Nuclease-mediated depletion biases in ribosome footprint profiling libraries

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ABSTRACT
Ribosome footprint profiling is a high-throughput sequencing–based technique that provides detailed and global views of translation in living cells. An essential part of this technology is removal of unwanted, normally very abundant, ribosomal RNA sequences that dominate libraries and increase sequencing costs. The most effective commercial solution (Ribo-Zero) has been discontinued as a standalone product and a number of new, experimentally distinct commercial applications have emerged on the market. Here we evaluated several commercially available alternatives designed for RNA-seq of human samples and find them generally unsuitable for ribosome footprint profiling. We instead recommend the use of custom-designed biotinylated oligos, which were widely used in early ribosome profiling studies. Importantly, we warn that depletion solutions based on targeted nuclease cleavage significantly perturb the high-resolution information that can be derived from the data, and thus do not recommend their use for any applications that require precise determination of the ends of RNA fragments.

Keywords: RNA-seq; Ribo-Zero; rRNA depletion; ribo-seq; ribosome profiling

INTRODUCTION
Ribosome footprint profiling (Ingolia et al. 2009, 2019; McGlinncy and Ingolia 2017) provides nucleotide-precision snapshots of ribosome positions transcriptome-wide. This technique has been used by many groups to study a wide variety of biological problems, ranging from the mechanisms of translation regulation and mRNA quality control to questions about viral infection, stem cell differentiation, and mouse neurobiology. The technique works by sequencing the short (~30 nt) ribosome-protected mRNA fragments (footprints) that result from nuclease digestion of a cellular lysate (Steitz 1969; Kozak and Shatkin 1976; Wolin and Walter 1988). This nuclease digestion process inevitably causes widespread nicks in the ribosomal RNA (rRNA), producing an abundance of short rRNA fragments that copurify with intact ribosomes and comigrate with the ribosome footprints on polyacrylamide gels. Thus, the resulting sequencing data sets largely consist of rRNA sequences (Ingolia et al. 2009), limiting the sequencing depth of useful ribosome footprints.

Some early studies were able to mitigate contamination with rRNA fragments by gel purifying a very tight distribution of ~30 nt RNA fragments (the expected size of a ribosome footprint) to avoid prominent rRNA fragments (Guo et al. 2010), an approach which will not work for all species and samples. Cutting a narrow range of fragments also excludes other lengths of ribosome footprints, which report on the often substantial populations of ribosomes lacking A-site tRNAs (Lareau et al. 2014; Wu et al. 2019), trapped on truncated mRNAs (Guydosh and Green 2014; Arribere and Fire 2018; D’Orazio et al. 2019), or engaged in collisions (Guydosh and Green 2014).

Later studies used subtractive hybridization with biotinylated oligos to deplete the most abundant rRNA fragments (Brar et al. 2012; Ingolia et al. 2012), but variation in sample and library preparation can lead to differences in the number and identity of the most abundant contaminants, often requiring design of new probes with each modification of protocol. Commercial rRNA depletion solutions for RNA-seq are generally designed to work on intact or lightly fragmented rRNA, whose fragments are present in equimolar ratios. These assumptions do not apply to the rRNA fragments in ribosome profiling libraries, which are hugely biased by nuclease digestion and size selection. The standard rRNA depletion reagent for ribosome profiling became Ribo-Zero, which consisted of internally...
biotinylated and fragmented RNA antisense to the entire rRNA (Sooknanan 2009) and was able to deplete a broad range of fragments from ribosome profiling libraries, albeit with varying success (McGlincy and Ingolia 2017).

The recent discontinuation of Ribo-Zero as a standalone product (though currently available for Human/Mouse/Rat as a component of some RNA-seq kits) led us to test several alternative commercial solutions designed for rRNA depletion of human samples. We found that the commercial products that we tested were at best only modestly effective at depleting rRNA from ribosome profiling libraries. More importantly, we found that methods utilizing targeted nucleolytic degradation of rRNA, such as by RNaseH, caused off-target trimming and degradation of ribosome footprints. Broadly speaking, these treatments reduce the fraction of mappable sequences, perturb global gene expression measurements and blur positional information.

RESULTS

In order to identify an alternative rRNA depletion option for ribosome footprint profiling we tested a small number of commercial depletion technologies designed for RNA-seq of human samples, alongside the remainder of our existing Ribo-Zero stock (hereafter referred to as legacy Ribo-Zero). In brief, our protocol (McGlincy and Ingolia 2017; Wu et al. 2019) consists of RNaseI digestion of cellular lysates, followed by pelleting ribosomes over a sucrose cushion and isolation of fragments by PAGE. To prepare them for sequencing (Fig. 1A), these fragments are then ligated to a 3′ preadenylated DNA linker, depleted of rRNA fragments, reverse transcribed, circularized, and PCR amplified for Illumina sequencing. For this comparison, we prepared ribosome footprints from the same HEK293T cell lysate in duplicate, cutting broadly to isolate 15 to 40 nt fragments, which include the standard 30 nt ribosome footprints, as well as the ~15–18 and 21 nt fragments that report on mRNA cleavage (Guydosh and Green 2014) and unoccupied A sites (Wu et al. 2019), respectively. We split these each into four identical groups after 3′ linker ligation (Fig. 1A), to be depleted with Ribo-Zero (Illumina), RiboCop (Lexogen), NEBNext (NEB), or left undepleted. RiboCop is an affinity depletion method that uses complementary biotinylated DNA oligos targeting the rRNA. The NEBNext rRNA depletion kit uses antisense DNA oligos that tile the entire rRNA to target RNaseH-mediated degradation of rRNA. As described above, Ribo-Zero consists of biotiny- lated rRNA fragments complementary to entire rRNAs (Sooknanan 2009).
After sequencing the resulting libraries, we found that the undepleted libraries contained 6.6 ± 0.13% of reads mapping to coding regions (CDS) (Fig. 1B; Supplemental Table S1). Legacy Ribo-Zero increased this to 16.6 ± 0.33%, while RiboCop and NEBNext yielded 9.78 ± 0.12% and 10.79 ± 0.95%, respectively. While all of these approaches increased the number of usable reads, we noticed that the NEBNext kit produced a large fraction of short and unmappable fragments (“<10 nt” and “unmapped” in Fig. 1B). To investigate this further, we looked at the fragment size distribution in these libraries (Fig. 1C). Ribosome footprint profiling in human cells normally produces fragments of ∼30 nt, though the length distribution can be affected by RNaseI digestion conditions such as salt concentration (Ingolia et al. 2012). The undepleted libraries from this experiment have a modal fragment length of 32, with the majority of fragments between 28 and 36 nt in length. The libraries depleted by RiboCop and legacy Ribo-Zero have indistinguishable size distributions; importantly, however, the RNaseH-mediated depletion by the NEBNext protocol resulted in a broad distribution of fragment sizes spanning nearly the entire range of the 15 to 40 nt size selection. These data indicate that substantial off-target RNaseH cleavage has occurred. Since we performed RNA depletion immediately after linker ligation (Fig. 1A), we surmised that this trimming was due to an abundance of leftover DNA linker, which contains six random nucleotides at its 5’ end that can anneal to our ribosome footprints and mediate RNaseH cleavage. To test this hypothesis, we repeated the profiling protocol, this time in K562 cells, with NEBNext depletion prior to linker ligation. We also tested the protocol with two different ranges of gel-purified fragments, 20–34 nt, which correspond to the footprint of a single ribosome, as well as 20–60 nt, which also includes the larger footprints of collided disomes (Guydosh and Green 2014), but which typically results in more rRNA contamination. The resulting depleted libraries contained substantially fewer short and unmappable reads than those produced with depletion post-ligation (compare Fig. 1D to Fig. 1B; Supplemental Table S1), suggesting that many of the short and unmappable fragments in our initial experiment were the result of widespread cleavage mediated by contaminating DNA linker. These data sets reveal a much tighter size distribution than post-ligation depletion, with peaks of ribosome-protected fragments 29 and 21 nt in length (Fig. 1E), which correspond to ribosomes with occupied and unoccupied A sites, respectively (Wu et al. 2019). However, there was still noticeable trimming of the ribosome footprints, particularly the 29 nt fragments, which we presume is off-target activity from the DNA probes in the NEBNext kit.

We next tried three additional commercial rRNA depletion kits. These included a prerelease version of Ribo-Zero plus (Illumina) (a targeted depletion method similar to NEBNext that utilizes a proprietary nuclease), QIAseq FastSelect (Qiagen) (a pool of locked nucleic acids that block reverse transcription), and riboPOOL (siTOOLS biotech) (a cocktail of biotinylated oligos against the rRNA). We performed Ribo-Zero plus before linker ligation, and QIAseq and RiboPOOL depletion immediately before reverse transcription (Fig. 1A). We performed variations of Ribo-Zero plus depletion with supplementary oligos against abundant rRNA fragments (O), as well as supplemented with formamide (F) during probe annealing, to better denature highly structured rRNA fragments. For riboPOOL depletion, we also performed a variation where we omitted the final heating step. None of these methods proved effective at increasing the fraction of CDS-mapping fragments (Fig. 1F; Supplemental Table S1). Although Ribo-Zero plus decreased the fraction of mRNA-mapping reads, it appears that CDS-mapping reads were degraded, and the library contained an abundance of unmappable or intronic reads. These effects are evident in the CDS-mapping footprint length distribution (Fig. 1G), which shows a broadening of the 21 and 29 nt peaks and a visible increase in the fraction of 23–25 nt reads, which likely are degradation products of the longer 28 nt fragments, that correlates with the extent of depletion. This fraction increases from 7% in the undepleted sample to 20% in the Ribo-Zero plus OF library. We interpret this result as indicating that off-target activity is a general feature of nuclease-mediated rRNA depletion.

Since the NEBNext kit was modestly successful at rRNA depletion, we tested if these depleted libraries were still suitable for making gene-level inferences of ribosome footprint density. The RPM (reads per million CDS-mapping reads) values for our undepleted or NEBNext-depleted libraries were reproducible (Pearson $r^2 = 0.900$ and 0.922) between pseudoreplicates (pseudo because we isolated different fragment sizes) despite their modest sequencing depth, but many transcript RPMs differed between the depleted and undepleted libraries (Pearson $r^2 = 0.788$ and 0.780), with a small number of genes showing extreme deviations in RPM (Fig. 2A). These differences indicate that even aggregate gene-level measurements are perturbed by RNaseH-mediated depletion. This perturbation was also seen in the Ribo-Zero plus depleted samples (Fig. 2B), but not in samples depleted with legacy Ribo-Zero or riboCop (Fig. 2C), suggesting that it is a general feature of nuclease-mediated depletion.

An important feature of ribosome profiling data is the nucleotide-resolution positional information, which provides information on ribosome movement at specific codons or groups of codons (Ingolia et al. 2009, 2011; Stadler and Fire 2011; Zinshteyn and Gilbert 2013; Nedialkova and Leidel 2015), as well as the A site status (occupied or empty) of the ribosome, which is inferred from the length of the footprint (Wu et al. 2019). These analyses require the identification of the A, P, and E site locations within each footprint, as well as the length class (∼21, 28, etc.) of that footprint. Site identification is generally accomplished by
averaging reads of a given length relative to all start codons, with the knowledge that ribosomes initiate with the start codon in the P site. Both NEBNext and Ribo-Zero plus depletion blurred the accuracy of site assignment (Fig. 3) as evidenced by reduced peaks and shallower troughs in the average gene plots.

Ribosome profiling libraries with perturbed read lengths could still be usable for positional analysis if the read trimming happens in a defined way. To see how the alterations of read lengths affect the positional resolution of ribosome profiling, we summed the total number of footprint 5′ ends for each read length relative to the start codons of all coding regions (Supplemental Figs. S1, S2). Ribosome profiling libraries generally have a peak of footprint 5′ ends mapping ~12 nt upstream of start codons. In the 5′-aligned footprint heatmaps in Supplemental Figures S1, S2, diagonal lines from the start codon peak are characteristic of footprints from initiating ribosomes that are shortened by nibbling at the 5′ end, while vertical lines indicate variation in the 3′ end, which maintains the position of the 5′ end while shortening the footprint. The presence of both vertical and diagonal lines emanating from the start codon peaks of the NEBNext-depleted libraries (Supplemental Figs. S1B, S2) indicates that the reads are trimmed from both ends. Since it is impossible to determine from which end a particular read is truncated, this trimming is unlikely to be computationally correctable.

In a final experiment, we decided to revisit previously used methods for rRNA depletion based on antisense biotinylated oligos (Ingolia et al. 2012). Based on the sequencing data from our undepleted K562 samples, we designed a set of six biotinylated oligos complementary to 37% of rRNA fragments (Supplemental Table S2). We used these oligos to subtract rRNA fragments from adapter-ligated footprints generated from HEK293T cells. This step increased the fraction of CDS-mapping reads in our library from 5.7% to 8.7%, with little or no perturbation of fragment lengths, positions, or gene-level fragment counts (Fig. 4). These promising results suggest that further efforts for ribosome profiling rRNA depletion should be aimed at optimizing custom oligo-based methods.

**DISCUSSION**

In summary, we have tried several rRNA depletion methods for ribosome profiling. While we do not intend this to be a comprehensive survey of the available options, we found all of the commercially available options that we tested lacking compared to legacy Ribo-Zero. For those with an immediate need some of these kits (most notably RiboCop) can be successfully used to increase usable reads for human samples, but we strongly caution against the use of any methods that use nuclease cleavage, as they blur the detailed mechanistic information that can be gained.
from the lengths of ribosome footprints (Guydosh and Green 2014; Wu et al. 2019). Moreover, most of these reagents are only available for use in a limited number of species. The best way forward for rRNA depletion in ribosome profiling may be the previously used custom oligo depletion method (Ingolia et al. 2012), which is less expensive and easier to optimize. In yeast library preparations, we have had reasonable success using previously published oligo depletion methods (Guydosh and Green 2014), consistently achieving 40% CDS-mapping reads with libraries cut from 20-34 nt (data not shown). However, our results have been less effective for human samples (Fig. 4), indicating that further species-specific and probably sample-specific optimization in either depletion protocols or depletion oligo design is required. Fortuitously, a recently published analytical framework (Ribo-ODDR) for designing depletion oligos has the potential for substantial enrichment of CDS-mapping reads (Alkan et al. 2020). Other groups have had success using double-strand specific DNase, combined with a high-temperature reannealing of the final DNA library that only leaves the most abundant sequences (the rRNA fragments) double-stranded (Chung et al. 2015), or with targeted oligos against cDNA for the most abundant rRNA contaminants (Archer et al. 2014, 2016), but these methods have not to our knowledge been tested on a broad range of fragment lengths, nor with improved library preparation methods such as randomized unique molecular identifiers (McGlincy and Ingolia 2017). We note that enzymatic depletions performed at the cDNA or dsDNA stage have the particular advantage that they cannot change the ends of a cloned fragment, but only completely remove it from the sequenced pool, since removal of the ligated adaptors will prevent sequencing. Recent reports indicate that optimization of RNase digestion conditions (Sharma et al. 2019) as well as judicious choice of RNase enzyme (Gerashchenko and Gladyshev 2017) can also reduce the rRNA content of ribosome profiling libraries, though many procedures using these different nuclease obscure the high-resolution information on ribosome position and A site status. Finally, it is important to note that off-target RNaseH activity will not only affect the conclusions of ribosome footprint profiling experiments, but of any protocol that requires precise knowledge of the 5’ and 3’ ends of RNA fragments.

MATERIALS AND METHODS

Cell culture conditions

Mammalian cell lines were grown in cell culture incubators at 37°C in the presence of 5% CO2. HEK293T cells were cultured in Dulbecco’s modified Eagle medium (DMEM) +10% fetal bovine serum (FBS) (Gibco). K562 cells were grown in Roswell Park Memorial Institute (RPMI) 1640 media + 10% FBS.

Cell lysis and ribosome footprint profiling

Our protocol is adapted from McGlincy and Ingolia (2017). Ten centimeter dishes of adherent HEK293T cells were briefly washed with PBS and lysed by addition of 0.5 mL of mammalian footprint
For each sample, 100 µL of Ribo-Zero magnetic beads were pelleted on a magnetic stand, washed 2× in 100 µL water and resuspended in 30 µL Magnetic Bead Resuspension Solution with 0.5 µL RiboGuard RNase inhibitor. Two microliters of Ribo-Zero reaction buffer and 5 µL Ribo-Zero Removal Solution was added to adapter-ligated RNA in 13 µL water. Solution was incubated in a thermomixer at 68°C for 2 min, followed by 15 min at 37°C with 400 RPM shaking, then 5 min on ice. Full 20 µL of this mixture was added to 30 µL of resuspended magnetic beads, vortexed for 10 sec, and incubated for 15 min in a thermomixer at 25°C. Beads were pelleted on a magnetic stand, and supernatant was kept, and isopropanol precipitated.

rRNA deletion methods overview

For Figure 1A, B, Legacy Ribo-zero (Illumina), RiboCop (Lexogen), and NEBNext (NEB) depletions were performed after linker ligation according to the manufacturer’s recommendations. For Ribo-Zero depletion, we omit the final heating step in the manufacturer’s protocol, as this has been suggested to improve depletion of small fragments (McGlincy and Ingolia 2017). For subsequent figures NEBNext and Ribo-Zero plus (Illumina) depletion were performed right after fragment size selection, while all other methods were performed after linker ligation. These methods were performed according to manufacturer recommendations, except that for some experiments with Ribo-Zero Plus we included 45% formamide in the hybridization reaction (indicated as F), as well as supplementary oligos (indicated as O), provided by Illumina, designed against abundant rRNA contaminants.

rRNA depletion with riboCop

For each sample, 100 µL of Ribo-Zero magnetic beads were pelleted on a magnetic stand, washed 2× in 100 µL water and resuspended in 30 µL Magnetic Bead Resuspension Solution with 0.5 µL RiboGuard RNase inhibitor. Two microliters of Ribo-Zero reaction buffer and 5 µL Ribo-Zero Removal Solution was added to adapter-ligated RNA in 13 µL water. Solution was incubated in a thermomixer at 68°C for 2 min, followed by 15 min at 37°C with 400 RPM shaking, then 5 min on ice. Full 20 µL of this mixture was added to 30 µL of resuspended magnetic beads, vortexed for 10 sec, and incubated for 15 min in a thermomixer at 25°C. Beads were pelleted on a magnetic stand, and supernatant was kept, and isopropanol precipitated.

rRNA depletion with NEBNext rRNA depletion kit (human/mouse/rat)

One microliter of NEBNext rRNA Depletion Solution and 2 µL Probe Hybridization Buffer were added to 12 µL size-selected fragments (or adapter-ligated fragments for Fig. 1A) in a PCR tube and mixed by pipetting. Samples were then annealed to probes by heating to 95°C for 2 min in a thermal cycler with a heated lid set to 105°C, cooled to 22°C at 0.1°C/sec and held at 22°C for 5 min. A mixture of 1 µL water, 2 µL RNase H Reaction buffer and 2 µL NEBNext RNase H was added to the annealed mixture and mixed by pipetting. The mixture was incubated in a thermal cycler with heated lid at 37°C for 30 min, then immediately placed on ice. To this mixture 30 µL of DNase I
digestion master mix (22.5 µL water, 5 µL DNase I reaction Buffer, 2.5 µL DNase I) was added and incubated at 37°C for 30 min, followed by isopropanol precipitation.

**rRNA depletion with Ribo-Zero plus reagent (human/mouse/rat)**

To 3.15 µL of gel purified RNA fragments, we added the following: 4 µL of premixed (3 µL DB1 + 1 µL DP1), 1 µL supplementary probes when indicated, 6.75 µL formamide when indicated, water to 15 µL. Samples were then annealed to probes by heating to 95°C for 2 min in a thermal cycler with a heated lid set to 105°C, and cooled to 37°C at 0.1°C/sec. Annealed samples were spun down and 5 µL of (1 µL RDE + 1 µL RDB) was added, mixed, and incubated in a thermal cycler at 37°C for 15 min to digest RNA. Samples were spun down and 10 µL of (3 µL PRE + 7 µL PRB) was added, mixed, and incubated in a thermal cycler at 37°C for 15 min to digest DNA, and placed on ice, followed by isopropanol precipitation.

**rRNA depletion with QIAseq FastSelect**

Pellets of adaptor-ligated RNA were resuspended in (10 µL water + 4 µL ProScript II buffer + 1 µL Qiagen fastselect [THS-001Z-24]), transferred to a PCR strip tube, and annealed in a thermal cycler with a heated lid for 2 min each at 75°C, 70°C, 65°C, 60°C, 55°C, 37°C, 25°C, 4°C. Reverse transcription was then performed as usual.

**rRNA depletion with riboPOOL (protocol_v1-5)**

Lyophilized riboPOOL (human) was resuspended to 100 µM. Pellets of adaptor-ligated RNA were resuspended in (14 µL water + 5 µL hybridization buffer [10 mM Tris-HCl pH 7.5, 1 mM EDTA, 2 M NaCl] + 1 µL riboPOOL, 1 µL superasin RNase inhibitor [Thermo]). The mixture was heated to 68°C in a thermomixer for 10 min, and the thermomixer was set to 37°C and allowed to slowly cool, allowing probe annealing over 30 min. Washed 80 µL of Dynabeads MyOne Streptavidin C1, 0.1 M NaOH 0.05 M NaCl (Thermo Fisher #65001) 2× in 100 µL (0.1 M NaOH 0.05 M NaCl), then 1× in (0.1 M NaCl), then resuspended in 160 µL (5 mM Tris-HCl [pH 7.5] 0.5 mM EDTA 1 M NaCl) and split into two tubes. Annealed samples were added to first aliquot of beads and incubated at 37°C for 15 min, followed by a 50°C incubation for 5 min (this heating was omitted where indicated). Both bead aliquots were pelleted, supernatant from second aliquot was removed and replaced with supernatant from first aliquot. Beads were incubated once more at 37°C for 15 min, followed by a 50°C incubation for 5 min (omitted where indicated). Beads were pelleted and supernatant transferred to a new tube twice.

**rRNA depletion with custom biotinylated oligos**

To design depletion oligos, we visualized the location of the most abundant contaminating rRNA in the UCSC Genome Browser and ordered 60 nt 5’ biotinylated oligos antisense to the most abundant fragments. An oligo mix solution was made by diluting the six depletion oligos (Supplemental Table S2) in 4× SSC buffer to a concentration of 2.5 µM for each oligo. A total of 10 µL of the oligo mix solution was added to 10 µL of sample solution (ligation products precipitated and resuspended in nuclease-free water). The sample and oligo mixture were heated at 80°C for 2 min in a thermomixer, which was then set to 25°C to gradually cool the mixture and anneal oligos to the sample RNA. During cooling, 150 µL of MyOne Streptavidin C1 dynabeads (ThermoFisher) per depletion were washed thrice with 150 µL of 1× Binding/Washing buffer (5 mM Tris, pH 8.0, 1 M NaCl, 0.5 mM EDTA), and then beads were resuspended in 30 µL 2× Binding/Washing buffer. The cooled sample was added to the prepared beads and incubated at 25°C for 15 min with shaking at 500 rpm in a thermomixer. After incubation, beads were precipitated with a magnetic stand, and supernatant was removed, and isopropanol precipitated to recover RNA.

**Data analysis**

Raw ribosome footprint reads were trimmed of the four random nt from the 5’ end with seqtk (https://github.com/lh3/seqtk), then trimmed of 3’ adaptor sequence (NNNNNCACTCGGGCAC CAAGGAC) using Skewer (Jiang et al. 2014). Reads longer than 10 nt were aligned to PhiX-174, human rRNA and human non-coding RNA sequences using STAR (Dobin et al. 2013) with –outFilterMismatchNmax 2 –outSAMmultNmax 1. Unmapped reads were mapped to the hg38 human genome (GENCODE release 27), using STAR with parameters –alignSJoverhangMin 1 –alignIntronMax 1000000 –alignSjOverhangMin 3 –outFilterType BySJout –outFilterMultiMapNmax 200, –outFilterScoreMin OverLread 0 –outFilterMatchNminOverLread 0 –outFilterMatchNmin 0, –outFilterMismatchNmax 3. To generate a reference transcriptome with a single transcript per gene for alignment, the GENCODE v27 gff file was filtered for those transcripts with an APPRIS score (Rodriguez et al. 2013) between 1 and 4 (inclusive) and then the transcript with the longest CDS (and longest transcript if tied) was chosen. For multiply mapping reads, the primary alignment was used for all analyses. For Figure 1A,C,E, reads were assigned the smallest feature that they overlapped based on the comprehensive GENCODE v27 annotation gff file. For RPM correlation plots, only genes with an average of 64 or more counts and a minimum of 1 RPM between all libraries in the same panel were included. The Python code for the analysis pipeline and for figure generation are available at https://github.com/borisz264/rRNA_depletion_2020. Sequencing data have been deposited in GEO with accession GSE147324.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

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