PREOMICS

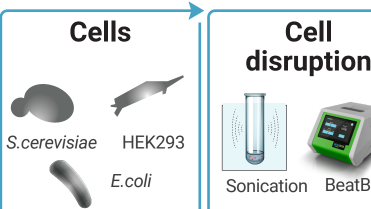
Efficient, reproducible, high-throughput lysis of cells on the BeatBox platform for protein analysis in 96-well plates

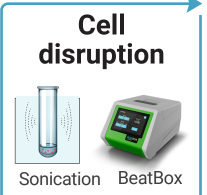
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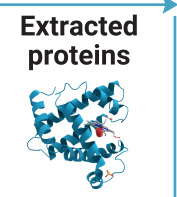
Spotlights

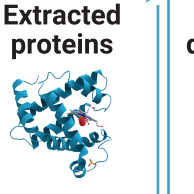
- Up to 96 samples homogenized in 10 min¹
- Small footprint, quiet operation at a steady temperature for the modern lab
- From eukaryotic to prokaryotic cell types
- From thousands to ten million of cells per well
- Seamless integration with iST workflows²

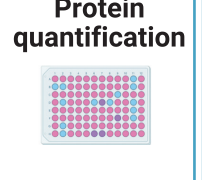
Materials & Methods

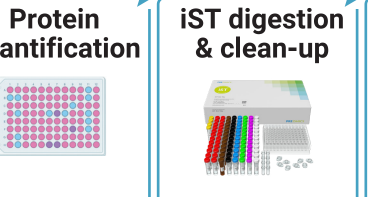










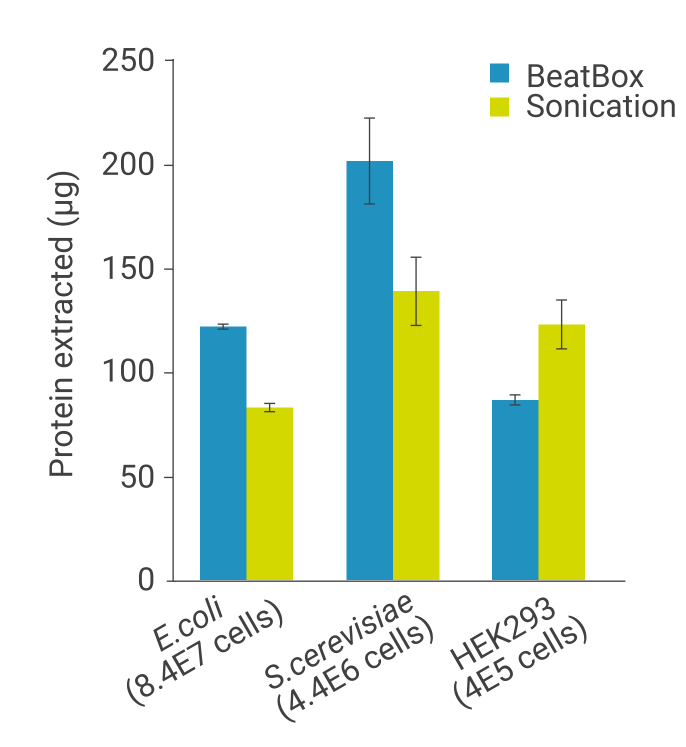


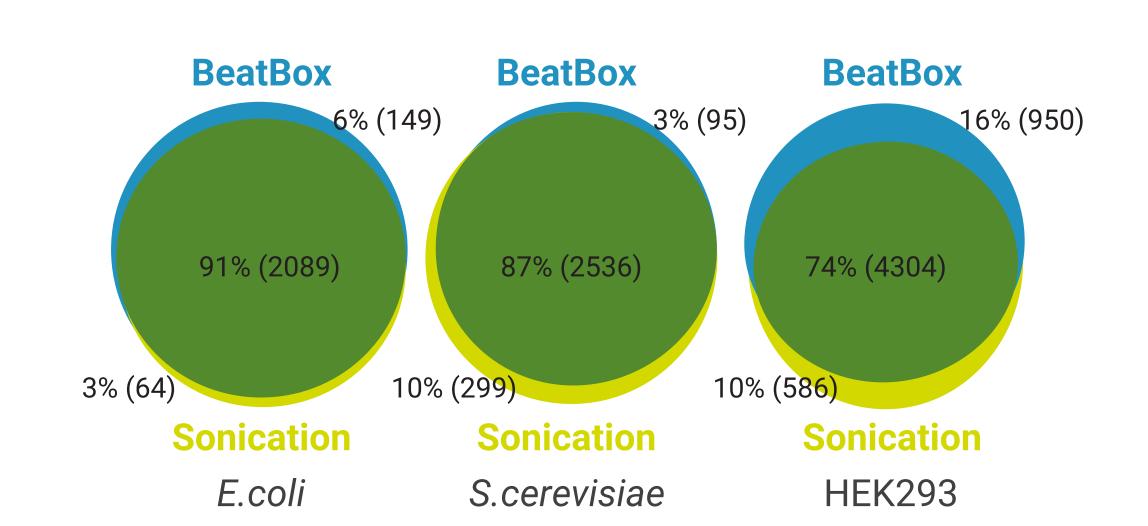




- Input: S.cerevisiae (4.4E6 cells), E.coli (8.4E7 cells) and HEK293 (4E5 cells)
- **Sonication**: Boiling step (95 °C, 10 min) followed by 10 sonication cycles, 30 sec on, 30 sec off in iST LYSE buffer
- BeatBox: No boiling step except for yeast (95 °C, 10 min), followed by 10min homogenisation with standard power setting in iST LYSE buffer
- Sample digestion/peptide clean-up: iST workflow
- Protein assay: Micro BCA™ Protein Assay Kit (ThermoFisher Scientific)
- MS and data analysis: EASY-nLC™ 1200 TimsTOF Pro, MaxQuant (v 2.0.1.0), STRING (v11.5)

Results & Discussion





Shared GO **BeatBox specific GO Sonication specific GO** Intracellular Organelle membrane Catalytic complex Intracellular organelle Microtubule cytoskeleton Autophagosome Microtubule organizing center Organelle Intracellular membrane-bounded Centrosom Membrane-bounded organelle TRAPP complex Cytoplasm Endosome Nucleus Ubiquitin ligase complex Nucleoplasn Cytosol Endomembrane system Bounding membrane of organelle Nuclear lumen Organelle envelope Protein-containing complex Cellular anatomical entity Vesicle tethering complex Transferase complex Lysosomal membrane Intracellular organelle lumen

Protein yields

- BeatBox significantly improves protein yields for E.coli and S.cerevisiae
- For HEK cells, BeatBox provided better protein IDs despite lower protein yields

Protein identification rate

- Similar protein groups IDs were obtained for each species
- HEK293 cell line shows a higher difference between extracted proteins by BeatBox and the sonication technique
- CVs below 3% for protein groups and peptides IDs

Specific Gene Ontology Cellular Component

 HEK293 cell line lysed by BeatBox show solid improvements in the enrichment of membrane-related protein groups

Conclusions

- BeatBox provides high-throughput, efficient, and reproducible lysis across various cell types
- Similar and in-depth proteomic data were obtained using both homogenization techniques
- Equal protein IDs and superb reproducibility were achieved with 4-time less cells in the sample
- Enrichment of cellular component protein groups was measured in BeatBox lysed cells.

References

¹Technical note: BeatBox: Tissue homogenization simplified (www.preomics.com/resources)

²Application note: High-throughput homogenization technique for deeper analysis of multiple mouse tissue proteomes (www.preomics.com/resources)



