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Selective recovery of RNAs from bacterial pathogens after their internalization by human host cells

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Highlights

• Method for bacterial RNA recovery from host cells.
• Easy and inexpensive procedure to selectively recover bacterial RNAs from composite cell mix.
• Key influence on beads size and composition during extraction.

Abstract

Selective RNA extractions are required when studying bacterial gene expression within complex mixtures of pathogens and human cells, during adhesion, internalization and survival within the host. New technologies should be developed and implemented to enrich the amount of bacterial RNAs since the majority of RNAs are from the eukaryotic host cells, requiring high read depth coverage to capture the bacterial transcriptomes in dual-RNAseq studies. This will improve our understanding about bacterial adaptation to the host cell defenses, and about how they will adapt to an intracellular life. Here we present an RNA extraction protocol to selectively enrich the lowest bacterial RNA fraction from a mixture of human and bacterial cells, using Zirconium beads, with minimal RNA degradation. Zirconium beads have higher capacity to extract bacterial RNAs than glass beads after pathogen internalization. We optimized the beads size and composition for an optimal bacterial lysis and RNA extraction. The protocol was validated on two human cell lines, differentiated macrophages and osteoblasts, with either Gram-positive (Staphylococcus aureus) or -negative (Salmonella typhimurium) bacteria. Relative to other published protocols, yield of total RNA recovery was significantly improved, while host cell infection was performed with a lower bacterial inoculum. Within the host, bacterial RNA recovery yields were about six-fold lower than an RNA extraction from pure bacteria, but the quality of the RNA recovered was essentially similar. Bacterial RNA recovery was more efficient for S. aureus than for S. typhimurium, probably due to their higher protection by the Gram positive cell walls during the early step of eukaryotic cell lysis. These purified bacterial RNAs allow subsequent genes expression studies in the course of host cell-bacteria interactions.

Key words: Bacterial & prokaryotic RNAs, host-pathogen interactions, Zirconium beads, bacterial internalization, RNA expression, selective RNA purification.
1. Introduction

Transcriptomic studies of internalized bacteria gene expression improve our understanding of bacterial adaptation to host cells defense, and intracellular life [1], [2]. Recent RNA-based technologies allow to tackle gene reprograming by intracellular bacterial pathogens when internalized by eukaryotic host cells [3]. Those dual RNA-seq technologies assess how bacterial pathogens fine-tune gene expression when intracellular. Also, information about how the infected host cells react during infection can be collected. A major technical challenge, however, is the huge excess of eukaryotic over bacterial RNA [4] that needs to be removed to obtain specific and reliable transcriptomic data on pathogen gene expression, when intracellular. To this endeavor, extraction methods have been developed to purify RNAs from prokaryotes or eukaryote organisms with higher efficiency and quality [5], [6]. For bacterial RNA extraction from a homogenous population of bacteria, collecting intact, non-degraded RNAs is relatively easy and include mechanical or enzymatic lysis in denaturing conditions, for cell opening [7], [8], followed by phenol extraction of nucleic acids [9] and DNase treatment.

When performed from complex mixtures of organisms, RNA extractions are complicated. However and surprisingly, very few methods are available to extract bacterial RNAs inside eukaryotic cells during infection [10], [11], [12]. Zirconia beads were previously proposed to purify total RNAs for sRNA discovery in bacteria [13]. These strategies provide low amounts of purified RNAs, insufficient for systematic analyses by qPCR or by Northern of bacterial gene expression inside host cells. These methods involve host cells lysis (phenol/ethanol mix; RLT buffer of Qiagen kit, or ice cold acetone: ethanol mix) to leave bacterial cells intact and subsequent total RNA purification. Also, most methods have an enrichment step of the bacterial RNAs with dedicated commercial kit, which increase the cost of the procedure [11], [12]. For S. aureus RNA extraction, most protocols use manufacturer kits [10] [11]. The protocol from [14] lyses bacteria in the bacteria/cellular debris pellets with Trizol and zirconia-silica beads (100µM diameter) into a high-speed homogenizer, to isolate total RNA. For S. typhimurium, the first described methods used a mix of 0.1% SDS, 1% acidic phenol and 19% ethanol (in water) to lyse infected epithelial cells [15], [16]. Then, S. typhimurium RNA was prepared using a total RNA purification kit (Promega SV total RNA purification kit). A second method used Trizol to extract RNAs, after host cells lysis with PBS- 0.1% Triton X-100 [17].

We developed and report here a novel extraction protocol to selectively enrich the minor RNA fraction from internalized bacteria while removing the majority of the eukaryotic RNA excess, using Zirconium beads. We provide evidences that beads sizes and their compositions during extraction have to be optimized to improve bacterial cell lysis for selective bacterial RNA extraction. Bacterial RNA extractions were performed on two human cell lines upon S. aureus internalization, THP1-differenciated macrophages and Saos-2 osteoblasts. We also tested the new method for THP1-differenced macrophages infected with S. typhimurium. We anticipate our bacterial RNA purification protocol to be
applicable for various bacterial-host cell gene expressions, for subsequent transcriptomic studies.

2. Methods

2.1 Bacteria, human cells and cultures.

RNA extraction of Staphylococcus aureus (Newman) [18] and Salmonella typhimurium strains were selected as Gram-positive and -negative representatives. The two strains were grown at 37°C in Brain Heart Infusion broth (BHI, Oxoid, Dardilly, France) for S. aureus, and in Lysogeny Broth (LB, Sigma, Saint-Louis, USA) containing 300mM NaCl for S. typhimurium.

THP1 human monocytes were from ATCC (Rockville, USA) and maintained at 37°C with 5% CO₂ in RPMI1640 (Thermo Fisher Scientific, Waltham, USA) containing 10% FCS (GE Healthcare Hyclone, Chicago, USA). The THP1 cells were treated with 20ng/ml phorbol 12-myristate 13-acetate (PMA, Sigma, St Louis, USA) for 3 days for differentiation into macrophages. The THP1 monocytes were differentiated into adherent macrophages, as evidenced by light microscopy. Saos-2 human osteoblasts (ATCC, Rockville, USA), are maintained at 37°C with 5% CO₂ in McCoy medium (Thermo Fisher Scientific's, Waltham, USA) containing 15% FCS (GE Healthcare Hyclone, Chicago, USA).

2.2 Bacterial internalization and human cell lysis.

Human THP1-differenciated macrophages and Saos-2 osteoblasts were infected at a multiplicity of infection (MOI) of 1:10 and 1:30, respectively, with either S. aureus or S. typhimurium. Experiments were conducted in 6-wells plates with 5.10⁵ THP1-differenciated macrophages, or 4.10⁵ Saos-2 osteoblasts by well. Internalized assays of S. aureus were carried out for 2 h at 37°C, with 5% CO₂ in RPMI 1640 containing 10% of human SAB for THP1-differenciated macrophages and in McCoy medium with 15% of human SAB for Saos-2 osteoblasts. Internalization was stopped on ice and cells were washed three times with ice-cold PBS. All the non-internalized bacteria were removed by culturing the human cells overnight in a medium supplemented with 50 µg/ml gentamycin (Sigma, St Louis, USA).

Internalized assays of S. typhimurium were carried out for 2 h at 37°C, with 5% CO₂ in RPMI 1640 containing 10% of human SAB for THP1-differenciated macrophages. After washing the cells three times with PBS, all the non-internalized bacteria were removed by culturing the human cells for 90 min in a medium supplemented with 100 µg/ml gentamycin (Sigma, St Louis, USA). Subsequently, for both internalized assays, the medium was removed, cells were washed three times
with PBS and incubated with 500µL of PBS/SDS 1% for 15 min at room temperature for cell lysis. Three or six wells were pooled to increase the amount of RNA extracted for each condition, and then centrifuged. After centrifugation, supernatant containing intracellular material of human cells was discarded.

### 2.3 RNA extraction

A method is described for selective bacterial RNA extraction by phenol/chloroform after human cells lysis and mechanical disruption of the bacterial envelopes with beads. The mechanical lysis probably improves bacterial release from the eukaryotic cells.

#### 2.3.1 The beads

Four different types of beads were tested for their abilities to extract RNAs from internalized bacteria. Glass beads (Sigma Chemical Co., St. Louis, USA), used as references, were compared to four new types of beads (Dutscher, Brumath, France). Table 1 shows their compositions, diameters and densities. The variation of beads composition and density should influence RNA extraction. Also, the variation of beads size should impact bacterial crushing, especially those trapped into the eukaryotic cell debris. Each bead set was washed twice in purified water and then twice in an RNA extraction buffer (SDS 0.5%; 20 mM sodium acetate; 1 mM EDTA pH 5.5), before use.

<table>
<thead>
<tr>
<th>Beads</th>
<th>Compositions</th>
<th>Diameters (µM)</th>
<th>Densities (g/cm³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Zirconium</td>
<td>100</td>
<td>6.5</td>
</tr>
<tr>
<td>2</td>
<td>Zirconium</td>
<td>400</td>
<td>6.5</td>
</tr>
<tr>
<td>3</td>
<td>Steel</td>
<td>200</td>
<td>7.8</td>
</tr>
<tr>
<td>4</td>
<td>Zirconium oxyde</td>
<td>500</td>
<td>5.9</td>
</tr>
<tr>
<td>5</td>
<td>Glass</td>
<td>100</td>
<td>2.5</td>
</tr>
</tbody>
</table>

*Table 1: Tested beads, physical and chemical properties*

#### 2.3.2 RNA extraction from host cell-internalized bacteria.

1. Prepare the Fast Prep tubes (cryotubes): add 250µL beating beads (one of the five from Table 1), 250µL phenol (pH 4) and 250µL of a CHCl₃/Isoamyl alcohol mix (24:1).

2. Mix the bacteria/THP1-differentiated macrophages or the bacteria/Saos-2 osteoblasts lysates from the internalization assays in 500µL of RNA extraction
buffer. Transfer this solution into a cryotube containing the beating beads and lyse into a Fast Prep, 30sec, power 6.5. As negative control, the same experiment was done with a bacteria pellet only, without human cells.

(3) Centrifuge cells at 12000 rpm for 5 min at 4°C, and keep the supernatant.


(5) Centrifuge the mix at 12000 rpm for 5 min at 4°C, and keep the upper phase.


(7) Centrifuge the mix at 12000 rpm for 5 min at 4°C, and keep the upper phase.

(8) Precipitate the RNAs with 0.7 vol of isopropanol and 0.1 vol of sodium acetate 3M (pH 5.2), for 2h at -80°C.

(9) Centrifuge the mix at 12000 rpm for 30min at 4°C, discard the supernatant.

(10) Wash the pellet with 500µL of 70% ethanol.

(11) Centrifuge the RNAs at 12000 rpm for 5 min at 4°C, discard the pellet.

(12) Dry the pellet for 5 min into a Speed Vacuum and dissolve the pellet in RNase-DNase free water.

2.3.3 Human cell RNA extraction.

RNAs from the THP1-differenciated macrophages were extracted by using an RNeasy® Mini Kit (Qiagen, Venlo, Netherland) in accordance with the protocol from the supplier. No gentamycin treatment was applied to this sample, which only serves as an internal control (eukaryotic rRNAs).

2.4 Quality-control assessment of the extracted RNAs.

2.4.1 Gel electrophoresis

(1) Each sample of RNAs extracted from the internalized bacterial, prokaryotic and eukaryotic RNA control was diluted into 10µL final volume, and 2µL of 6X gel loading dye were added (NEB, Evry, France).

(2) Load and migrate the samples for 1h at 90V on 1% agarose gel, supplemented with ethidium bromide.
(3) Visualize the eukaryotic 28S and 18S rRNAs and the prokaryotic 23S and 16S rRNAs by UV transilluminator (Figs 1 and 2).

2.4.2 RNA6000 Nano assays

RNA Nano Chips (Agilent, Santa Clara, USA) were used to quantify RNAs and access RNA Integrity Numbers (RIN, Table 3). The RNA Nano Chips were prepared according to the protocol supplied with the RNA 6000 Nano assay kit (Agilent).

(1) Decontaminate the electrodes with an electrode cleaner chip filled with an RNaseZAP™ RNase decontamination solution (Ambion, Inc. cat. No. 9780) for 1 min and once with an electrode cleaner chip filled with RNase-free water for 10 sec.

(2) The chip was filled with the gel-Dye Mix (Agilent).

(3) 5µL of RNA markers mixture was then applied to each well, together with 1µL of a heat denatured (2 min at 70°C) sample or an RNA ladder (Agilent).

(4) The chip was vortexed and run on the Agilent 2100 Bioanalyzer.

2.5 Assessing the amount and ratio of prokaryotic versus eukaryotic RNAs.

The amount and proportion of eukaryotic versus prokaryotic RNAs in either the THP1-differenciated macrophages or Saos-2 osteoblasts host cells with internalized bacteria was independently monitored by (i) RT-qPCR, (ii) gel electrophoresis and (iii) bioanalyzer.

(1) Remove putative DNA remnants from the extracted RNAs using amplification-grade DNase I (Life Technologies, Carlsbad, USA).

(2) Synthesize cDNAs with a High-Capacity cDNA Reverse Transcriptase Kit (Life Technologies, Carlsbad, USA).

(3) Perform qPCR using a Real Master Mix SYBR Kit (5’ PRIME, Life Technologies, Carlsbad, USA) on a StepOnePlus Real-Time PCR system (GE Healthcare, Saint Aubin, France). The DNA Primers are listed on Table 2.
**Table 2:** Sequences of the DNA primers used to detect bacterial and host cells RNAs by qPCR.

For the gene expression studies targeting *invA* mRNA, *tmRNA*, 18S rRNA and *gapdh* mRNA, a Ct ‘cut-off value’ of 35 was chosen because obtaining a 3-cycle difference in target Ct, in comparison to a negative control Ct, except for the 16S rRNA expression where the Ct ‘cut-off value’ of 28 was selected. Before scoring any reaction, optimized qPCR conditions and melting curves were conducted by verifying that the Ct values and amplification curves were both acceptable. As the assay’s thermal cycling protocol involved 40 cycles, Ct 35 was the latest Ct value where the above requirements were acceptable [19].

### 3. Results and discussion.

After human cell internalization, bacterial RNA extraction allows studying bacterial gene expression within various host cells. Here we propose an easy and inexpensive procedure to increase bacterial RNA recovery while removing most of the eukaryotic RNA excess after host cell uptake. The technology was tested on two human cell types infected by either Gram-positive or -negative bacteria, used as model organisms. Various types of beating beads were tested to improve the purification of prokaryotic RNAs, especially those protected and trapped within the eukaryotic cells debris.
3.1 Beads testing and selection.

In this proposed method, *S. aureus* strain Newman was internalized by two different human cell lines, THP1-differentiated macrophages and Saos-2 osteoblasts. These uptakes were followed by cells lysis and total RNA extraction by using five different types of beads (Table 1). As shown in Figure 1, two samples were compared for each type of beads. Extraction controls were achieved with non-internalized bacteria (3.10^7) compared to macrophages-internalized bacteria (3.10^6 human cells with 3.10^7 bacteria; MOI 1:10). In Figure 1, we see two bands correspond to the 23S and 16S rRNAs extracted from the prokaryotic controls, as well as two bands corresponding to the 28S and 18S rRNAs extracted from the eukaryotic control. For RNAs extracted from internalized bacteria with beads 1 or 4, we detect the intact bacterial rRNAs, with no or low amounts of human rRNAs. However, beads 2, 3 and 5 are inappropriate for bacterial RNA recovery after host cell internalization (Figure 1). Beads 1 were selected based on the amount of RNAs collected estimated by (i) Nanodrop dosage, (ii) visual integrity on agarose gel and also according to the beads diameter that is identical to that of glass beads, adapted for bacterial lysis (Table 3 and Figure 1).

<table>
<thead>
<tr>
<th>Beads</th>
<th>Yields of prokaryotic control RNA recovery</th>
<th>Yields of total RNA recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beads 1</td>
<td>100µm Zirconium beads</td>
<td>35-40µg</td>
</tr>
<tr>
<td>Beads 2</td>
<td>400µm Zirconium beads</td>
<td>10-12µg</td>
</tr>
<tr>
<td>Beads 3</td>
<td>200µm Steel beads</td>
<td>24-40µg</td>
</tr>
<tr>
<td>Beads 4</td>
<td>500µm Zirconium Oxide beads</td>
<td>25-30µg</td>
</tr>
<tr>
<td>Beads 5</td>
<td>100µm Glass beads</td>
<td>20-40µg</td>
</tr>
</tbody>
</table>

Table 3. Total RNA recovery from bacteria internalized into human cells and the respective controls testing various beads for cell crushing. The experiment is the result of pooling 6 wells which correspond to 3.10^6 human cells and 3.10^7 bacteria. The RNA dosage were done by Nanodrop.
3.2 Comparison between Glass and Zirconium beads.

To investigate the efficiency of the zirconium beads compared to the glass beads, we performed the RNA extraction in parallel to compare these two types of beads. The extracted RNAs were analyzed by an electrophoresis on agarose gel (Figure 2). For the ‘non-infected macrophage’ control, an extraction was performed with RNeasy® Mini Kit (Qiagen, Venlo, Netherlands).

First, for the *S. aureus* prokaryotic control (Newman strain), two bands with no degradation correspond to the 23S and the 16S rRNAs, and attest RNA integrity (Figure 2A) with both the glass and zirconium beads. Second, for the non-infected conditions (THP1-differenciated macrophages), two bands are detected corresponding to the 28S and the 18S rRNAs, attesting the efficiency of intact eukaryotic RNAs recovery with the Qiagen kit (Figure 2A). This sample serves as positive control. For the internalized bacteria, we detected differences between the THP1 and Saos-2 host cells. For the macrophages-internalized bacteria, we purified intact bacterial RNAs with the Zirconium beads, but not with the glass beads (Figure 2A). For the Saos-2-internalized bacteria, however, we collected both the bacterial and eukaryotic rRNAs with the zirconium beads (Figure 2A). Our hypothesis is that during THP1 cells lysis, most of the eukaryotic RNAs are released and discarded since they locate in the supernatants after centrifugation, whereas the prokaryotic RNAs is probably sheltered inside bacteria (pellet). For the Saos-
2 cells, their lysis may be less effective, with a lower release and elimination of their (eukaryotic) RNAs.

For the RNAs extracted from Gram-negative *S. typhimurium* (control), we detected three bands corresponding to the 23S rRNA, which is cleaved in two smaller RNA fragments of 1.1 and 1.7kb [20], together with full-length 16S rRNAs for both the glass and zirconium beads, validating the control (Figure 2B). We used the same sample from Figure 2A for a eukaryotic control. For the internalized bacteria, we visualize a lower quantity of RNAs than for *S. aureus*, as well as more contaminating eukaryotic RNAs (Table 4 and 5). There is a higher yield of RNA recovery with the zirconium beads than for glass beads. The extraction is less effective for the Gram-negative bacteria than for the Gram-positive.

In conclusion, zirconium beads have higher capacity to extract bacterial RNAs than glass beads after pathogen internalization. This works better for Gram-positive (*S. aureus*) in comparison to Gram-negative bacteria (*S. typhimurium*). We assume that the improvement of extraction was due to the elevated density of zirconium (d=6.5) in comparison to glass (d=2.5). This characteristic may allow efficient bacteria crushing after their internalization by both THP1 and Saos-2 cells.

<table>
<thead>
<tr>
<th></th>
<th>RNA concentration (ng/µL)</th>
<th>Total RNA recovered (µg)</th>
<th>Loaded RNA (ng)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Prokaryotic control</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td>Glass</td>
<td>3250</td>
<td>97</td>
</tr>
<tr>
<td></td>
<td>Zirconium</td>
<td>6900</td>
<td>207</td>
</tr>
<tr>
<td><strong>Prokaryotic control</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>S. typhimurium</em></td>
<td>Glass</td>
<td>1350</td>
<td>21.6</td>
</tr>
<tr>
<td></td>
<td>Zirconium</td>
<td>1650</td>
<td>26.4</td>
</tr>
<tr>
<td><strong>Eukaryotic control</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>THP1 macrophages</td>
<td>Qiagen</td>
<td>716</td>
<td>21.5</td>
</tr>
<tr>
<td><strong>THP1-internalized</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><em>S. aureus</em></td>
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<td>0.06</td>
</tr>
<tr>
<td></td>
<td>Zirconium</td>
<td>1070</td>
<td>17</td>
</tr>
<tr>
<td><strong>Saos-2-internalized</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td>Glass</td>
<td>26</td>
<td>0.5</td>
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<td></td>
<td>Zirconium</td>
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<td>1.25</td>
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<tr>
<td><strong>THP1-internalized</strong></td>
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<tr>
<td><em>S. typhimurium</em></td>
<td>Glass</td>
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<td>190</td>
</tr>
<tr>
<td></td>
<td>Zirconium</td>
<td>128</td>
<td>1540</td>
</tr>
</tbody>
</table>

Table 4. Comparing total RNA recovery (Nanodrop dosage) between glass and Zirconium beads. The amounts of loaded RNA correspond to the data presented in Figure 2.
Figure 2: Monitoring the extracted RNAs from internalized bacteria by agarose gel electrophoresis stained with ethidium bromide. 23S and 16S rRNAs reveal the presence of prokaryotic RNAs. 28S and 18S rRNAs are the contaminating eukaryotic RNAs. The eukaryotic control represents RNAs extracted from THP1-differenciated macrophages cells. A, prokaryotic controls represent RNAs extracted from S. aureus Newman cells. B, prokaryotic controls represent RNAs extracted from S. typhimurium cells. The arrow points to the bacterial rRNA. The amounts of RNA loaded are indicated onto Table 4.

3.3 Quality control of the extracted RNAs.

Next, we performed a RNA chip to monitor the quality of the extracted RNAs after THP1 uptake (Table 5). The chip allows quantifying the extracted RNAs to assess their integrity. For each condition (prokaryote control, Saos-2 internalized bacteria and THP1 internalized bacteria), higher amount of RNAs was collected with the zirconium beads compared to the glass beads (Table 5). Also, the RNAs extracted from THP1 internalized S aureus with zirconium beads show a RNA Integrity Number (RIN) of 8.7, compared with the prokaryotic controls (RIN of 9.6). That RIN validated the integrity of the extracted RNAs, except for a slight degradation of 23S rRNA for the internalized S. aureus (Supplemental data Figure A.1). For the Saos-2 internalized bacteria, with the RNAs extracted by the Zirconium beads, we obtained a mixture of eukaryotic and prokaryotic RNAs. On the electrophoregram of the extracted RNAs from the Saos-2 internalized bacteria (Supplemental data Figure A.2), 4 peaks correspond to the eukaryotic and prokaryotic rRNAs, and one peak may correspond to a 23S rRNA degradation, as for the THP1 internalized bacteria. This electrophoregram attests of the integrity of the extracted RNAs, with each peak associated to an rRNA (Supplemental data Figure A.2). A summary of each extracts dropped on the chip is represented on Figure B.1 (Supplemental data).

We realized a second RNA chip to get information about the quality of our S. typhimurium extracts. The RIN of both glass and zirconium beads are 6.4 and 6.2 respectively. These values are lower than for S. aureus and can be due to the 23S rRNA cleavage [20] that we discussed earlier. A profile similar to that obtained by [15] was observed, with three bands. One of them corresponding to the 16S rRNA and the two others to 23S rRNA fragments (Supplemental data B.2). For the THP1-internalized S. typhimurium, a mix of eukaryotic and prokaryotic RNAs is detected (RIN of 5.2), which can be consider as acceptable in comparison to prokaryotic control RIN. On the electrophoregram (Supplemental data A.3), we could detect easily the prokaryotic rRNAs and validate their presence in our extracts.
Table 5. Qualitative and quantitative analyses of the extracted RNAs from *S. aureus* Newman or *S. typhimurium* uptake by human cells. The experiment results in pooling 6 wells, which correspond to $3 \times 10^6$ human cells and $3 \times 10^7$ bacteria. RIN: N/A = Not Attributed, ‘1’ corresponds to degraded RNAs, and ‘10’ to intact RNAs. The bacterial inoculum was 50µl ($5 \times 10^6$ CFU) per well of 2ml, corresponding to a total inoculum of 300µl.

### 3.4 Assessing eukaryotic and prokaryotic RNA expression from the extracts.

Gene expression studies were performed on total RNAs extracted with the 100µM diameter Zirconium beads. The expression levels of selected RNA genes were monitored by RT-qPCR in the RNAs extracted from THP1 internalized *S. aureus* Newman and *S. typhimurium*. The expression of several representative genes was measured: *gapdH* mRNA and 18S rRNA were recorded from RNAs extracted from THP1-differentiated cells, *tmRNA* from RNAs extracted from *S. aureus* Newman, *invA* from RNAs extracted from *S. typhimurium*, and 16S rRNA (extracted from both *S. aureus* and *S. typhimurium*) expression levels.

First, we studied the expression of 16S rRNA, a common gene between *S. aureus* and *S. typhimurium*. We designed a set of primers that could work on both bacteria, amplifying a conserved region of the RNA.

The Ct value of a *S. aureus* RNA control is lower than that of a *S. typhimurium* control, when monitoring 16s RNA expression (Fig 3A). This difference between the two strains is also detected when comparing the values obtained when intracellular. The Ct value of
16S rRNA detection increases for both bacteria when intracellular, as expected. We also monitored RNA expression for a gene specific for each bacterium. For *S. aureus*, we monitored *tmRNA* expression (Fig. 3B), which was reported as unfluctuating before and after *S. aureus* uptake [21], that is also observed here. For *S. typhimurium*, genus-specific *invA* mRNA was selected and was detected when the bacteria are internalized (Figure 3C). These expression data are in agreement with both the agarose gel and the chip experiments. Overall, our RNA extraction method is efficient for Gram-positive bacteria during host cell uptake but, to a lesser extent, for a Gram-negative bacterium. Probably the thicker, solid envelope of *S. aureus* better protects the RNA during the initial lysis steps of the eukaryotic cells than for *S. typhimurium*.

**Figure 3:** RNA expression levels of selected prokaryotic genes in internalized bacteria versus non-internalized bacteria by RT-qPCR. The primer set to amplify the 16S rRNA are identical for both bacteria (A), *tmRNA* is specific to *S. aureus* (B) and *InvA mRNA* is specific to *S. typhimurium* (C).
Monitoring the RNA expression levels of selected eukaryotic genes inform us about the amount of eukaryotic RNAs still present in the RNA extracts from internalized bacteria, compare with a non-infected ‘macrophages only’, sample. First, in Figure 4A, we studied 18S rRNA expression which is highly expressed (a Ct value of 7.6 in macrophages). In comparison, the Ct values of internalized *S. aureus* and *S. typhimurium* are higher (15.8 and 10.9) respectively. It indicates that the amount of eukaryotic rRNA was reduced in our internalized sample, with a less efficiency for *S. typhimurium*. For mRNA, we studied expression of *gapdh* mRNA and obtained the same profile than for the 18S rRNA (Fig 4B).

**Figure 4:** RNA expression levels of selected eukaryotic genes in internalized bacteria samples versus non-infected macrophages, obtained by RT-qPCR. 18S rRNA (A) was picked for its high expression and *gapdh* mRNA (B) since it is a commonly used eukaryotic referent gene [22].

As described in the introduction, the majority of RNA extraction methods, after host cells uptake of bacteria, use commercial kits for bacterial RNA recovery needed for subsequent analyses. In this report, we describe a simple and inexpensive method to get the same result. A PBS-1% SDS host cell lysis followed by zirconium beads beating and phenol extraction of RNAs. This method is simple and inexpensive. That method, however, will need optimization when modifying the host cells and bacteria. The method is more efficient on Gram positive bacteria, probably due to different cell walls. Gram negative bacteria cell walls are more fragile than Gram-positive bacteria, with premature lysis of *S. typhimurium* during the extraction leading to premature prokaryotic RNA elimination. A lighter bacterial lysis (using Triton X100), may improve bacterial RNA recovery from *S. typhimurium*. *S. aureus* RNA extraction is more efficient after their uptake by THP1-differenciated macrophages than by Saos-2-osteoblasts. Indeed, RNAs extracted from Saos-2 internalization contain a significant fraction of eukaryotic RNAs and need further optimization.
**Concluding remarks**

The data presented in this report provide a simplified and inexpensive RNA extraction procedure for scientists working on ‘host cell-bacteria’ interactions that expect to enhance bacterial RNA recovery over the eukaryotic RNA excess from the host cells. This method allows genome-wide expression study of the bacterial RNome during host uptake, without the need to use expensive kits removing the bulk of eukaryotic RNAs. The ‘proof of concept’ was obtained on a Gram-positive and negative bacterial pathogens used as model organisms, in two different human host cells. 100µM-diameter Zirconium beads are optimal to get reproducible, clean and intact bacterial RNA extractions from that initial heterogeneous cell mixture. Our protocol is improved from the others because we first use SDS to lyse most of the eukaryotic cells, and then ‘high density’ Zirconia beads to destruct the bacteria cells embedded within the human cell debris. Therefore, we removed most of the eukaryotic RNAs whereas extracting efficiently the bacterial RNAs from the internalized cells. The amount of eukaryotic mRNAs recovered is low, probably because they are degraded and discarded during the initial lysis step. Eukaryotic rRNAs, however, are detected but their quantity is acceptable for subsequent bacterial gene expression studies.

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**References.**


Supplemental Data.

Supplemental Figure A.1: Electrophoregram of the extracted RNAs after S. aureus internalized by THP1-macrophage cells. a =16S rRNA,  b = 23S rRNA degradation,  c= 23S rRNA.
Supplemental Figure A.2: Electrophoregram of the extracted RNAs after *S. aureus* internalization by Saos-2-osteoblast cells.

a = 16S rRNA, b = 18S rRNA, c = 23S rRNA degradation, d = 23S rRNA, e = 28S rRNA.

Supplemental Figure A.3: Electrophoregram of the extracted RNAs after *S. typhimurium* internalization by THP1-macrophage cells.

a = 23S rRNA fragment (1.1kb), b = 16S rRNA, c = 23S rRNA larger fragment (1.7kb), d = 18S rRNA, e = 28S rRNA.
Supplemental Figure B.1: Summary of the RNA chip with *S. aureus* samples

1 = *S. aureus* control extract with glass beads,
2 = *S. aureus* control extract with zirconium beads,
3 = non-infected macrophages extract with Qiagen kit,
4 = macrophages-internalized *S. aureus* extract with glass beads,
5 = macrophages-internalized *S. aureus* extract with zirconium beads,
6 = Saos-2-internalized *S. aureus* extract with glass beads,
7 = Saos-2-internalized *S. aureus* extract with zirconium beads.
**Supplemental Figure B.2:** Summary of the RNA chip with *S. typhimurium* samples.

1 = *S. typhimurium* control extract with glass beads,
2 = *S. typhimurium* control extract with zirconium beads,
3 = macrophages-internalized *S. typhimurium* extract with glass beads,
4 = macrophages-internalized *S. typhimurium* extract with zirconium beads,
5 = non-infected macrophages extract with Qiagen kit.