







Protocol	#6.1
Title	BOMB TNA extraction from mammalian cells using GITC lysis
Keywords	HT TNA isolation, carboxyl-beads, silica beads, GITC, mammalian cells
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.(2019), Bio-On-Magnetic-Beads (BOMB): Open platform for high-throughput nucleic acid manipulation. PLOS Biology,17(1), https://doi.org/10.1371/journal.pbio.3000107</i>
Online	https://bomb.bio/protocols/
Revision	V1.0 (10 th August 2018)

Summary

Isolation of total nucleic acid (TNA) from mammalian cells is a basic wet lab technique and the starting point for many analysis pathways. This protocol describes a high-throughput magnetic bead-based protocol to purify total nucleic acid (TNA) from (low protein) mammalian cells lysed in GITC buffer [1]. If RNase A is added at the beginning, only DNA will be isolated (see BOMB protocol #7.1). This protocol can also be coupled with an on-bead DNase I digest to extract only RNA (see BOMB protocol #8.2). It utilizes a sarkosyl and guanidinium-isothiocyanate (GITC) based lysis buffer and isopropanol to drive precipitation of the nucleic acid to the paramagnetic beads. Variations of this protocol exist for isolation from tissues, plants and yeast. Volumes can be adjusted, however, should remain at a consistent ratio of 2:3:4, beads:lysate:isopropanol. Below is a sensible volume for deep-well plates, however, lower volumes on 0.2 ml PCR plates are also possible (i.e. 40 µl of beads, 60 µl of lysate and 80 µl of isopropanol), and may further reduce costs.

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 (Sarkosyl, 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	18413002 5	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

Carboxyl-coated or silica-coated magnetic beads in TE buffer (160 µl per sample, 16 ml for a 96-well plate)

TE buffer (160 µl per sample, 16 ml per 96-well plate. Used for bead dilution, see above)

10 mM Tris pH 8.0

1 mM EDTA

Lysis buffer (stable for at least several weeks at RT. 240 µl per sample, 24 ml per 96-well plate)

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 (720 µl per sample, 72 ml for a 96-well plate)

80% ethanol (600 µl per sample, 60 ml for a 96-well plate)

Equipment and setup

Fume hood

Microtiter plate orbital shaker (e.g. IKA MS 3 basic)

Magnetic stand for 96-well plate (e.g. BOMB microplate magnetic rack)

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

Multichannel Pipettes

30 – 300 µl (e.g. Eppendorf, Eppendorf Research® plus 8-channel – PN: 3125000052)

50 – 1200 µl (e.g. VWR, Multi channel pipette, 8-channel – PN: 613-5422)







96-well PCR plate (e.g. Sarstedt, 96 PCR-Plate half skirt flat – PN: 170134)

1.2 ml 96-well deep well plates (e.g. Sarstedt, MegaBlock® 96 Well – PN: 82.1971.002)

Reservoirs (e.g. Roth, Rotilabo®-liquid reservoirs – PN: E830.2)

Seals (e.g. Bio-Rad, Microseal® 'B' Adhesive Seals – PN: MSB1001)

BOMB TNA extraction

Step	Sample collection	Time	<input checked="" type="checkbox"/>
1	Collect up to 10 ⁶ cultured mammalian cells in a single well of a deep-well plate		<input type="checkbox"/>
2	Pellet the cells via centrifugation at 500g and discard supernatant		<input type="checkbox"/>
	At this point, the pellets can be frozen at -20 °C and processed later		
Step	TNA purification		
3	Add 240 µl of lysis buffer , seal and shake at RT at 1400 rpm for 5 min		
	<i>The lysis buffer contains guanidine isothiocyanate, a chemical which can cause burns and sensitivity and produces hydrogen cyanide when mixed with bleach/acids</i>	10 min	<input type="checkbox"/>
4	Add 320 µl of isopropanol , seal and shake at RT at 1400 rpm for 5 min	10 min	<input type="checkbox"/>
5	Add 160 µl of coated magnetic beads (diluted 1:50 in TE from stocks), seal and shake at RT at 1400 rpm for 5 min	10 min	<input type="checkbox"/>
6	Settle the magnetic beads on a magnetic stand and discard the supernatant	5 min	<input type="checkbox"/>
	<i>Ensure that the beads are completely settled</i>		
7	Remove the plate from the magnetic stand and add 400 µl isopropanol . Shake at RT at 1400 rpm for 2 min	10 min	<input type="checkbox"/>
8	Settle the magnetic beads on a magnetic stand and discard the supernatant	5 min	<input type="checkbox"/>
9	Wash twice with 300 µl of 80% ethanol as above	10 min	<input type="checkbox"/> <input type="checkbox"/>
10	Remove the supernatant completely and dry the beads for approximately 5-10 min		
	<i>Make sure to remove all remaining ethanol. Silica-coated beads should be completely dried (at 50 °C), whereas carboxyl-coated ones should be only dried briefly (at RT)</i>	5 min	<input type="checkbox"/>
11	Add 70 µl of nuclease free water to the wells and shake for at least 5 min to resuspend (centrifuge shortly if beads stick to the walls)		
	<i>If the liquid is too viscous to pipette off and/or the beads don't settle, double the elution volume until it works</i>	5 min	<input type="checkbox"/>
12	Settle the magnetic beads on a magnetic stand and transfer the supernatant to a collection plate/tube	5 min	<input type="checkbox"/>
End	Measure concentration	~1.5 h (45 min hands-on)	
	Store @ -80 °C		

Troubleshooting

Problem	Solution
Beads sticking to the sides	<ul style="list-style-type: none"> Sonicate shortly in the sonic bath and/or push the beads down with a pipette tip
Beads stay at the bottom of the well when mixing	<ul style="list-style-type: none"> Use a good foil to cover the plate well and invert a few times
Elution very viscous	<ul style="list-style-type: none"> We observe this problem when too many cells were used for extraction. Add more elution buffer (concentration should still be very high), vortex strongly or pipette up and down. Heat up to 65 °C

Exemplary Results

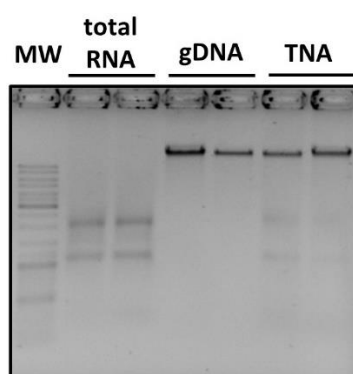


Fig 1: Quality control of BOMB TNA extraction. Total nucleic acids (TNA) was extracted from 500k HEK293 cells (two rightmost lanes). This TNA can then be digested with either DNase I to yield only RNA (lanes 3+4) or RNase A to extract the DNA (lanes 4+5). MW: GeneRuler DNA Ladder Mix, Thermo Scientific.

References

- 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