







Protocol	#4.3
Title	BOMB Gel extraction
Keywords	Gel-extraction, silica-beads, carboxyl beads
Authors	Oberacker P*, Stepper P*, Bond DM*, Höhn S, Focken J, Meyer V, Schelle L, Sugrue VJ, Jeunen GJ, Moser T, Hore SR, von Meyenn F, Hipp K, Hore TA# and Jurkowski TP#
Citation	<i>Oberacker et al., Bio-On-Magnetic-Beads (BOMB): Open platform for high-throughput nucleic acid manipulation. Submitted</i>
Online Revision	https://bomb.bio/protocols/ V1.0 (13 th August 2018)

Summary

The BOMB gel extraction protocol is designed to purify DNA of specific sizes from agarose gels [1]. The protocol assumes users know how to run an agarose electrophoresis gel and safely cut out the band they want. Following this, the gel slice is melted at 50 °C in 1.5 volumes of GITC buffer [2] (also used for BOMB protocols #6.1 and #6.3). DNA is then captured on either silica- or carboxyl-coated beads using isopropanol to trigger precipitation. Standard isopropanol and ethanol washes are used to further purify the DNA ahead of resuspension.

Chemicals

Name	Provider	PN	MW [g/mol]		Safety codes
Ethanol (C₂H₆O, 99.9%)	Honeywell/ Riedel-de Haën	34963	46.07	 Danger	H: 225-319 P: 210-240-305+351+338-403+233
Guanidine Isothiocyanate (GITC, C₂H₆N₄S)	Roth Chemicals	2628.4	118.16	 Warning	H: 302+312+332-412-EUH032 P: 273-280-302+352-304+340-312
Tris(hydroxymethyl)-aminomethane (Tris, C₄H₁₁NO₃)	Roth Chemicals	AE15.3	121.14	 Warning	H 315-319-335 P: 280-302+352-305+351+338-312
N-Lauroylsarcosine sodium salt (Sarkozyl, C₁₅H₂₉NO₃Na)	Sigma (Merck)	L9150-50G	293.38	 Danger	H: 315-318-330 P: 260-280-284-305+351+338-310
Antifoam 204	Sigma (Merck)	A8311-50ML	n.a.	n.a.	n.a.
Ethylenediamine tetraacetic acid disodium salt dihydrate (EDTA, C₁₀H₁₄N₂Na₂O₈ · 2 H₂O)	Roth Chemicals	8043.1	372.24	 Danger	H: 332-373 P: 260-314
Isopropanol (C₃H₈O)	Acros Organic	184130025	60.01	 Danger	H: 225-319-336 P: 210-233-240-305+351+338-403+235

Please consult appropriate MSDS information before working with these chemicals! Use lab coat, gloves and eye protection at all times! The chemicals are available from other providers as well. No preference is given to the indicated vendors.

Buffers and solutions

Silica-coated BOMB beads in TE (1:10 diluted from stock)

Or, 1:50 TE diluted carboxylate-coated magnetic beads can be used; for example, purchased from GE Healthcare (Sera-Mag SpeedBead Carboxylate-Modified Magnetic Particles, Hydrophilic, 15 ml, cat., 45152105050250)

TE buffer (stable for at least a year at RT)

10 mM Tris pH 8.0

1 mM EDTA

GITC buffer (stable for at least several weeks at RT)

Reagent	Concentrations	For 50 ml
GITC	4 M	23.64 g
Tris HCl pH 7.6-8.0	50 mM	2.5 ml of 1 M stock
Sarkosyl	2%	1 g
EDTA	20 mM	2 ml of 0.5 M stock
Antifoam (optional)	0.1 %	50 µl

adjust pH with HCl to 7.6-8.0 and adjust the volume with water to 50 ml

Isopropanol (~0.5 ml – 1 ml per sample)

80% ethanol (1 ml per sample)

Elution buffer (stable, can be stored for at least 1 year at RT; 30 µl per sample)

5 mM Tris-HCl, pH 8.5

Equipment and setup

Heat block (e.g. Roth, Rotilabo®- block thermostat H250 – PN: Y264.1)

Magnetic stand for 1.5 ml microcentrifuge tubes (numerous vendors available)





Pipettes

20 – 200 µl (e.g. Gilson Pipetman P200)

200 – 1000 µl (e.g. Gilson Pipetman P1000)

1.5 ml microcentrifuge tubes (numerous vendors available)

BOMB Gel extraction

Step	Melt gel slice	Time	<input checked="" type="checkbox"/>
1	Collect a gel slice containing the desired DNA fragment or amplicon into a 1.5 ml microcentrifuge tube		<input type="checkbox"/>
2	Weigh tube containing gel slice (and subtract mass of empty tube)		<input type="checkbox"/>
3	Add 1.5 volumes of GITC buffer to 1 volume of gel slice (e.g. 150 µl of GITC buffer for 100 mg gel slice)	1 min	<input type="checkbox"/>
	<i>The GITC buffer contains guanidine isothiocyanate, a chemical which can cause burns and sensitivity and produces hydrogen cyanide when mixed with acid</i>		
4	Incubate gel slice in GITC buffer at 50 °C for 10 min or until fully dissolved (mixing by inversion during incubation helps accelerate process)	10 min	<input type="checkbox"/>
Step	Task - DNA capture and purification		
5	Calculate total volume of gel/GITC mix (e.g. 150 µl of GITC buffer + 100 mg of gel slice = 250 µl total)		<input type="checkbox"/>
6	Add 50 µl of TE-diluted beads to a <u>new</u> 1.5 ml microcentrifuge tube and top up to 1.67 volumes of original gel slice size with TE buffer (e.g. for 100 mg gel slice, combine 50 µl TE-diluted beads + 117 µl of 1x TE buffer)	1 min	<input type="checkbox"/>
7	Add 3.34 volumes of isopropanol to the TE-diluted beads from step 6 (e.g. for 100 mg gel slice, add 334 µl of isopropanol)	1 min	<input type="checkbox"/>
8	Add melted gel-GITC mix from step 4 to the beads-isopropanol tube from step 7 and mix by pipetting. Incubate for 5 min while beads capture DNA	5 min	<input type="checkbox"/>
9	Settle the magnetic beads on a magnetic stand and discard the supernatant	2 min	<input type="checkbox"/>
	<i>Ensure that the beads are completely pelleted</i>		
10	Add 500 µl isopropanol to wash the beads, wait for 30 s and discard the supernatant	2 min	<input type="checkbox"/>
11	Add 500 µl 80% ethanol to wash the beads, wait for 30 s and discard the supernatant	2 min	<input type="checkbox"/>
12	Repeat step 11 once for a total of 2 washes	2 min	<input type="checkbox"/>
13	Remove the supernatant completely and dry at RT for approximately 5 min	5 min	<input type="checkbox"/>
	<i>Make sure to remove all remaining ethanol</i>		
14	Add 20-30 µl of elution buffer or ddH₂O to elute the DNA from the beads, mix well for 5 min (shake at 1300 rpm) and spin down briefly	5 min	<input type="checkbox"/>
15	Pellet the beads magnetically and remove the eluted DNA into a fresh tube	2 min	<input type="checkbox"/>
End	Measure concentration and proceed	~35 min (20 min hands-on)	
	Store @ -20 °C or 4 °C		

Troubleshooting

Problem	Solution
Beads sticking to the tube	<ul style="list-style-type: none"> Sonicate shortly in a sonic bath and/or centrifuge briefly
Beads stay at the bottom of the tube when mixing	<ul style="list-style-type: none"> Close lid and invert tube a few times
Low DNA concentration	<p>Low DNA concentration can have multiple reasons</p> <ul style="list-style-type: none"> Gel slice incompletely solubilised: After addition of GITC buffer to gel slice, mix by inverting tube 5 times every 2-3 min during 50 °C incubation (DNA will remain in any undissolved agarose) Remaining ethanol: before elution you have to ensure that the beads are completely dry. Their colour will turn from shiny black to brittle brown. If this was not the case, dry the beads again and elute a second time

Exemplary Results

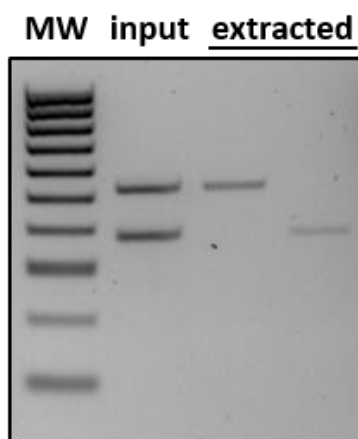


Fig 1: Gel extraction using BOMB. The second lane contains the product of a *CDH1* PCR using female chicken gDNA as a template, according to Fridolfsson and Ellegren [3]. The two rightmost lanes contain the gel extracted 519 bp and 373 bp bands from lane two using the BOMB protocol with carboxyl-coated magnetic beads. The volumes loaded are proportional (i.e. the right hand 2 lanes represent the efficiency of capture from the left hand lane). MW: Hyperladder IV (Bioline).

References

- Vogelstein B, Gillespie D. Preparative and analytical purification of DNA from agarose. *Proc Natl Acad Sci.* 1979;76: 615–619. doi:10.1073/pnas.76.2.615
- Chirgwin JM, Przybyla AE, MacDonald RJ, Rutter WJ. Isolation of biologically active ribonucleic acid from sources enriched in ribonuclease. *Biochemistry.* 1979;18: 5294–5299. doi:10.1021/bi00591a005
- Fridolfsson A-K, Ellegren H. A Simple and Universal Method for Molecular Sexing of Non-Ratite Birds. *J Avian Biol.* 1999; doi:10.2307/3677252