries 127 were constructed using the TruSeq[®] DNA Sample Preparation Kit (Illumina, San Diego, CA, USA) 128 following the manufacturer's recommendations. The amplicon libraries were sequenced (250x2 base pairs, 129 pair-end) on the Illumina MiSeq System platform at the Research Resources Centre at the University of 130 Illinois. 131

	RT kit	Primers	Thermo cycling		Reaction mix	
Manufacturer			Temperature [°C]	Time [min]	Reactant	Amount
		Random Hexamers				
Promega	Im-Prom II	(500 ng/reaction)	70	5	DTT	10 µM
		× 5 ,	4	5	Tris-HCl	50 mM
			25	5	KC1	75 mM
			42	60	MgCl ₂	2.5 mM
			70	15	dNTP	0.5 mM
					RNAse inhibitor	0.5 µl/20 µl
		Random Hexamers				
Promega	Im-Prom II	(500 ng/reaction)	70	5	DTT	10 µM
Ū.			4	5	Tris-HCl	50 mM
			25	5	KC1	75 mM
			55	60	MgCl ₂	2.5 mM
			70	15	dNTP	0.5 mM
					RNAse inhibitor	0.5 µl/20 µl
ThermoFisher	SuperScriptIV	Random Hexamers (2.5 µM)	65	5	DTT	5 µM
	1 1	× • • •	0	1	Tris-HCl	50 mM
			23	10	KC1	50 mM
			55	10	MgCl ₂	4 mM
			80	10	dNTP	0.5 mM
					RNAse inhibitor	0.5 µl/20 µl
		Random Hexamers				
TGIRT	TGIRT-III	(500 ng/reaction)	65	5	DTT	5 µM
		,	0	1	Tris-HCl	10 mM
			23	10	EDTA	1 mM
			58	120	MgCl ₂	4 mM
			80	10	dNTP	0.5 mM
					RNAse inhibitor	0.5 µl/20 µl

Table 2. A summary of conditions applied to the different reaction conditions. .

132 Sequence analysis

Resulting paired end sequences were merged using the CASPER program (Kwon et al., 2014), and the
 resulting merged reads were clustered using the UPARSE pipeline according to the recommended settings
 (Edgar, 2013). The resulting OTU representative sequences were taxonomically assigned with the SINA

incremental aligner using a lowest common ancestor algorithm (Pruesse et al., 2007) and the SILVA

database version 132 (Quast et al., 2013). All sequences retrieved in this study were uploaded to European

¹³⁸ Nucleotide Archive (https://www.ebi.ac.uk/ena) submission number PRJEB32237.

139 Class enrichment plot data preparation

To explore whether G-C content contributed to differences in relative abundances of different taxa, we ran
the following analysis: for each reaction condition we calculated pairwise comparisons at the class level:
we normalized the proportional enrichment in each respective reaction conditions following Equation 1
(Figure 2 and Supplementary Figure S2), where the A and B represent a class at the different conditions.

44
$$Class_{normalized} = \frac{A-B}{A+B}$$
 (1)

The error bars represent a standard deviation, which have been calculated as a standard deviation of each category and normalized according to the Equation 2. The δA and δB represent the standard deviation of class A and B. Details of deriving this equations are specified in Supplementary Equation S1.

48
$$\delta Class_{normalized} = \frac{2}{(A+B)^2} \sqrt{B^2 \delta A^2 + A^2 \delta B^2}$$
(2)

149 Statistical analysis

All data analysis was performed in R v3.4.3 (R Core Team, 2018). The dataset was sub-sampled (rarified) 150 to an even depth of 9000 sequences per sample using python numpy package v1.15.4 (Van Der Walt et al., 151 2011). The subsampling removed five samples from the dataset. Additionally, three more samples were 152 removed as outliers ("https://gitlab.com/stovicek/rt article/blob/master/data preparation.ipynb). In order 153 to equalize the number of replicates, two random samples were removed from the TGIRT dataset. This 154 reduced the number of replicates to four samples per experimental category. The sample diversity was 155 analyzed using the vegan package v2.5-2 (Oksanen et al., 2018). The data was visualized using the R 156 package ggplot2 package v2.2.1 (Wickham, 2016) and python package matplotlib (Hunter, 2007). 157

158 RESULTS

159 Sample preparation and diversity analysis

After removing obvious outliers, four samples from each condition were analyzed. The summary of the analysis and all the code used to produce each figure is included in the Supplementary File S1 and S2. The changes in bacterial diversity among the tested conditions were expressed using species count, Pielou's evenness index (Pielou, 1967) and Shannon diversity index (Shannon and Weaver, 1949). Despite observed trends in the diversity indices, no statistically significant differences were detected (Supplementary Figure S1, Supplementary diversity_statistics.ipynb).

166 Relative abundance plot

A relative abundance of major taxonomic classes is depicted in the Figure 1. Each column is an av-167 erage of four biological replicates. OTUs (operational taxonomic units) that were not taxonomically 168 assigned at this level are summarized as "Unclassified" ($\approx 2\%$). Various patterns (detailled below) 169 were detected among the experimental conditions: some patterns could be attributed to differences in 170 reaction temperature (which ranged from 42 °C to 55 °C and 57 °C). Other patterns could be linked to 171 enzyme type. RT reactions with SupeScript IV and TGIRT RTs yielded no significant differences in 172 class relative abundances. However, transcription with ImProm-II RT at a similar temperature (≈ 55 °C) 173 yielded different abundances: specifically, the abundances of Alphaproteobacteria, Bacteroidia, Deltapro-174 teobacteria, Oxyphotobacteria, Rubrobacteria and Verrucomicrobidae decreased. However, Chloflexia, 175 Gammaproteobacteria and Thermoleophilia abundances increased when their ribosomes were transcribed 176 with ImProm-II RT (Figure 1, Supplementary Table S1). When transcription occured at lower temperature 177 (42 °C), relative abundances of Bacilli, Deltaproteobacteria and Oxyphotobacteria were enriched, while 178 Actinobacteria, Chloroflexia and Acidimicrobia were depleted under the same conditions (Figure 1 and 179 Supplementary Table S1). 180

181 Class enrichment plot

The Figure 2 also depicts the weighted average of the G-C content in each class. The proportional 182 comparison is interpreted as follows: A value of zero in the proportional comparison represents the 183 taxonomic class count that is exactly equal between the two compared groups (Figure 2). A value of 1 184 or -1 is assigned when a given taxonomic class is only present in one category and absent in another, 185 respectively. In general, there was a tendency towards lower G-C content lower temperature of ImProm-186 II (Figure 2,a, and Supplementary Figure S2, a and b). No statistically significant differences were 187 detected between the profiles resulting from RT of SuprScript-IV or TGIRT (Supplementary Figure S2, 188 d). The taxa Gemmatimonadetes, Fibrobacteria and Thermoanaerobaculia were relatively insensitive 189 to the RT conditions. However the majority of the classes were enriched in some conditions. (I.) the 190 rate of RT was only sensitive to temperature for the classes Alphaproteobacteria, Gemmatimonadetes, 191 Fibrobacteria, Thermoanaerobaculia, TK10, and Blastocatellia (Figure 2, a). These groups tend to have 192 extreme GC content (both high and low). (II.) Gammaproteobacteria, Planctomycetacia and Phycisphaerae 193 are relatively insensitive to the reaction temperature (Figure 2, a), but their abundances vary with different 194 RT enzymes (Figure 2, b, Supplementary figure S2). 195

196 Class enrichment statistics

We calculated a linear regression, where the response variable was the relative proportion of each class 197 between two tested categories, and the explanatory variable was the G-C content (the assumptions tests 198 and plots can be found in the Supplementary file S2). The linear regression assumptions were tested: 199 in case of two condition pairs (ImProm-II at 42 °C & SuperScript-IV as well as ImProm-II at 55 °C & 200 SuperScript-IV), the assumption conditions were not met (Supplementary file S2). Since we cannot 201 confidently discard the null hypothesis in these cases, we do not consider these two tests significant. 202 Therefore, we are considering only the ImProm-II 42 °C & ImProm-II 55 °C as well as ImProm-II 42 °C & 203 TGIRT as a significant outcome. Differences in the remaining cases cannot be explained by the weighted 204 G-C content alone (Table 3). 205

206 DISCUSSION

As high throughput sequencing has become increasingly accessible in recent years, researchers urgently call for method standardization to allow for accurate cross-study comparisons (Pan et al., 2010; Blago-

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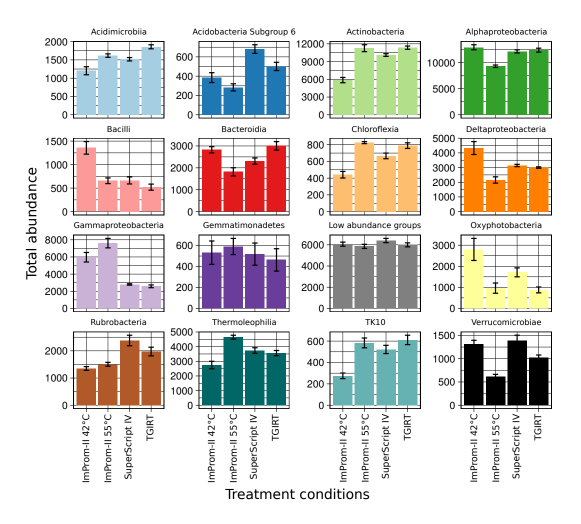


Figure 1. Relative abundance of main classes across each tested condition. Only top 15 % of the most abundant classes are represented and the rest is summarized in the 'Low abundance' category. The x axis show different enzymes and conditions. The y axis shows an average relative abundance. Each category is an average of 4 samples.

datskaya and Kuzyakov, 2013). With this motivation, researchers developed new platforms that offer 209 protocols and standardized methods for data acquisition from DNA resources, such as the Earth Micro-210 biome Project (http://www.earthmicrobiome.org/) that standardizes DNA amplicon sequencing. Despite 211 the success with standardizing methods and protocols for DNA-based analysis, to this date, there is 212 RNA-based methods have not been standardized, despite the discrepancies repeatedly reported between 213 the RNA and DNA based analyses (Blazewicz et al., 2013; Carini et al., 2016; Dlott et al., 2015) and 214 the plethora of methods used in these studies (Table 1). The analysis of rRNA adds specific biases to 215 high throughput sequencing analysis, such as reduced template stability compared to DNA, RT priming 216 bias, and linearity of RT reaction (Bustin and Nolan, 2004). These biases need to be either minimized or 217 standardized. 218

In this study, we focused on one crucial step in the RNA analysis that was previously overlooked: the transcription of RNA to cDNA (Table 1). Several biases connected to the RT reaction have been described for RT-qPCR, such as quantification of expressed genes (Bustin and Nolan, 2004, 2017; Zhang and Byrne, 2015) in RNA-Seq, i.e., primer related bias of expressed transcripts (Hansen et al., 2010). Yet, the role of RT in diversity patterns was not yet investigated in the context of high throughput sequencing of ribosomes. Here, we focus on the role of enzyme and reaction temperature in shaping the diversity and composition of ribosome-based communities. We have compared four RT enzymes commonly used in soil

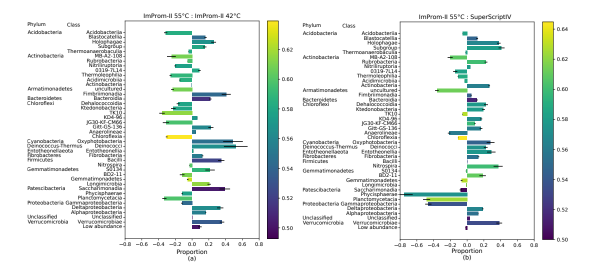


Figure 2. A proportional comparison of most abundant classes between the ImProm-II at 42 $^{\circ}$ C and 55 $^{\circ}$ C (a) and the ImProm-II at 55 $^{\circ}$ C and SuperScriptIV at 55 $^{\circ}$ C (b). Enrichment is expressed such that a class that is equally proportional in both conditions, has a value of 0. If the class shows in one condition but is absent from another, its value would be equal to 1 or -1 respectively. Furthermore, a weighted average of the GC content of each class is expressed as the bar color. Each value is an average of 4 biological replicates.

Condition 1	Condition 2	Adjusted R2	t value	p value	Significance	Note
ImProm-II 42 °C	ImProm-II 55 °C	0.429	5.366	4.89E-06	***	
ImProm-II 42 °C	SuperScript IV	0.1373	2.692	0.0127	*	•
ImProm-II 42 °C	TGIRT	0.2032	3.231	0.00264	**	
ImProm-II 55 °C	SuperScript IV	0.0841	-2.097	0.0431	*	•
ImProm-II 55 °C	TGIRT	0.03624	-1.546	0.131		
SuperScript IV	TGIRT	0.03634	1.548	0.130		

Table 3. The linear regression statistics.

We used a GC content as an explanatory variable of a the class enrichment. The rows marked with a \bullet do not fulfill all test assumptions (see supplementary diversity_statistics.ipynb).

microbiology (Table 1) at two distinct temperature modes (42 °C and 55 °C–57 °C). Then we analyzed temperature and RT enzyme-related effects on the resulting community profiles (Table 2).

Under different reaction conditions, we detected differences in the relative abundance of bacterial 228 classes portrayed by different reaction conditions (Figure 1). Some observed differences can be attributed 229 to the combined effect of reaction temperature and average template G-C content (Figure 2 and Supple-230 mentary Figure S2). As expected, this effect is clearest when we applied the same enzyme (ImProm-II) 231 at two reaction temperatures (42 $^{\circ}$ C and 55 $^{\circ}$ C), then the G-C content had the highest prediction power 232 $(t = 5.366, p = 4.8 \times 10^{-5}, Table 3)$. Likewise, in every comparison of RT enzymes at low and high 233 temperature, G-C content affected the relative abundance of taxonomic classes with statistical significance. 234 Although the RT reactions are commonly performed at 42 °C (Table 1), our results indicate that this 235 reaction temperature is too low to allow successful RT of some soil community taxa, in particular species 236 with higher G-C content. 237

When transcription was performed with different enzymes under similar reaction temperatures, relative 238 abundances of taxa differed notably (Figure 1 and 2). Although the RTs of SuperScript-IV and TGIRT 239 originate from different organisms, they yielded similar taxa abundances. Reactions with ImProm-II 240 yielded different profiles. These differences cannot be explained by the G-C content (Table 3), but could 241 be attributed to ribosome properties and the efficiency of RT. The ribosomes extracted from the soil 242 environment were diverse and probably differ in their secondary and tertiary structures (Yilmaz et al., 243 2006) post-transcriptional modifications (Schwartz and Motorin, 2017). Thus RTs kinetics would differ. 244 The discrepancies reported in this study raise further questions: how would one decide which enzymes 245 or temperatures best reflect the active community composition? It has been suggested that the total 246

²⁴⁷ community could be used as a reference to accurately deduce the diversity. Furthermore, this study was
 ²⁴⁸ performed on desert soil samples and the effects in other environments remain to be determined.

We previously demonstrated that the total and active communities in the Negev soil used in this study differ in both abundance profiles and community composition during the dry season, while during the wet season, no differences were detected (Baubin et al., 2019). During the dry season, a "phantom taxa"

(Klein et al., 2016), Deinococcus-Thermus, comprised $\approx 30\%$ of the total soil community (Baubin et al.,

²⁵³ 2019) but was undetected in the active community of the dry season by any of the methods used here.

These results suggest that the DNA-based total community may differ from the RNA-based community and thus cannot be used as a reliable reference for diversity. Furthermore, our results underline a need to

256 standardize and specify the RT conditions that allow cross-study comparisons. The scale of the effect

of RT conditions on the RNA-based community might vary with a studied biome. Dependending on

obvious factors such as GC content (discussed above), as well as poorly studied factors such as ribosomal

²⁵⁹ post-transcriptional modifications. Therefore, we recommend verifying each case separately before

²⁶⁰ attempting a cross-study comparison.

261 CONCLUSION

We have tested commonly utilized RT enzymes at assorted temperatures and observed marked differences 262 in the output community structure. These differences were attributed to RT type and reaction conditions. 263 We suggest that RT reaction conditions may dictate the diversity of a given community and therefore 264 the exact conditions should be detailed in full [i.e., the common notation, "according to manufacturer 265 instructions" does not provide sufficient information (Table 1)]. Furthermore, RT should be performed at 266 a sufficiently high temperature to minimize the G-C bias, preferably at 55 $^{\circ}$ C. Lastly, we suggest that the 267 same RT enzyme should be used across comparable studies, since we detected discrepancies between 268 RT enzymes performing at equivalent conditions (Figure 2, b). Here, we highlight, for the first time, 269 the need for standardisation and careful consideration of RT reaction conditions in studies describing 270 ribosome-based diversity and community composition. 271

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