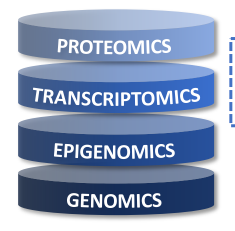
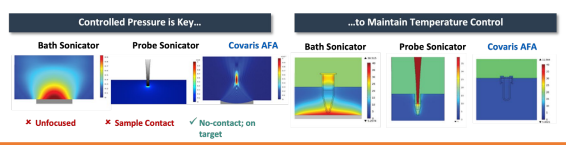
MultiOmics Enabled by AFA® Technology



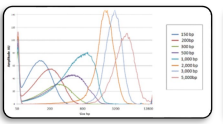
Connecting genotype to phenotype means combining DNA, epigenetics, RNA, protein, or other molecular measurements into a full cellular readout provides researchers with novel scientific insights that cannot be found from single omic methods alone.

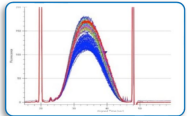
Bigger picture with MultiOmics—MultiOmics goes beyond the genome to unlock deeper biological insights. Using every piece of molecular data available can accelerate biological discoveries and transform our understanding of human health.

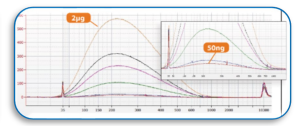


AFA employs highly controlled bursts of focused high-frequency acoustic energy to efficiently and reproducibly process samples in a temperature-controlled and non-contact environment.

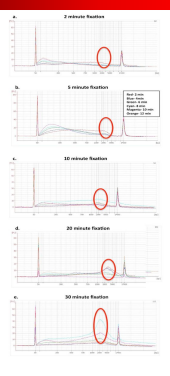
Gold Standard for DNA Shearing Enabling Comprehensive Genomic Profiling







Chromatin Shearing with AFA



Proteomics Sample

- Sample preparation in proteomics largely relies on manual handling Seamless integration of these steps into an comated process (autoSP3 + AFA) noves shortcomings ucibility across different tiss High reprod types and steps Handle detergents, including SDS, adds great flexibility Efficient lysis of fresh and FFPE tissue (1 and 4% SDS) without the need for further peptide clean-up before LC-MS Fast workflow (3.5 h for 96 samples (1 h for ultrasonication, 2.5 h for autoSP3)) up Allows for processing 2-3 plates (or up 300 samples) per working day with m hands-on time

Metabolomics Sample

Combining Proteome and Metabolome Analysis in Plasma/Serum bolomic and proteomic analyses of human plasma and n samples harbor the power to advance our geted metabolic profiling (497 metabolites) and teomics (572 proteins) on the same samples re e and temperature as the strongest influencers Rapid handling and low temperatures (4°C) are imp 2] 1

Monk et al, Cell Systems, 2016, 3(3), 238-251; Davis et al, Communications Biology, 2022, 5, 1301; Rasmussen et al, https://orcid.org/0000-0002-7710-8912; Cyupers et al, Infection, Genetics, and Evolution, 2018, 62, 170-178; Covaris Application Note M020016; Hagen M. Gregner et al, bioRxiv, 2022; Torsten Müller et al, Molecular Systems Biology, 2020,