

Evaluation of Omega Mag-Bind® TotalPure NGS Beads for DNA Size Selection

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BACKGROUND & OBJECTIVE

Carboxyl-coated paramagnetic beads such as AMPure, SPRISelect, and Omega Mag-Bind are frequently used in molecular biology to purify DNA from within reaction mixtures. Beads are typically supplied in a solution containing polyethylene glycol (PEG) and sodium chloride (NaCl). These components cause the negatively-charged DNA to condense and bind to the positively-charged beads. Once the supernatant is removed from the magnetized bead pellet, purified DNA is obtained by washing the beads in ethanol and eluting the DNA into elution buffer.

The nature of the reversible immobilization of DNA onto the beads is dependent upon the amount of PEG and NaCl in solution. Bead suppliers typically recommend a bead:sample ratio (v/v) of **1.8 to I** (referred to as **1.8x**) in order to recover DNA fragments above 100bp in length, at the exclusion of unincorporated dNTPs and primers. Using ratios lower than 1.8x means there is less PEG/NaCl in solution, which prevents small DNA fragments in the mixture from condensing and binding to beads in the solution. Thus, by using ratios other than the recommend 1.8x, DNA fragments of differing lengths will be preferentially bound, which permits the targeted the recovery of desired lengths of DNA. This behavior can be utilized to target a desired size cutoff, i.e. an approximate size division between the "small" and "large" fragments.

The objective of this report is to determine the specific DNA fragment sizes that can be recovered at various applied ratios of Omega Mag-Bind® TotalPure NGS magnetic beads.

MATERIALS

DNA Sources: O'GeneRuler **Low Range** DNA Ladder, 0.1 μg/μL (Thermo Fisher #SM1203, Lot 00593385)

Contains fragment sizes (bp): 25, 50, 75, 100, 150, 200, 300, 400, 500, 700

O'GeneRuler 100bp Plus DNA Ladder, 0.1 µg/µL (Thermo Fisher #SM1153, Lot 00136737) Contains fragment sizes (bp): 100, 200, 300, 400, 500, 600, 700, 800, 900, 1000, 1200, 1500, 2000, 3000

O'RangeRuler **500bp** DNA Ladder, 0.05 µg/µL (Thermo Fisher #SM0643, Lot 00607550)

Contains fragment sizes (bp): 500, 1000, 1500, 2000, 2500, 3000, 3500, 4000, 4500, 5000, 5500, 6000

Magnetic Beads: Mag-Bind TotalPure NGS Beads (Omega Bio-Tek #M1378-02, Lot TPN031417JCM22873)

Quantification: Quant-iT dsDNA High Sensitivity Assay Kit (Thermo Fisher #Q33120, Lot 1966973)

SpectraMax M5E Microplate Reader (Molecular Devices #M5E)

Fragment Analysis: High Sensitivity NGS Fragment Analysis Kit (Advanced Analytical #DNF-474-500)

Fragment Analyzer Automated CE System (Advanced Analytical)

METHOD

- I. The 3 DNA ladders listed above were combined as follows:
 - + I part of O'GeneRuler Low Range @ 0.1 μg/μL
 - + I part of O'GeneRuler 100bp Plus @ 0.1 μg/μL
 - + 2 parts of O'RangeRuler 500bp @ 0.05 μg/μL
 - = 4 parts of mixed ladder @ 66.67 ng/µL

- 2. The mixed ladder was diluted to approx. 15 ng/ μ L in Qiagen elution buffer (EB) (10 mM Tris-Cl, pH 8.5) and distributed into aliquots of 50 μ L each
- 3. Aliquots of mixed ladder were size-selected using Omega Mag-Bind® TotalPure NGS beads with a standard bead cleanup protocol consisting of these main steps (see Table I & Figure I for more details):
 - a. Mixing of 50 µL mixed ladder with Omega beads at 23 different beads:sample ratios (Table I)
 - b. Magnetization to separate the supernatant from the beads
 - c. Removal of the supernatant
 - d. Washing the beads twice with 150 μ L of 80% ethanol
 - e. Air drying the beads
 - f. Eluting the DNA in 50 µL of Qiagen EB
- **4.** From each of the 23 size selection reactions, both the **supernatant fraction** (containing smaller unbound fragments) and the **bead fraction** (containing larger bound fragments) were retained, and their bead cleanups were performed in parallel (Figure 1).

Keeping both fractions in each reaction allows the separate recovery of both the small fragments and the large fragments on either side of the targeted size cutoff. Thus, each of the 23 ratio tests produced two separate eluted DNA samples.

We establish the naming convention for these size-selection cleanups according to which fragments are removed from the final eluted product:

- Small Fragment Removal (SFR) Cleanup =
 - Removes smaller fragments (in the supernatant)
 - Keeps larger fragments (bound to beads)
- Large Fragment Removal (LFR) Cleanup =
 - Removes larger fragments (bound to beads)
 - Keeps smaller fragments (in the supernatant)
 NOTE: In order for the smaller fragments to be purified from the supernatant, a second addition of beads is required to bind the small DNA in solution (Figure 1).
- 5. All final eluted samples, along with the non-size selected mixed ladder, were characterized as follows:
 - Quantified to measure total DNA yield with Quant-iT dsDNA High Sensitivity Assay Kit
 - Fragment analyzed to determine recovery of the various DNA lengths with High Sensitivity NGS Fragment Analysis Kit

These data were then normalized to the results from the 1.8x bead test (referred to as the "control" ratio as this is the ratio recommended by bead suppliers) to determine percent yields.

6. Each of the 23 tests were repeated independently, for a total of 2 replicates of each test condition. All data reported here are the means of the two replicates for each data point.

Table 1. Conditions used for size selection tests (*n*=2 for each condition). The control ratio of 1.8x (supplier recommended) is indicated in **red**.

Bead: Sample Ratio (v/v)		I st DNA	A binding ste	Þ		2 nd DNA binding step							
	Volume of sample used (µL)	Volume of beads added (µL)	Volume of supernatant removed (µL)	Fate of supernatant	_	Volume of beads added to the previous supernatant (µL) in order to achieve the control bead:sample ratio of 1.8x	Volume of total PEG/NaCl in solution (µL)	Volume of final supernatant removed (μL)	Fate of supernatant				
0.30×	50	15	65		•	75	90	140					
0.35×	50	17.5	67.5			72.5	90	140					
0.40x	50	20	70	Kept and	treated as a arallel sample or recovering both the large & small fractions separately, a a 2nd DNA	70	90	140 140					
0.45x	50	22.5	72.5			67.5	90						
0.50×	50	25	75	for recovering		65	90	140					
0.55×	50	27.5	77.5			62.5	90	140					
0.60x	50	30	80			60	90	140 140 140	Discouded				
0.65×	50	32.5	82.5	fractions		57.5	90						
0.70×	50	35	85			55	90						
0.75×	50	37.5	87.5	binding step		52.5	90	140	Discarded				
0.80x	50	40	90	[as per		50	90	140					
0.85×	50	42.5	92.5	columns to the		47.5	90	140					
0.90×	50	45	95	right 🔿		45	90	140					
0.95×	50	47.5	97.5	and indicated in Figure 1]		42.5	90	140					
1.00x	50	50	100			40	90	140					
1.20x	50	60	110			30	90	140					
1.40×	50	70	120			20	90	140					
1.60x	50	80	130			10	90	140					
1.80x (control)	50	90	140										
2.00×	50	100	150										
2.25x	50	112.5	162.5	Discarded									
2.50x	50	125	175										
3.00x	50	150	200										

Figure I. Diagram of method to perform SFR and LFR using Omega beads. The method used in this report to keep both fractions is indicated in **red**. An optional method to perform double-ended selection is in **purple**.

Combine beads with sample at a ratio to achieve the desired cutoff; Vortex to mix; Quick spin down to collect droplets Incubate solution @ room temp for 5 mins Ist DNA binding step Place on magnet until supernatant is clear; Remove supernatant (contains smaller fragments + PEG + NaCl) **SMALL FRAGMENT REMOVAL (SFR): LARGE FRAGMENT REMOVAL (LFR):** Remove smaller fragments (in the supernatant) Remove larger fragments (bound to beads) **Keep larger** fragments (bound to beads) **Keep smaller** fragments (in the supernatant) **Discard** clear supernatant; Keep bead pellet on magnet **Transfer** clear supernatant to new tube Add additional Omega beads (in order to Method to recover <u>BOTH</u> fractions separately: re-bind small fragments in the supernatant) to bring up the total volume of PEG/NaCl Keep the supernatant and 2nd DNA input here as a parallel in solution to achieve a full 1.8x bead ratio binding sample for LFR step Incubate @ room temp for 15 minutes; Add freshly-prepared 80% ethanol to bead pellet Place on magnet until supernatant is clear (minimum volume = original volume of beads + sample); Incubate @ room temp for 30 seconds; Ethanol Remove & discard clear supernatant Remove & discard Ist ethanol wash washes (contains only PEG + NaCl at this point); Keep bead pellet on magnet Repeat wash as above for a total of 2 washes Add freshly-prepared 80% ethanol to bead pellet (minimum volume = original volume of beads + sample); Quick spin down; Place on magnet; Incubate @ room temp for 30 seconds; Remove any residual ethanol from tube washes Remove & discard Ist ethanol wash Open caps; Allow beads to air dry on Dry magnet @ room temp for 2 minutes Repeat wash as above for a total of 2 washes beads Quick spin down; Place on magnet; Remove tubes from magnet; Remove any residual ethanol from tube Add desired volume of elution buffer; Cap tubes; Vortex to fully resuspend beads; Quick spin down to collect droplets Open caps; Allow beads to air dry on Dry Elute beads magnet @ room temp for 2 minutes DNA Incubate @ room temp for 3-5 minutes; Place on magnet until supernatant is clear Remove tubes from magnet; Add desired volume of elution buffer; Cap tubes; Vortex to fully resuspend beads; **Transfer** clear supernatant Quick spin down to collect droplets **Elute** (eluted larger fragments) to new tube DNA Incubate @ room temp for 3-5 minutes; Optional method for **DOUBLE-ENDED** size selection: Place on magnet until supernatant is clear Instead of adding beads to achieve the full 1.8x bead ratio (which will recover ALL fragments > 100bp), add just enough beads to achieve a final ratio for your desired lower end **Transfer** clear supernatant cutoff; e.g., to achieve a 0.7x or 0.8x rather than the full 1.8x (eluted smaller fragments) to new tube

RESULTS & DISCUSSION

Data from these tests indicate that different ratios of Omega Mag-Bind® TotalPure NGS beads can be used to precisely target any desired size cutoff (i.e. approximate size division between the "small" and "large" fragments) for the separation of DNA fragments \leq 3000 bp. The recovery of the small or large fragments on either side of the cutoff is accomplished simply by keeping either the bead-bound fraction or the supernatant fraction (Figures 2 & 3; Table 2).

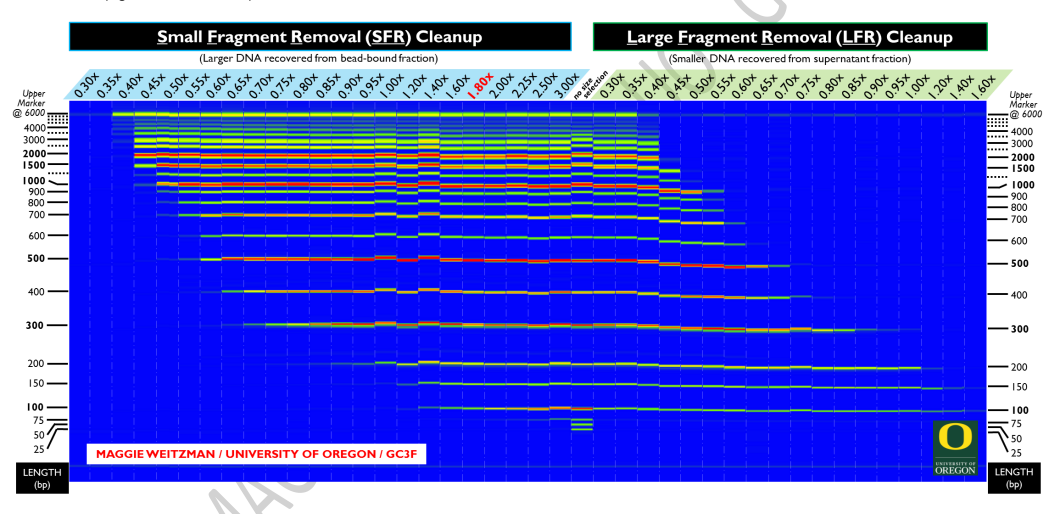


Figure 2. Pseudogel image from the fragment analysis of DNA recovered after size selection. Note: Data from only 1 of the 2 test replicates is shown here.

Table 2. Heatmap of percent of DNA removed for various fragment lengths (versus 1.8x control, in **red**). n=2 for each condition *Note: Negative percentages are reported as 0%.*

9	% OF D	NA -			FRAGMENT LENGTH (bp)																				
	% OF D	VA R	EMO	/ED	100	150	200	300	400	500	600	700	800	900	1000	1200	1500	2000	2500	3000	3500	4000	4500	5000	5500
	0.30		n/a		99	100	100	100	100	100	100	100	100	99	100	99	100	100	100	100	99	100	99	99	99
	0.35	ζ .	n/a	UF	100	100	100	100	99	100	100	100	100	100	100	99	100	100	99	100	99	99	99	99	99
	0.40	ζ	3,000	AN	99	100	100	100	100	100	100	100	100	100	100	99	99	98	95	86	71	52	39	30	48
	0.45	ζ .	1,000	CLEANUP	100	100	100	100	100	99	99	98	98	96	94	80	48	1	0	0	0	0	0	0	0
	0.50	‹	700) C	99	100	100	100	99	98	96	92	82	66	50	9	0	0	0	0	0	0	0	1	3
	0.55	ر ا	500	REMOVAL (SFR)	99	100	99	99	96	95	85	68	38	20		9	7	7	8	15			9	12	9
	0.60	fragments	450	<u>5</u>) 7	99	99	99	98	91	78	44	19	8	7	5	7	7	5	6	6	9	7	8	2	0
	0.65	ַ אַ וּ	350	٧A	99	99	99	95	73	26	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0.70	₹] ge	300	0	99	99	98	85	35	5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	0.75	<u>.</u> [280	EN	97	99	96	76	23	15	6	5	5	6	4	5	7	5	2	5	6	5	2	4	11
	0.80	"large"	260		98	99	96	59	17	22	13	13	13	13	12	12	14	12	11	12	15	16	14	17	23
	0.85	<u>a</u> [240	<u>F</u> RAGMENT	98	97	90	27	0		0	0	0	0	0	0	0	0	0	0	1	1	0	0	0
	0.90	₫ 🖥	220	ME	97	96	84	15	2	9	4	3	3	4	2	3	6	3	4	5	7	9	5	7	4
	3 0.95	and	200	٩G	97	95	77	5	0		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
) 1.00	ી ≒ૄ	175	<u>F</u> R/	96	93	65	11	9	15	10	9	9	9	8	3	10	8	4	5	7	8	9	15	3
	1.00 1.20 1.40 1.60	"small"	150		90	67	12	0	0	19	0	0	0	0	0	0	0	0	0	0	0	8	2	6	0
	5 1.40		125	<u>S</u> MALL	72	27	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0			8
		e l	100	SN	46	15	8	8	9	12	10	10	10	10	10	12	11	12	12	14	14	15	13	15	15
	当 <u>1.80</u>	etween	100		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
	1.80 0.30 0.35 0.40 0.45	et 🗠	n/a	JP	0	3	4	5	6	4	9	8	8	8	7		8	9			11	16	9	19	14
	S 0.35		n/a	N	14	10	9	12	13	13	14	15		15	14	17	16	16	18	18	20	22	14	20	25
	0.40	(dq)	3,000	CLEANUP	21	19	18	20	20	22	20	21	19	20	19	21	20	21	24	31	44	57	67	75	77
	0.45	∐	1,000	C	21			15	15	21	18	18	19	20	22	34	59	95	99	99	99	99	99	99	99
	0.50	⊣	700	(LFR)	28	19	18	18	21	30	26	29	36	48	61	89	98	100	100	100	99	99	99	99	99
	0.55	45	500	<u>I</u>	26				13	20	24	35	67	85	93	99	99	99	99	99	99	99	99	99	98
	0.60		450	AL	24	13	11	12	18	36	51	88	97	99	99	99	99	99	99	99	99	99	99	99	99
	0.65	- 1	350	9	36	22	21	25	40	70	93	99	100	100	100	100	100	100	99	99	99	99	99	99	99
	0.70	•	300	Ž	27		13	23	65	94	98	99	99	99	99	99	99	99	99	99	98	98	98	97	98
	0.75	⊈ ©	280	RE	29	17	0	38	85	98		99	99	99	99	99	99	99	99	99	99	99	99	99	99
	0.80	- L	260	NT	33	13	0	54	95	100	100	100	100	100	100	100	100	100	100	100	99	100	100	100	100
	0.85		240	ΛEI	29	8	12	72	97	99	98	99	100	100	99	100	99	99	99	99	99	99	100	100	100
	0.90		220	LARGE FRAGMENT REMOVAL	38	19	30	88	99	99	99	99	99	99	99	99	99	99	99	99	99	99	99	99	99
	0.95	_	200	RA	45	22	36	93	99	100	99	99	99	99	99	99	99	99	99	99	99	99	99	99	99
	1.00	_	175	ΕF	40	16	44	97	99	99	99	99	99	98	99	99	99	99	99	99	99	99	99	99	99
	1.20	_	150	3G	51	40	88	99	99	99	99	99	99	97	99	99	99	99	99	99	99	99	99	99	99
	1.40	_	125	₽.	60	72	97	99	99	99	99	99	99	99	100	99	99	99 MAGG	SIE WEIT	ZMAN/	UNIVE	RSITY O	F OREG	100 ON / GC	100 3F
L	1.60	(100	1	81	91	99	100	100	100	99	99	100	100	100	100	100	100	100	100	100	100	100	100	100



0 to 2

RESULTS, cont'd

Table 3. Total DNA recovery for SFR, LFR, and summed fractions (versus 1.8x control, in red). n=2 for each condition

Summary of key findings:

When combining DNA yields from the SFR & LFR fractions (for the 18 ratios ≤1.8x), the following amounts of DNA were recovered (Table 3):

- 95-100% total recovery for 13 of the 18 tests
- 91-94% total recovery for 5 of the 18 tests

These yields indicate that Omega beads have a high DNA recovery rate and can be considered reliable for performing bead cleanups and size selection.

Less than 1% of the total DNA was retained when using a ratio under 0.40x (i.e. 0.35x or 0.30x). This is likely because there is simply not enough PEG/NaCl in solution to condense the DNA. Due to extremely low yields, these lowest bead ratios are not recommended for size selection.

The effect of size selection on the 6000 bp ladder band could not be evaluated because the Upper Marker reagent in the fragment analyzer kit (size 6000 bp) occurs at the same location on the electropherogram.

	Recovered from (larger fragme			Recovered fro (smaller fragm	·	
Bead: Sample Ratio (v/v)	Mean DNA concentration (ng/µL) ± SD (n=2)	% Total DNA yield	•	Mean DNA concentration (ng/µL) ± SD (n=2)	% Total DNA yield	Sum of yields from SFR + LFR
0.30×	0 ± 0	0%		12.4 ± 0.3	98%	99%
0.35×	0 ± 0	0%		12.2 ± 0.1	97%	97%
0.40×	1.0 ± 0.2	7%		11.6 ± 0.3	92%	99%
0.45×	4.0 ± 0.2	31%		8.1 ± 0.2	64%	95%
0.50×	5.7 ± 0.3	44%		6.6 ± 0.1	52%	96%
0.55×	6.8 ± 0.3	52%		5.2 ± 0.1	41%	93%
0.60×	7.8 ± 0.4	60%		4.3 ± 0.1	34%	93%
0.65×	8.8 ± 0.2	67%		3.3 ± 0.1	26%	93%
0.70×	10.3 ± 0.2	79%		2.5 ± 0	20%	99%
0.75×	10.2 ± 0.2	78%		2.1 ± 0.1	16%	95%
0.80×	10.7 ± 0.4	82%		1.7 ± 0.1	14%	96%
0.85×	11.4 ± 0.2	88%		1.5 ± 0	12%	100%
0.90×	11.2 ± 1.1	86%		1.2 ± 0.1	9%	95%
0.95×	11.5 ± 0.3	88%		1.0 ± 0.2	8%	96%
1.00×	11.8 ± 0.3	91%		1.0 ± 0.1	8%	98%
1.20×	12.5 ± 0.1	96%		0.5 ± 0.1	4%	100%
1.40×	11.9 ± 0.5	91%		0.3 ± 0.0	2%	93%
1.60×	II.7 ± 0.7	90%		0.1 ± 0.1	1%	91%
I.80x (control)	13.0 ± 0.6	100%				
2.00×	13.1 ± 0.7	101%				
2.25x	13.5 ± 0.6	103%				
2.50x	13.4 ± 0.5	103%				
3.00x	13.8 ± 0.3	106%				

For most NGS library cleanups where the objective is to remove fragments <200bp (leftover primer, adapter dimers, small insert libraries, etc.), a bead ratio of **0.8x** would be adequate in most cases.

Ratios at 1.8x and below are intended to recover fragments greater than 100 bp in length (as per bead supplier protocols). To determine whether DNA fragments less than 100 bp can also be retained with Omega beads, several ratios above the recommended 1.8x were tested using the SFR method (Figure 1). It was found that more of the 25, 50, and 75 bp fragments could indeed be recovered when increasing the ratio above 1.8x. In fact, a 3.00x ratio recovered more than twice as many 25 bp fragments and nearly 6 times as many 75 bp fragments as a 1.8x ratio (Table 4).

RESULTS, cont'd

Table 4. Yield of small DNA fragments for ratios above the 1.8x control (in red). n=2 for each condition

Bead: Sample Ratio	•		fragment	_	% yield of small fragment lengths (versus non-size selected ladder)							
(v/v)	25 bp	50 bp	75 bp	100 bp	25 bp	50 bp	75 bp	100 bp				
I.80x (control)	100%	100%	100%	100%	2%	0%	7%	34%				
2.00x	118%	123%	149%	143%	2%	0%	10%	48%				
2.25×	98%	70%	236%	207%	2%	0%	16%	70%				
2.50×	146%	102%	344%	238%	3%	0%	24%	80%				
3.00x	209%	163%	587%	273%	4%	1%	41%	92%				

However, even a ratio of 3.00x could only recover 4% of the 25 bp fragments that were present in the original non-size selected DNA ladder (Table 4). Thus, while increasing the bead ratio beyond the recommended 1.8x can provide a higher recovery of smaller fragments, it is still a relatively small recovery when compared to the total amount of small DNA present in the original sample. For applications where recovery of these small fragments is more critical, increasing the ratio even further beyond 3.00x could result in higher yields.

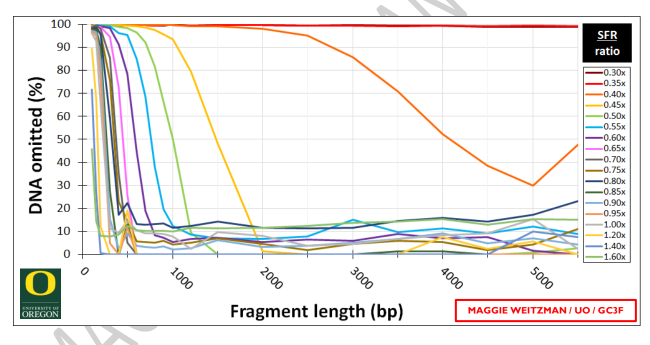


Figure 3.
Percent of DNA
removed
at various
fragment
lengths for
SFR tests
(versus 1.8x
control, in
red). n=2
for each
condition

Note: Negative percentages are reported as 0%

Extrapolating from these data, ratios at smaller increments than the 0.5x increments shown here could be used to target cutoffs at intermediate intervals. For example, a ratio of 0.57x would result in an expected cutoff around 600bp. At the GC3F, we routinely use such smaller increment ratios for size selecting DNA.

We also routinely combine the SFR and LFR methods into a double-ended size selection protocol (Figure 1), such that the two size cutoffs can be made in a single tube. Removing DNA from both ends of the spectrum allows the targeted recovery of fragments from only within a specific range (e.g., from 400-800bp, from 275-625bp, etc.). We find Omega beads to be immensely useful and an essential part of our NGS workflows!