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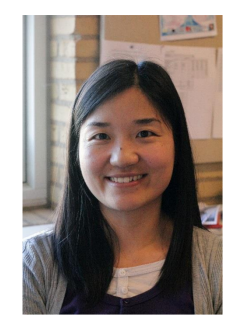
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# **RNA-seq profiling of pathogens in prosthetic joint infection**



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### Introduction

Prosthetic joint infections (PJIs) are serious complications of joint alloplasties. These device-associated infections are typically caused by bacteria growing in biofilms, which protect them from antimicrobial agents and host immune responses in poorly vascularized joints. Our current knowledge of how pathogens cope with the complex conditions within the host is based on *in vitro* studies and animal models, which can differ substantially from their behavior in the human host. Studying gene expression of pathogenesis. RNA sequencing (RNA-seq) is capable of characterizing the entire transcriptome, both quantitatively and qualitatively, of an organism.

### Conclusions

This study indicates that RNA-seq is a challenging but powerful tool to profile the gene expression of pathogens *in vivo*.

#### Aim

To study the gene expression of pathogens directly in PJI in a human

## Results

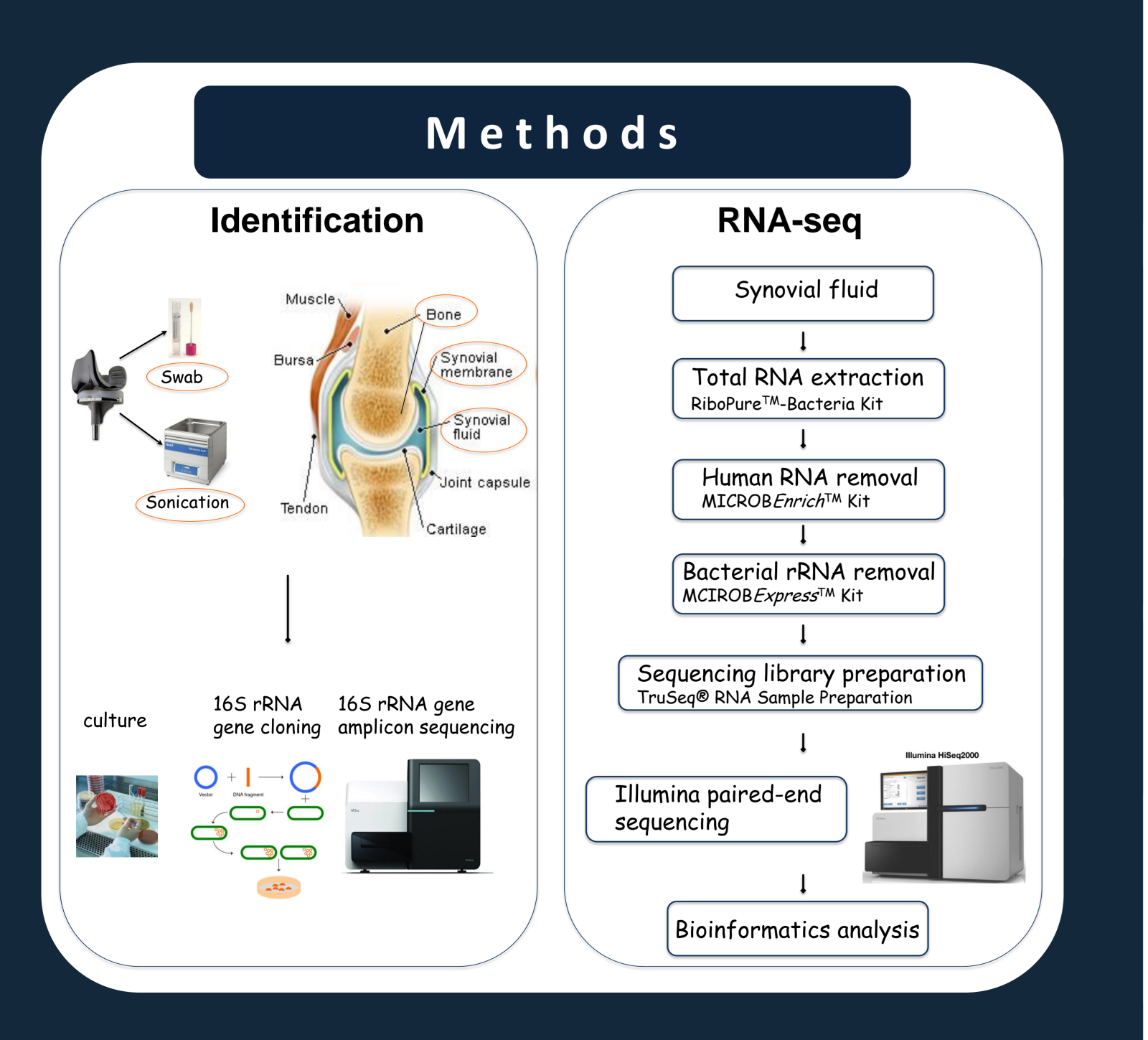
#### Identification

Monomicrobial infection: *Staphylococcus aureus* was identified by 3 independent methods (culture, 16S rRNA gene cloning, and 16S rRNA gene amplicon sequencing).

#### **Mapping statistics**

Number of reade Number of paired reade

# host using RNA-seq.



	Number of reads	Number of paired reads
Total reads	23.000.000	11.500.000
S. aureus reads <sup>1</sup>	270.000 (1.17%)	113.000 (0.98%)
<i>S. aureus</i> mRNA reads	17.000 (0.074%)	7.000 (0.061%)

<sup>1</sup>Reference genome: *S. aureus* N315, 2.84 Mb, 2694 genes, 2613 proteins

#### Top 100 genes (ordered by RPKM)

- 52 Essential genes
- 22 Hypothetical proteins
- 8 Infection related genes

#### Virulence factors:

- hlgA, hlgB and hlgC : form pores in immune cells
- **Sbi**: interferes with the host complement system

#### Virulence regulation genes: saeS and saeR Agr quorum sensing signal $\rightarrow$ $\stackrel{+P}{\operatorname{SaeR}} \rightarrow$ Virulence genes S. aureus cell

membrane

### Acknowledgement

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