ABSTRACT
This protocol describes a clean up and size selection method for nucleic acids (tested on DNA) to deplete and remove fragments below 1 - 2 kb. The success of this depends on the cleanliness of your sample (it doesn't have to be super clean but a whole lot of contaminants make working with the beads more difficult, diluting the sample out before usage can help with that).

The concentrations of PEG and NaCl and the volume of the beads solution are crucial for recovery and proper removal of small fragments. As a basic guideline it can be said: more PEG and NaCl - higher recovery but hence less removal of small fragments and the other way round. I found for my samples (eucalyptus) that with 1 volume of the beads solution respectively to DNA sample I'm on the safe side recovery wise, but if I want to make sure to get rid of more smaller fragments I use 0.8 volumes.

So in numbers that means:
Final reaction concentration of PEG8000:
1 V: 5.5%  
0.8 V: 4.8%

Final reaction concentration of NaCl:
1V: 0.8 M  
0.8V: 0.7 M

EXTERNAL LINK
https://doi.org/10.1111/1755-0998.12938

THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION
Before starting

Before doing this clean up on your whole (maybe very precious) sample I would recommend testing it out on a smaller amount (300 - 500ng) to make sure the beads work for you.

- For making the beads solution stock take Sera-Mag SpeedBeads (GE Healthcare, PN 24152105050250) out of the fridge and let warm up to room temperature

- For further clean ups always take the beads stock solution out of the fridge at least 15 minutes before usage to let it warm up to room temperature and mix (vortex) very well until solution looks homogeneous and no bead clumps are visible anymore (that is very important)

- Always make fresh 70% Ethanol

- Preheat your elution buffer of choice (TE-Buffer, Tris 10 mM, Water..) to 50° until usage

Make beads stock solution

1 For 10 mL beads stock solution:
<table>
<thead>
<tr>
<th>Final stock</th>
<th>Input</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 mM Tris-HCl</td>
<td>1 M 100 μl</td>
</tr>
<tr>
<td>1 mM EDTA pH 8</td>
<td>0.5 M 20 μl</td>
</tr>
<tr>
<td>1.6 M NaCl</td>
<td>5 M 3.2 ml</td>
</tr>
<tr>
<td>11% PEG 8000</td>
<td>50% (w/v) 2.2 ml</td>
</tr>
<tr>
<td>0.4% beads (v/v)</td>
<td>100% 40 μl</td>
</tr>
<tr>
<td>Milliq Water</td>
<td>4.44 ml</td>
</tr>
</tbody>
</table>

Frist combine only Water, Tris-HCl, EDTA and NaCl in a 50 mL tube.

2 Vortex Ser-Mag SpeedBeads (GE Healthcare, PN 24152105050250) very well and pipette 40 μl into a 1.5 ml tube, put it on the magnetic rack and wait until solution has cleared up and all beads have bound to the back of the tube.

3 Wash beads by removing supernatant and adding 1.5 ml milliq water.

4 Take tube of the magnet, mix well, spin down in a microcentrifuge and put back on the magnet.

5 Wait for beads to assemble at the back of the tube.

6 Pipette off and discard supernatant.

7 Repeat washing (steps 3 - 6) 3 more times.

8 After pipetting of the supernatant the last time take off tube from the magnet and add 40 μl of the previous (step 1) prepared stock solution, mix well, spin down and pipette everything into the remaining stock solution in the 50 mL tube and mix.

9 Now the 2.2 ml 50% PEG can be added to the stock solution, which after vortexing very well is ready for use.

   Be careful to actually pipette 2.2 ml as solution is very viscous, but the final concentration of PEG is crucial for the clean up to work properly.

Clean up

10 Bring your DNA sample in a 1.5 ml tube to comfortable pipetable volume (I usually do it with some volume between 20 - 200 μl) and if you know already that your sample contains a lot of contaminants and/or DNA (hence a really viscous solution) diluting it out and splitting into two tubes can make life easier.

Citation: Miriam Schalamun, Benjamin Schwessinger (07/01/2017). DNA size selection (>1kb) and clean up using an optimized SPRI beads mixture. https://dx.doi.org/10.17504/protocols.io.idmca46.

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(A lot of contaminants can also clump around the beads which makes binding to the magnet slower and sometimes pipetting off the supernatant really difficult)

11 Make sure to know the exact volume of your sample and add 1 V of that (or 0.8 V for removal of more smaller fragments but also higher risk of less recovery) of well homogenised, room temperature beads solution and mix by flicking the tube

12 Place tube on a rotor (or mixer if you don't have one) and mix for 10 minutes

13 Spin down tube in the microcentrifuge and place on the magnet

14 Wait until beads have moved to the back of the tube and the solution becomes clear

   (depending on viscosity of the solution that can take between 1 min to much longer like 1h, if after that the beads look like they are stuck in a big cloud and just don't properly bind to the magnet I add same amount of buffer and beads (the ratio always has to stay the same!) mix again and the put it back on the magnet - usually that solved the problem. But you probably will never have that problem, I just worked with really contaminated samples (plants))

15 Remove and discard supernatant

16 Wash beads with fresh 70% Ethanol by adding 1 - 1.5 mL to the opposite side than where the beads bind and wait 30 seconds

17 Remove and discard Ethanol

18 Repeat washing once more (steps 15 - 18)

   (Don't remove tube from magnet during the washing steps until here)

19 For the last removal of Ethanol make sure that all the Ethanol is removed, therefore take tube off the magnet, spin down for a second and place back onto the magnet, like that also the last drops of Ethanol can be pipetted off

20 Let beads air dry for a maximum of 30 seconds or else elution will be difficult

21 Add 50 ul (or in whatever final volume and concentration the sample is needed) of preheated to 50°C 10 mM Tris (or TE-Buffer)

22 Make sure the beads are resuspend properly by flicking the tube gently and spinning it down - the solution will be...
homogeneous and brown

(I put it in a 50°C Thermoblock for about 20 seconds to encourage the elution reaction)

23 Spin down the tube before placing it on the magnet again and wait until the beads have bound the back of the tube (that can again take its time especially if the sample contains really long DNA fragments) Depending on the sample that will take between 1 minute to a few hours, I usually wait 5 - 10 minutes

24 When the solution has cleared up completely, pipette the supernatant to a fresh tube and discard beads.

Quality control

25 Measure the DNA concentration with a Qubit. The recoveries for HMW DNA > 20kb should lie between 60 - 90%

26 Run a 0.8% agarose gel

80 ng DNA per sample, normalised to 10 ul per lane run for 45 min in 1 x TBE buffer
Lanes:
control: untreated 10kb hyperladder
A: 0.45 AMPure Beads XP beads (Beckman Coulter). This is the standard dilution used in PacBio and Oxford Nanopore protocols.
1: 1 volume of beads solution from this protocol
0.9: 0.9 volumes
0.8: 0.8 volumes