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# Auto-Mag<sup>®</sup> X-Pure Size Select

Version 2.1

Magnetic beads-based chemistry for DNA purification, DNA fragment size selection and cleanup for NGS

**Catalog Number: S003-01, S003-02, S003-03, S003-04**

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## Disclaimers and Safety Information

**This kit is designed for research use only.** All biological samples are considered potentially infectious. When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDSs). MSDS can be downloaded from the “Product Documents” tab when viewing the product kit. Download MSDS at [www.amdbiotech.com](http://www.amdbiotech.com). Information in this document is subject to change without notice.

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## Product Introduction

Auto-Mag® X-Pure Size Select is based on paramagnetic bead technology, and designed for PCR amplicons cleanup, DNA fragments or RNA purification, and sheared DNA size selection and cleanup in the library construction process for next generation sequencing (NGS).

For PCR products purification, the purification protocols provide the procedures for PCR purification in single tube or 96/384 well format. Auto-Mag® X-Pure Size Select reagent utilizes an optimized buffer to selectively bind DNA fragments >100 bp to the beads. Excess primers, nucleotides, salts, and enzymes can be removed using a simple washing procedure and PCR products are eluted with low salt elution buffer or water.

In the DNA size selection, size selection is required to produce a uniform distribution of fragments around an average size. The protocols of DNA size selection describe procedures of DNA size selection by adjusting volume ratio of Auto-Mag® X-Pure Size Select reagent to sheared DNA sample and the desired size DNA fragment are selected and recovered by a simple binding, washing and elution steps.

Auto-Mag® X-Pure Size Select is an RNase free reagent and suitable for RNA cleanup and concentration. The highly purified RNA can be used directly for downstream applications, such as PCR and RT-PCR reactions, RNase protection assays, Transfection for RNAi experiments, Antisense RNA (aRNA) amplification, Probes for microarray or microarray, cDNA synthesis and labeling, as well as RNA-Seq library preparations, etc.

Auto-Mag® X-Pure Size Select is suitable for both manual and fully automated processing and the highly purified DNA can be used directly for downstream applications.

## Features:

- Designed for “bottle swap” with no protocol change against major competitor.
- High recovery of PCR amplicons or dsDNA fragments >100bp
- Single, or double-sided size selection capability with uniform fragments size distribution for NGS
- Rapid and reliable clean-up total RNA, including miRNA, siRNA, and aRNA
- Flexibility of starting sample volumes with the ability to scale and consistent results
- Compatible with manual and automated processes
- Significant cost savings

## Kit Contents

Product Number	S003-01	S003-02	S003-03	S003-04
Auto-Mag® X-Pure Size Select	5 ml	50 ml	250 ml	500 ml

## Storage and Stability

Auto-Mag® X-Pure Size Select is shipped at room temperature and is stable for at least 12 months from the date of purchase when stored at 2-8°C. Contents of the kit should never be frozen at any time.

## Preparation of Reagents

1. Prepare 80% Ethanol for DNA Wash. (Prepare from absolute ethanol. Do not use denatured alcohol).

Fresh prepare 80% ethanol, keep cover tight and use in one week.

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2. Molecular biology grade water, Tris (10 mM, pH 8.0) or TE (10 mM Tris, pH 8.1 mM EDTA) for DNA elution

## Additional Information

### 1. Specifications

Features	Specification
Isolation Technology	Magnetic Beads
Sample Sources	Fragmented DNA, PCR product
Starting amount	Scalable
DNA recovery	>90% recovery for DNA >100bp
Downstream Application	NGS, PCR, Cloning, Nucleic Acid Labeling, Mutation detection, genotyping, etc.
Processing format	Automated or Manual
Storage	2°C - 8°C

### 2. Performing manually without access to a magnet

The compatible magnetic separation device is required to pellet the magnetic particles when performing purification processing. If performing the protocol manually without access to a magnet, sample tubes or plates can be centrifuged for 30 seconds (single tubes: full speed; plates: 3,000 x g) to pellet the magnetic particles and discard the liquid. All processes are to be carried out at room temperature (15–25 °C).

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## Auto-Mag® X-Pure Size Select Protocols

Auto-Mag® X-Pure Size Select can selectively bind DNA fragments based on the ratio of Auto-Mag® X-Pure Size Select reagent to sample. Adjusting the ratio will control to eliminate smaller or larger fragment sizes not within the target range and the desired size DNA Fragment are selected and recovered for fragment library preparation of next-generation sequencing (NGS)

### Materials and Equipment to Be Supplied by User

- Single-tube format: Nuclease-free 1.5-2.0 ml tube, and Magnetic Rack Separator for tube. (Any vendor of choice).
- 96-well format: 96 well thermal cycling plate, or 300µl round bottom microtiter plate, or 1.2 ml deep well microtiter plate and appropriate magnetic separation device ([www.fishersci.com](http://www.fishersci.com) or any vendor of choice).
- Laboratory mixer, vortex, or equivalent.
- 80% ethanol (Prepare from absolute ethanol. Do not use denatured alcohol).
- RNase/DNase free Elution Buffer (AMD-B232 [www.amdbiotech.com](http://www.amdbiotech.com)), or Molecular biology grade water, Tris (10 mM, pH 8.0) or TE (10 mM Tris, pH 8.1 mM EDTA) for DNA elution.
- Well calibrated pipettor and Disposable pipette tips.

### Before Starting

- Prepare 80% Ethanol for DNA wash steps according to the instructions of Preparation Reagents on page 3.
- Bring the Auto-Mag® X-Pure Size Select reagent bottle to room temperature for at least 30 minutes before use.
- Shake thoroughly the Auto-Mag® X-Pure Size Select reagent to fully resuspend the magnetic beads.

### Sample preparation.

- DNA samples should be fragmented double-stranded DNA and dissolved in molecular biology grade water or lower salt buffer solution.
- For best results, the sample volume should be  $\geq 50\mu\text{l}$ . A lower volume will decrease pipetting accuracy, therefore increasing selection point variability.
- For a Left Side-Size Selection, the majority of DNA fragment size distribution should be larger than the selected cutoff point.
- for a Right-Side Size Selection, the majority of DNA fragment size distribution should be smaller than selected cutoff point.
- For a Double Size Selection, most of size distribution should be centered between the selection points and in general, the range of DNA fragments may be no smaller than 100 bp and no larger than 800 bp.

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## Protocol for fragmented DNA Double Size Selection

For NGS library preparation, sheared DNA fragments need to be cleaned up and selected appropriate size. To perform Double Size Selection, the upper cut-off is performed first to remove larger fragments and then, the subsequent lower cut-off binds all fragments greater than the specified size to the beads whilst unwanted smaller fragments remain in the supernatant and are discarded. At this point, fragments within the desired range are bound to the beads and recover after the standard washing and elution steps.

To determine the DNA fragment selection range and corresponding ratio of Auto-Mag® X-Pure Size Select reagent, refer to Table 1.

Table 1: Reference conditions for DNA Double Size Selective for NGS library preparation

The average size of selection (bp)	150-220	200-300	250-320	280-350	300-400	400-550	500-700
1 <sup>st</sup> ratio of Auto-Mag® X-Pure Size Select / DNA	1.0x	0.9x	0.8x	0.7x	0.6x	0.55x	0.5x
2 <sup>nd</sup> ratio of Auto-Mag® X-Pure Size Select / DNA	0.2x	0.2x	0.2x	0.2x	0.2x	0.15x	0.15x
Total ratio	1.2x	1.1x	1.0x	0.9x	0.8x	0.7x	0.65x

1. Completely float the Auto-Mag® X-Pure Size Select reagent until it appears homogeneous in color.
2. Transfer 50µl sheared DNA sample into a 1.5-2.0 ml RNase-DNase free tube or well of 96-well plate. If sample is less than 50µl, add reagent grade water, or 10mM Tris-HCl pH8.0, to ensure the starting volume is at least 50µl.
3. Reference Table 1, first add the required volume of Auto-Mag® X-Pure Size Select reagent to the sample for remove unwanted larger DNA fragment.

*Note: Volume of sample \* ratio = volume of Auto-Mag® X-Pure Size Select reagent*

*For example: for 250-320bp fragment select, 50µl sample \* 0.8 ration = 40µl of Auto-Mag® X-Pure Size Select NGS reagent*

4. Mix the Auto-Mag® X-Pure Size Select reagent and the samples thoroughly by pipette 10 times or vortex for 10 seconds.

*Note: Insufficient mixing will lead to inconsistent size selection results. Make sure to mix well. Pipette mixing is preferable as it tends to be more reproducible. If a 96 well plate and vortex is used, the plate must be sealed with a plate seal before vortex.*

5. Incubate samples at room temperature for 5 minutes.
6. Place the sample tube or plate on a compatible magnetic separation device for 5 minutes or until the solution is completely clear. The Auto-Mag® X-Pure Size Select bead settles to the magnet.

*Note: The higher sample volume and higher Auto-Mag® X-Pure Size Select ratio will require a longer settling time.*

7. With the sample tube or plate on the magnet, transfer clear supernatant, which contains the Size Selected sample, into a new tube or plate. The reaction tube or plate with the remaining beads can be discarded.

*Note: Do not disturb the attracted magnetic beads while transferring the supernatant. As the undesired larger fragment sizes are associated with the beads, significant bead transfer will cause tailing into the larger size range.*

8. Reference Table 1, add an additional required volume of Auto-Mag® X-Pure Size Select reagent to the

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supernatant from step 6. This will bind the DNA fragments in the supernatant to the new Auto-Mag® X-Pure Size Select NGS beads.

*Note: Volume of sample \* 0.2 ratio = volume of Auto-Mag® X-Pure Size Select reagent*

*For example: for 250-320bp fragment select, 50µl sample \* 0.2 = 10µl of Auto-Mag® X-Pure Size Select reagent. The total ratio of Auto-Mag® X-Pure Size Select reagent suspension to the original sample is 1.0 x now. (40µl + 10µl) / 50µl = 1.0x.*

9. Mix the Auto-Mag® X-Pure Size Select reagent and the samples thoroughly by pipette 10 times or vortex for 10 seconds.

*Note: Insufficient mixing will lead to inconsistent size selection results. Make sure to mix well. Pipette mixing is preferable as it tends to be more reproducible. If a 96 well plate and vortex is used, the plate must be sealed with a plate seal before vortex.*

10. Incubate samples at room temperature for 5 minutes.

11. Place the sample tube or plate on a compatible magnetic separation device for 2 minutes or until the solution is completely clear. The Auto-Mag® X-Pure Size Select bead settles to the magnet.

*Note: The higher sample volume and higher Auto-Mag® X-Pure Size Select ratio will require a longer settling time.*

12. Remove and discard the clear supernatant, which contains the unwanted smaller DNA fragments.

*Note: Do not disturb the attracted magnetic beads. As the desired library is associated with the beads, any bead loss during this step will result in a reduced yield.*

13. With the samples still on the magnet, add 180µl of 80% ethanol to each sample and incubate at room temperature for 30 seconds. Do not disturb the attracted beads. Mixing is not necessary.

14. With the sample still on the magnet, Remove and discard the ethanol supernatant by pipetting. Do not disturb the attracted beads.

15. Repeat steps 13-14 for the second 80% ethanol wash.

16. Dry the beads by incubating the plate at room temperature for 5 minutes with the sample still on the magnetic separation device. Remove any residue liquid with a pipettor.

*Note: It is critical to completely remove all traces of alcohol but take caution in not over drying the beads as this will reduce the yield.*

17. Remove the sample tubes or plate from the magnet. Add >20µl of elution buffer (reagent grade water, or TE buffer) and mix by pipet up and down 10 times.

*Note: Depending on the downstream application, you can add any volume of Elution Buffer to elute the DNA. However, Elution volume should be large enough so that the liquid level is high enough for the beads to settle to the magnet.*

18. Incubate the sample at room temperature for 5 minutes.

19. Place the sample back on the magnet and wait 2 minutes or until the magnetic beads are completely cleared from solution.

20. Transfer the eluate (size selected DNA fragments) to an appropriate storage vessel and keep at -20°C for long term storage or for subsequent applications.

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## Protocol for removing adapter dimers from NGS library preparation samples

In the NGS library preparation, Adapter dimers contain full-length adapter sequences that are able to bind and cluster on the flow cell and generate sequencing data. they can negatively impact sequencing data quality and may even cause a run to stop prematurely. If adapter dimers are present in the library, it need to perform an additional clean-up step.

1. Completely float the Auto-Mag® X-Pure Size Select reagent until it appears homogeneous in color.
2. Conform the sample volume, Transfer sample into a 1.5-2.0 ml RNase-DNase free tube or well of 96-well plate.
3. Reference Table 2, Add 0.9x volume of Auto-Mag® X-Pure Select reagent to the adapter addition reaction sample.

Table 2: Some common adapter addition reaction volumes and suggested Auto-Mag® X-Pure Size Select volumes.

Adapter Addition Reaction Sample Volume (μl)	Auto-Mag® X-Pure Size Select Volume Needed (μl) *
20	18
25	2.5
35	31.5
50	45
65	58.5

*(Volume of Auto-Mag® X-Pure Size Select reagent per reaction) = 0.9 X (sample volume).*

4. Mix the Auto-Mag® X-Pure Size Select reagent and the samples thoroughly by pipette 10 times or vortexing for 10 seconds.
- Note: If a 96 well plate and vortexing is used, the plate must be sealed with a plate seal before vortexing.*
5. Incubate samples at room temperature for 5 minutes for maximum recovery.
  6. Place the sample tubes or plate on a compatible magnetic separation device for 2 minutes or until the solution clears. Beads will pull to the side of the well.
  7. With the sample still on the magnet, remove and discard the supernatant by pipetting. Do not disturb the attracted beads.
  8. With the sample on the magnet, add 200μl of 80% ethanol to each sample and incubate at room temperature for 30 seconds. Mixing is not necessary.

9. With the sample still on the magnet, remove and discard the supernatant by pipetting.

*Note: If the total volume of sample plus reagent exceeds 200μl, then use a wash volume of at least the volume of sample plus reagent.*

10. Repeat steps 8-9 for a second 80% ethanol wash.
11. Dry the beads by incubating the plate at room temperature for 5 minutes with the samples still on the magnetic separation device. Remove any residue liquid with a pipettor.

*Note: It is critical to completely remove all traces of alcohol but take caution in not over drying the beads as this will significantly decrease elution efficiency.*

12. Remove the sample tubes or plate from the magnet. Add >20μl of elution buffer (reagent grade water, or TE

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buffer) and mix by pipet up and down 10 times.

*Note: Depending on the downstream application, you can add any volume of Elution Buffer to elute the DNA. However, Elution volume should be large enough so that the liquid level is high enough for the beads to settle to the magnet.*

13. Incubate the sample at room temperature for 2 minutes.
14. Place the sample back on the magnet and wait 2 minutes or until the magnetic beads are completely cleared from solution.
15. Transfer the eluate to an appropriate storage vessel and keep at -20°C for long term storage or for subsequent applications.

### Protocol for PCR Amplicon Cleanup: 96 Well Format, or Single-Tube Format

1. Completely float the Auto-Mag® X-Pure Size Select reagent until it appears homogeneous in color.
2. Confirm the volume of PCR reaction and determine whether a plate transfer is necessary for 96 well plate, or transfer the sample to a new 1.5-2.0 ml tube.

*Note: PCR plates generally have a maximum volume of 200 µl. If the reaction volume multiplied by 2.8 exceeds the volume of the PCR plate, transfer PCR reaction to a 300ul round plate or a 1.2 ml deep-well plate.*

3. Reference Table 3, add the appropriate volume of Auto-Mag® X-Pure Size Select reagent to the PCR sample.

Table 3: PCR sample volumes and suggested Auto-Mag® X-Pure Size Select volumes.

PCR Reaction Volume (µl)	Auto-Mag® X-Pure Size Select Volume at 1.8x (µl) *
10	18
20	36
25	45
50	90
100	180
<i>(Volume of Auto-Mag® X-Pure Size Select reagent per reaction) = 1.8 X (PCR reaction volume).</i>	

4. Mix the Auto-Mag® X-Pure Size Select reagent and the samples thoroughly by pipette 10 times or vortexing for 10 seconds.

*Note: This step binds DNA fragments 100 bp and larger to the magnetic beads.  
If a 96 well plate and vortexing is used, the plate must be sealed with a plate seal before vortexing.*

5. Incubate samples at room temperature for 5 minutes for maximum recovery.
6. Place the sample tubes or plate on a compatible magnetic separation device for 2 minutes or until the solution clears. Beads will pull to the side of the well.
7. With the sample still on the magnet, remove and discard the supernatant by pipetting. Do not disturb the attracted beads.
8. With the sample on the magnet, add 200µl of 80% ethanol to each sample and incubate at room temperature for 30 seconds. Mixing is not necessary.

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9. With the sample still on the magnet, remove and discard the supernatant by pipetting.

*Note: If the total volume of sample plus reagent exceeds 200µl, then use a wash volume of at least the volume of sample plus reagent.*

10. Repeat steps 8-9 for a second 80% ethanol wash.

11. Dry the beads by incubating the plate at room temperature for 5 minutes with the samples still on the magnetic separation device. Remove any residue liquid with a pipettor.

*Note: It is critical to completely remove all traces of alcohol but take caution in not over drying the beads as this will significantly decrease elution efficiency.*

12. Remove the sample plate or tubes from the magnet. Add 20-50µl of elution buffer (reagent grade water, or TE buffer) to each sample and mix by pipet up and down 10 times.

*Note: Prewarming the elution buffer to 55°C can increase the yield.*

13. Incubate the sample at room temperature for 2 minutes.

14. Place the sample back on the magnet and wait 2 minutes or until the magnetic beads are completely cleared from solution.

15. Transfer the eluate to an appropriate storage vessel and keep at -20°C for long term storage or for subsequent applications.

### Protocol for Total RNA Cleanup (single-tube or 96-well format)

**This Protocol will purify and recover all of RNA molecules including miRNA, siRNA, and aRNA from pre-purified total RNA or enzyme reactions.**

1. Shake thoroughly the Auto-Mag® X-Pure Size Select reagent to fully resuspend the magnetic beads.
2. Confirm the volume of RNA sample and transfer the sample to a 1.5-2.0 ml tube, or the well of 96 well plate (96 well format)

*Note: PCR plates generally have a maximum volume of 200µl. If the RNA sample volume multiplied by 5 exceeds 200µl, transfer sample to a 500ul round plate or a 1.2 ml deep-well plate.*

3. Add the appropriate volume of Auto-Mag® RNA-Pure reagent and 100% isopropanol according to the sample volume shown in table.

Sample Volume (µl)	Auto-Mag® X-Pure Size Select Volume (µl)*	100% Isopropanol (µl)**
10	18	20
25	45	50
50	90	100
100	180	200
<i>*(Volume of Auto-Mag® X-Pure Size Select reagent needed per reaction) = 1.8 X (RNA sample volume)</i>		<i>**Volume of isopropanol needed per reaction = 2 X (RNA sample volume)</i>

4. Mix the total reaction volume by pipetting up and down 10 times or vortex for 30 seconds.

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*Note: Pipette mixing is preferable. If a 96 well plate and vortex is used, the plate must be sealed with a plate seal before vortex.*

5. Incubate samples at room temperature for 5 minutes for maximum recovery.
6. Place the sample tubes or plate on a compatible magnetic separation device for 2 minutes or until the solution clears. Beads will pull to the side of the well.
7. With the sample still on the magnet, remove and discard the supernatant by pipetting. Do not disturb the attracted beads.
8. With the sample on the magnet, add 200 $\mu$ l of 80% ethanol to each sample and incubate at room temperature for 30 seconds. Mixing is not necessary.
9. With the sample still on the magnet, remove and discard the supernatant by pipetting.

*Note: If the total volume of sample plus reagent exceeds 200 $\mu$ l, then use a wash volume of at least the volume of sample plus reagent.*

10. Repeat steps 8-9 for a second 80% ethanol wash.
11. Dry the beads by incubating the plate at room temperature for 5 minutes with the samples still on the magnetic separation device. Remove any residue liquid with a pipettor.

*Note: It is critical to completely remove all traces of alcohol but take caution in not over drying the beads as this will significantly decrease elution efficiency.*

12. Remove the sample plate or tubes from the magnet. Add 20-50 $\mu$ l of elution buffer (reagent grade water, or TE buffer) to each sample and mix by pipet up and down 10 times.

*Note: Prewarming the elution buffer to 55°C can increase the yield.*

13. Incubate the sample at room temperature for 2 minutes.
14. Place the sample back on the magnet and wait 2 minutes or until the magnetic beads are completely cleared from solution.
15. Transfer the eluate to an appropriate storage vessel and keep at -20°C for long term storage or for subsequent applications.

## **Customized Protocol and Programmed Procedure for Automation Purification**

To obtain a custom protocol for DNA fragment size selection of a specific fragment size; or automating this procedure on a liquid handler or a magnetic processor, please contact AMD Biotech for instrument-specific instructions or additional processing procedures.

Phone: 404-290-5063 (in US), Email: [support@amdbiotech.com](mailto:support@amdbiotech.com)

## Troubleshooting Guide

Please use this guide to troubleshoot any problems that may arise. For further assistance, please contact technical support via Phone: 1-404-290-5063 (in US), Email: [support@amdbiotech.com](mailto:support@amdbiotech.com)

Observation	Possible Causes	Comments
Low yield/ Incorrect recovery of purification	Recovery was measured by Spectrophotometry Absorbance. This causes the recovery to appear lower than it is.	Run sample on an agarose gel to double check the recovery measurement or use a quantitative double-stranded DNA assay reagent such as PicoGreen assay.
	Bead Loss	If beads get aspirated into tips during supernatant removal, the nucleic acid bound to these beads will also be lost. Aspirate slowly and remove as much of the first supernatant as possible without disturbing the bead.
	Insufficient Mixing	Mixing thoroughly during the initial bind mix and elution mix is critical. to ensure the beads get sufficiently resuspended.
	Large Reaction Volume	Large volume reactions can benefit from an extended binding and separation time. Increase binding time to 10 minutes and ensure all beads are separated before removing the supernatant.
	Low Elution Volume	A small elution volume leads to a decrease in recovery. This is because a small amount of elution buffer always stays behind coating the beads. To increase the elution volume.
Fragment size incompatible	Insufficient ratio	Use volume ratios outlined in this manual.
	Insufficient ethanol concentration used for washing step	Use freshly prepared 80% ethanol. Over time ethanol becomes more diluted through evaporation and absorption of atmospheric water. Therefore, parts of the DNA pellet go into solution and DNA fragments are washed away.
	Elution buffer volume insufficient	Bead pellet must be covered completely with elution buffer
	Beads over dried	Do not dry beads for longer than 15 minutes at room temperature. Over drying of beads may result in lower elution efficiencies.
Downstream applications are unsuccessful	Carry-over of ethanol from washing step	Be sure to remove all the ethanol after the final wash step. Dry beads 5-10 minutes at room temperature.
Carry-over of beads	Time for magnetic separation too short	Increase separation time to allow the beads to be attracted to the magnetic pins completely

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## Ordering Information

Product Description	Catalog No.	Size
Auto-Mag® X-Pure Size Select	S003-01	5 ml
	S003-02	50 ml
	S003-03	250 ml
	S003-04	500 ml
Auto-Mag® DNA Elution Buffer	B228-01	50 ml
	B228-02	250 ml
	B228-03	500 ml

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