




A Sysmex Group Company



SureSeq™ for Research Use Only (RUO*)



NGS products for
haematological cancers
and solid tumours

An expanding portfolio of NGS panels for research into haematological and solid tumour cancers, as well as library preparation kits for the accurate detection of a wide range of genetic aberrations.

SureSeq™ NGS panels have been designed in collaboration with recognised cancer experts to detect key aberrations implicated in a wide range of haematological and solid tumour cancers. More so, you can always modify each panel to what's relevant to your research with SureSeq™ myPanel, our regularly updated, expert-curated library of pre-optimised cancer content. Simply mix and match the gene, exonic or intronic content you need to create an NGS cancer panel that meets your exact requirements.

Our unique panel design coupled with hybridisation-based enrichment offers unparalleled coverage completeness and uniformity, allowing accurate detection of low-frequency SNVs and indels, as well as structural aberrations such as ITDs, PTDs, CNVs, LOH and translocations. Combined with our various Universal NGS Complete workflow products, SureSeq™ alleviates the burden of running multiple assays and streamlines your research, delivering comprehensive results using a single NGS workflow.







Interpret, OGT's powerful and easy-to-use NGS analysis solution, provides effortless translation of all your NGS data into meaningful results and is complimentary with all SureSeq™ NGS panels.

* Not for use in diagnostic procedures.







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Discover library preparation kits and Interpret NGS analysis software for the accurate detection of a wide range of genetic aberrations

SureSeq™ haematological panels


| | | |
|--|--|--|
| CLL + CNV panel  | Core MPN panel  | Pan-Myeloid panel  |
| Custom AML panel  | Germline Breast Cancer + CNV panel  | Myeloid Plus panel  |

SureSeq™ solid tumour panels

| | | |
|--|--|--|
| Custom Breast Cancer panel  | Breast Cancer + CNV panel  | Custom Colorectal Cancer panel  |
| Custom Melanoma panel  | Custom Prostate Cancer panel  | Custom Cancer panel  |

Interpret software

Interpret NGS analysis software



Library preparation products

| | |
|---|---|
| Universal NGS Complete workflow solution  | FFPE DNA Repair Mix*  |
|---|---|

Last update August 2022

*The SureSeq™ FFPE DNA Repair Mix can only be purchased in conjunction with SureSeq™ NGS panels, not as a standalone product.

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