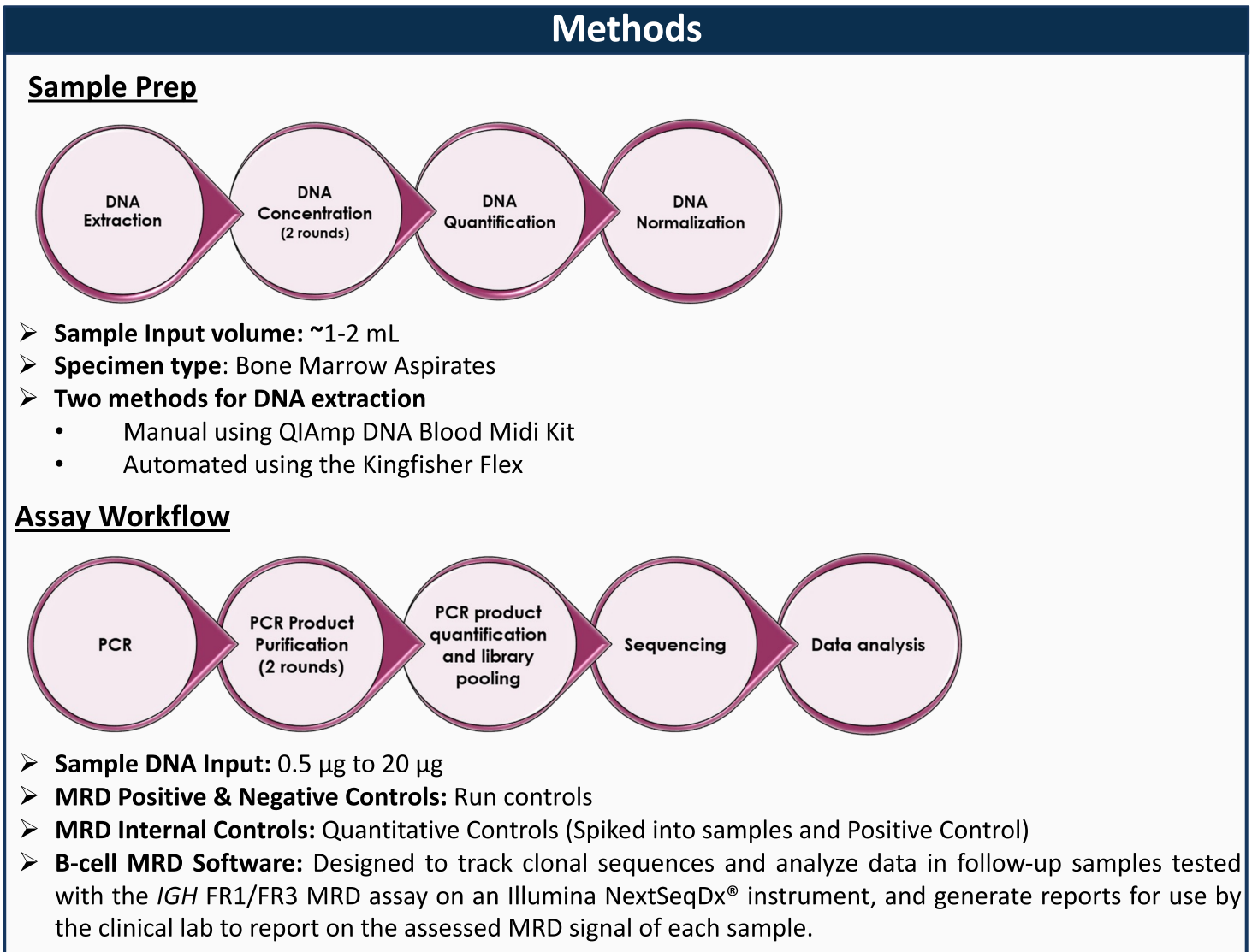
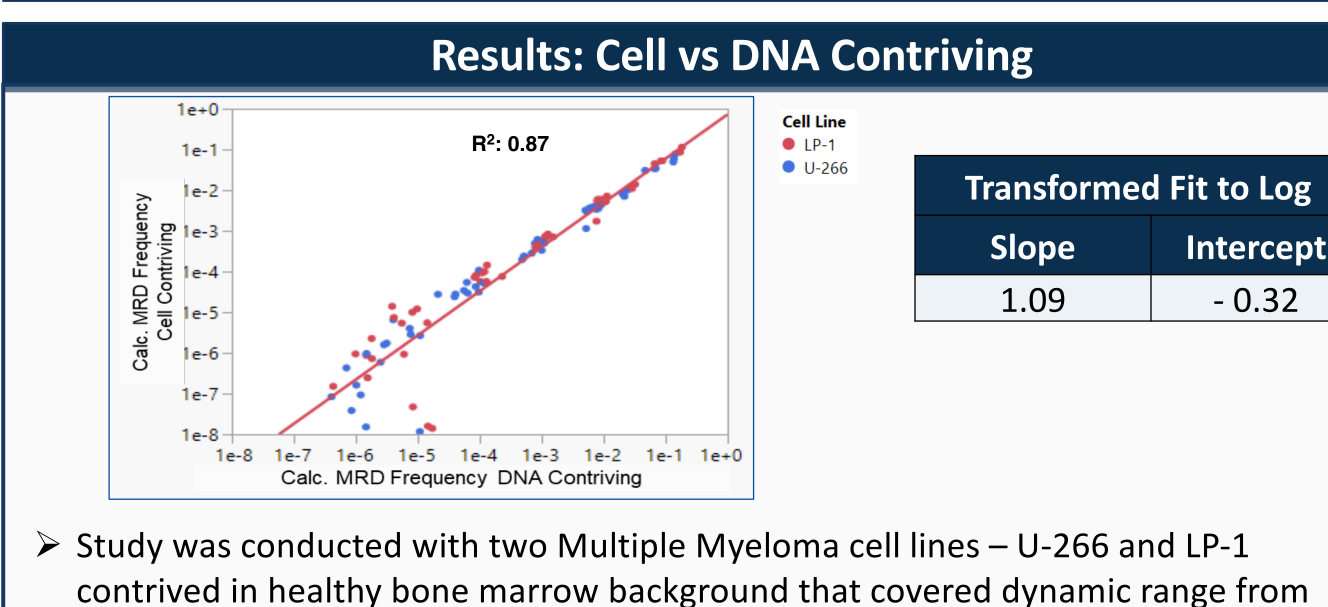
### Validate an NGS Based Measurable Residual Disease (MRD) Assay for B-Cell Lymphoproliferative Diseases

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#### Background

MRD assessment in B-cell lymphoproliferative diseases has proven utility in assessing response to therapy, refining treatment and predicting clinical outcome in patients previously treated with a combination of treatments (Garcia-Marco et al. Haematologica 2019; Pui et al. Leukemia 2017). Next-generation sequencing (NGS) based approaches for MRD detection have provided high sensitivity and specificity to identify and track measurable residual disease using clonal immunoglobulin heavy chain (IGH) gene rearrangements (Rawstron, Leukemia, 2016). Here, we report a sensitive NGS-based B-cell MRD Assay to detect MRD in B-cell malignancies such as acute lymphoblastic leukemia (ALL), chronic lymphocytic leukemia (CLL), multiple myeloma (MM) and mantle cell lymphoma (MCL) with *IGH* FR1 and FR3.





2.0E-01 to 3.0E-07. > MRD detection of cell and DNA contrived samples was observed to be equivalent with R<sup>2</sup> of 0.87 and slope of 1.09 in regards to calculated (calc.) MRD Frequency (i.e Clonal MRD Frequency)

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### **Results: LOB, LOD, and LOQ**

Transformed Fit to Log						
Slope	Intercept					
1.09	- 0.32					

> LOB Results: No clonal sequence was detected in 10 healthy bone marrow samples tested at 0.5 µg and 20 µg (N= 180 per master mix lot) with both *IGH* FR1 and FR3 gene targets. Thus, LOB was zero.

#### IGH FR1- LOD/LOQ Results

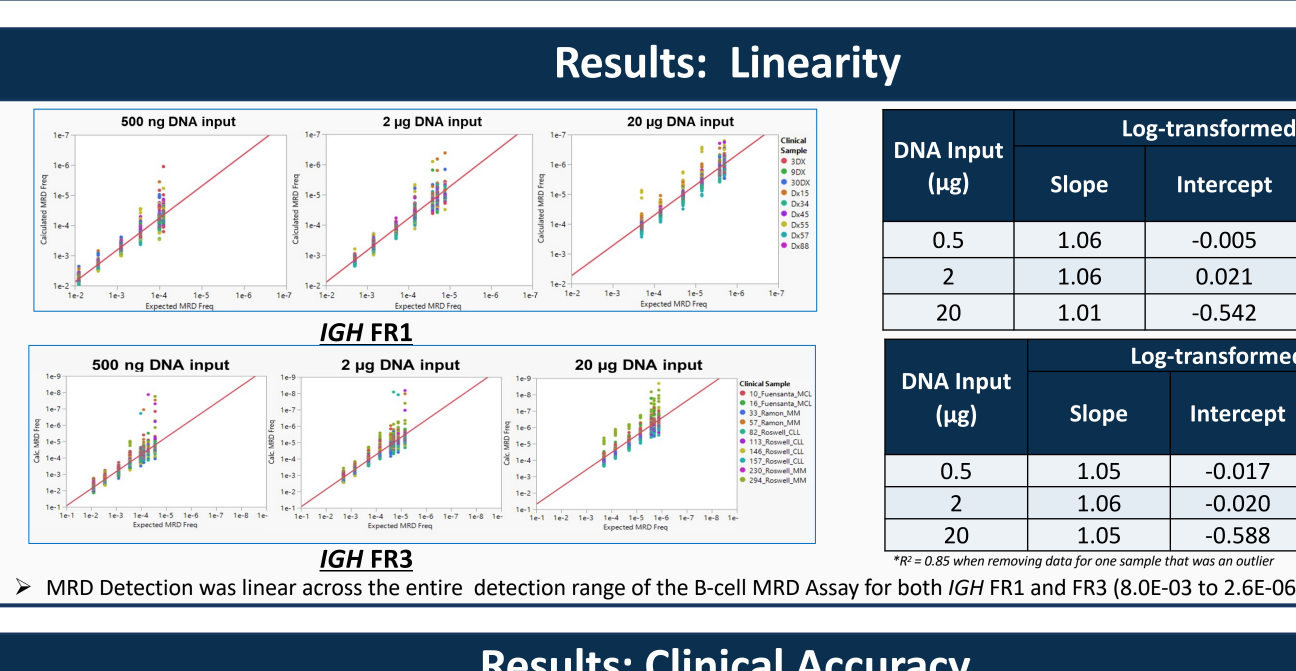
DNA Input	Clonal cell estimates	<b>Clonal MRD Frequency</b>	Clonal cell estimates	Calculated (Clonal) MRD Frequency
(µg)	LOD [95% CI]	LOD [95% CI]	LOQ [95% CI]	LOQ [95% CI]
0.5	7.0 [4.9 - 12.1]	9.1E-05 [6.4E-05 - 1.6E-04]	5.4 [3.8 - 8.5]	7.0E-05 [4.9E-05 - 1.1E-04]
2	4.7 [3.5 - 7.7]	1.5E-05 [1.1E-05 - 2.5E-05]	4.6 [2.6 - 7.9]	1.5E-05 [8.5E-06 - 2.6E-05]
20	5.4 [4.0 - 8.5]	1.8E-06 [1.3E-06 - 2.7E-06]	7.3 [5.4 - 9.2]	2.4E-06 [1.7E-06 - 3.0E-06]

 $\succ$  LOD is highest value at 20 µg : 7.0 cells which is equivalent to 2.3E-6  $\succ$  LOQ is highest value at 20 µg: 7.3 cells which is equivalent to 2.4E-6

#### IGH FR3 - LOD/LOQ Results

DNA Input (µg)	Clonal cell estimates LOD [95% CI]	Clonal MRD Frequency LOD [95% CI]	Clonal cell estimates LOQ [95% CI]	Calculated (Clonal) MRD Frequency LOQ [95% CI]
0.5	3.9 [3.0-7.2]	5.11E-05 [3.9E-05 – 9.4E-05]	6.4 [5.1-8.0]	8.3E-05[6.6E-05-1.0E-04]
2	4.7 [3.7-7.4]	1.54E-05[1.2E-05 - 2.4E-05]	12.1 [5.2-28.2]	3.9E-05[1.7E-05-9.2E-05]
20	5.7 [4.2-10.1]	1.84E-06[1.4-06 – 3.3E-06]	51.4*[6.7-396.9]	1.7E-05[2.2E-06-1.3E-04]

 $\succ$  LOD is highest value at 20 µg : 5.7 cells which is equivalent to 1.8E-6  $\blacktriangleright$  LOQ is highest value at 20 µg: 51.4 cells which is equivalent to 1.7E-5 \*LOQ is 14.4 cells which is equivalent 4.7E-6 when removing data for one sample that was an outlier



- > 10 Healthy bone marrow and 20 contrived MRD clinical samples (CLL, MM, and MCL) were tested with B-Cell MRD Assay for the *IGH* FR1 and FR3.
- $\succ$  The clonal sequences detected during baseline testing with the reference method (clonality assay) were used to track MRD in contrived clinical samples.
- FR1 and FR3 gene targets.

N =:	220	IGH FR1				
N =.	220	Detected	Not Detected			
FR1 Clonality	Clonal	20	0			
Assay	Non-Clonal	0	200			

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DNA Input	Lo	g-transformed	Fit
μg)	Slope	Intercept	R <sup>2</sup>
0.5	1.06	-0.005	0.91
2	1.06	0.021	0.88
20	1 01	1.01 -0.542	
20	1.01	0.542	0.85
	Lc	og-transformed	
DNA Input (μg)	Lc		
DNA Input		og-transformed	Fit
DNA Input (µg)	: Slope	og-transformed Intercept	Fit R <sup>2</sup>

### **Results: Clinical Accuracy**

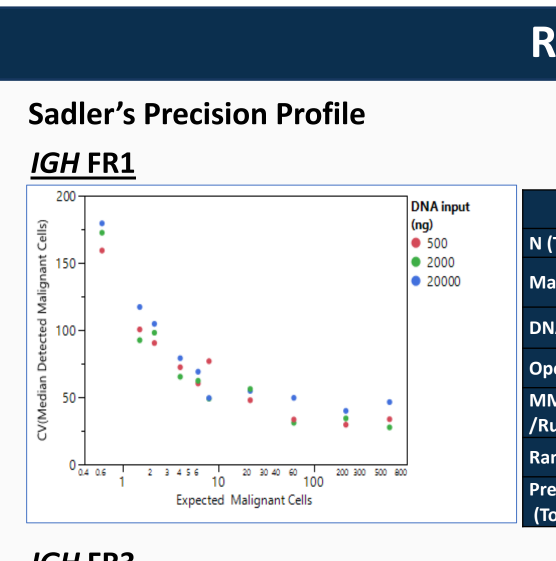
> Positive Percent agreement (PPA) and Negative Percent Agreement (NPA) was 100% for both IGH

NI - 2	20	IGH FR3			
N =2	20	Detected	Not Detected		
FR3 Clonality	Clonal	20	0		
Assay	Non-Clonal	0	200		

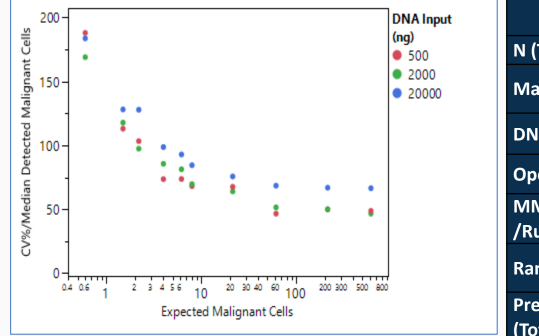
Results: Carryover and Crossover Contamination								
Study	Read Cutoff (MRD Detection)	Minimum Threshold Reads	Rate 95% CI ± 2 sd	Average Rate				
Crossover	≥ 2 reads	≥165,160	1.21E-05	1.11E-06				
Carryover	≥ 2 reads	≥ 19,730	1.01E-04	7.91E-06				

- clonal sequence in a previous or within the same run.

- positive.



#### IGH FR3



> Study was conducted with two master mix lots, 2 operators, 2 instruments, 3 DNA inputs with 9 clinical samples (CLL, MM and MCL). CV% decreases with increase in Clonal Malignant cell level

- precision with %CV ranging from 27.6% to 63.5% at the LoD.
- precision with %CV ranging from 46.6% to 89.4% at the LoD.
- study that included 20 clinical specimens (CLL, MM and MCL) and 10 healthy donors.

## H053

#### Results Carryover and Crossover Contamination

> The clonal sequence of a sample is searched in a previous run for carryover analysis and within the same run for crossover contamination. No two samples should have the same

> If target reads of a sample are greater than minimum threshold reads , a Z-score is calculated to determine the probability (p-value) for a sequence to be detected.

• If p-value < 0.05 then the sample is considered detected, it's a true positive

• If p-value  $\geq 0.05$  then the sample is considered not detected, and signal is a false

#### **Results: Precision**

		%CV per Malignant Cell Level								
(Total data points)	216	216	216	216	216	216	216	216	216	216
alignant Cell Input	615.4	215.4	61.5	21.5	8.0	6.2	4.0	2.2	1.5	0.6
NA Input	12.9	10.0	14.3	10.4	0.0	5.3	13.3	0.0	22.6	24.1
perator	3.4	0.0	0.0	0.0	5.1	0.0	0.0	0.0	0.0	3.8
M Lot/Instrument tun/Day	0.0	0.5	0.0	0.0	7.7	2.0	9.4	0.0	4.6	0.0
andom Error	35.4	34.3	37.2	52.9	59.0	63.6	71.4	97.2	102.3	173.8
recision Total Variation)	37.8	35.8	39.8	53.9	59.2	63.8	72.7	97.2	104.7	175.5

		%CV per Malignant Cell Level								
(Total data points)	216	216	216	216	216	216	216	216	216	216
alignant Cell Input	615.4	215.4	61.5	21.5	8.0	6.2	4.0	2.2	1.5	0.6
NA Input	25.8	30.5	30.4	19.5	30.8	26.5	26.4	24.6	25.9	30.2
perator	13.3	14.7	0.0	7.0	13.1	16.3	0.0	0.0	6.5	0.0
M Lot/Instrument tun/Day	10.7	11.1	9.4	8.6	14.6	0.0	0.0	0.0	0.0	35.1
andom Error	41.9	44.2	46.2	60.3	64.0	72.5	78.3	100.6	110.4	178.4
ecision otal Variation)	52.1	56.8	56.1	64.3	73.6	78.9	82.6	103.6	113.6	184.4

#### Conclusions

> IGH FR1 exhibited a LoB at 0, LoD at 2.3E-06, LoQ at 2.4E-06, good linearity with R<sup>2</sup>>0.88 across 4.5 logs and

IGH FR3 exhibited a LoB at 0, LoD at 1.8E-06, LoQ at 1.7E-05, good linearity with R<sup>2</sup>>0.74 across 4.5 logs and

> The B-cell MRD Assay demonstrated excellent clinical performance; yielding 100% agreement in an accuracy