



## Genomic DNA Clean & Concentrator<sup>®</sup>-25

Clean and concentrate large-sized DNA from any enzymatic reaction or impure preparation.

#### Highlights

- Quick (5 minute) spin column recovery of large-sized DNA (e.g., genomic, mitochondrial, plasmid (BAC/PAC), viral, phage, (wga) DNA, etc.) from any enzymatic reaction or impure preparation (e.g., Proteinase K digestion). No messy precipitations!
- Unique spin column for low volume (≥35 µl) elution of ultra-pure, highyield DNA.
- · Eluted DNA is ideal for PCR, restriction endonuclease digestion, Sanger and Next-Gen sequencing, etc.

Catalog Numbers: D4064, D4065



Scan with your smart-phone camera to view the online protocol/video.







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# **Product Contents**

| Genomic DNA Clean &<br>Concentrator <sup>®</sup> -25 | <b>D4064</b><br>(25 Preps.) | <b>D4065</b><br>(100 Preps.) | Storage<br>Temperature |
|--|-----------------------------|------------------------------|------------------------|
| ChIP DNA Binding Buffer                              | 50 ml                       | 2 x 50 ml                    | Room Temp.             |
| DNA Wash Buffer <sup>1</sup>                         | 6 ml                        | 24 ml                        | Room Temp.             |
| DNA Elution Buffer                                   | 4 ml                        | 10 ml                        | Room Temp.             |
| Zymo-Spin™ IIC-XLR Columns                           | 25                          | 100                          | Room Temp.             |
| Collection Tubes                                     | 50                          | 100                          | Room Temp.             |
| Instruction Manual                                   | 1                           | 1                            | -                      |

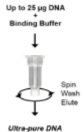
<sup>&</sup>lt;sup>1</sup> Ethanol must be added prior to use as indicated on **DNA Wash Buffer** label.

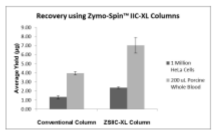
## **Specifications**

- **DNA Purity** High-quality (*A*<sub>(260/280)</sub> ≥ 1.8) high molecular weight DNA ideal for ligation, sequencing, labeling, PCR, microarray, transfection, transformation, and restriction digestion procedures.
- **DNA Size Limits** Capable of purifying small DNA fragments >50 bp and large sized DNAs up to 200 kb.
- **DNA Recovery** Typically, up to 25 µg total DNA per column can be eluted into ≥35 µl of low salt **DNA Elution Buffer** or water.
- Sample Sources DNA from impure preparations of genomic DNA (e.g., Proteinase K digestions), plasmid DNA (including BAC), viral DNA, and whole genome amplified (wga) DNA. Can also be used for the purification of low molecular weight DNA (50 bp to 10 kb) from PCR, endonuclease digestion, post-RT cDNA synthesis, etc. Suitable for isolated DNA stored in DNA/RNA Shield (page 6).
- **Product Detergent Tolerance** ≤ 5% Triton X-100, ≤ 5% Tween-20, ≤ 5% Sarkosyl, ≤ 1% SDS.

## **Product Description**

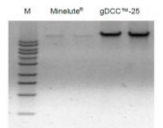
The Genomic DNA Clean & Concentrator<sup>®</sup>-25 (DCC<sup>®</sup>) is for quick (5 minute) recovery of ultra-pure, large-sized DNA (*e.g.*, genomic, mitochondrial, plasmid (BAC/PAC), viral, phage, (wga) DNA, etc.) from any enzymatic reaction or impure preparation (*e.g.*, Proteinase K digestion). There is no need for organic denaturants, chloroform, or messy precipitations: simply add the specially formulated ChIP DNA Binding Buffer to a sample and then transfer the mixture to the supplied Zymo-Spin<sup>™</sup> Column. Eluted DNA is suitable for sequencing, PCR, endonuclease digestion, and other enzymatic procedures. The product is also compatible with smaller DNAs (50 bp to 10 kb) from PCR, digestions, crude plasmid preparations, cDNA synthesis, *etc*.





Five minute Genomic DCC®-25 procedure.

Zymo-Spin<sup>™</sup> IIC-XLR Columns result in superior yields compared to conventional columns. Genomic DNA extracted using the Zymo-Spin<sup>™</sup> IIC-XLR Column results in higher yields from HeLa Cells and Porcine Whole Blood.



High molecular weight DNA is efficiently purified using the Genomic DCC<sup>®</sup>-25. Lambda ( $\lambda$ ) phage DNA (48.5 kb) was purified (in duplicate) from input material using Qiagen's Minelute<sup>®</sup> and the Genomic DCC<sup>®</sup>-25 (gDCC<sup>®</sup>-25). The gDCC<sup>®</sup>-25 resulted in yields > 40% compared to the Minelute<sup>®</sup>. Eluted DNAs were analyzed in a 0.8% (w/v) TAE/agarose/EtBr gel. The size marker "M" is a 1 kb ladder (Zymo Research).

## Formats

|          | Genomic<br>DCC™-10  | Genomic<br>DCC™-25   | ZR-96 Genomic<br>DCC™-5 |
|----------|---------------------|----------------------|-------------------------|
|          |                     |                      |                         |
| Column   | Zymo-Spin™<br>IC-XL | Zymo-Spin™<br>IIC-XL | Zymo-Spin™<br>I-96-XL   |
| Capacity | 10 µg/ prep.        | 25 µg/ prep.         | 5 µg/ prep.             |
| Elution  | ≥ 10 µl             | ≥ 35 µl              | ≥ 15 µl                 |

## Applications

| Post-PCR DNA Clean-up                    | Efficient desalting of DNA with the removal of DNA polymerases, primers, and free dNTPs.   |
|--|--|
| DNA Clean-up From<br>Enzymatic Reactions | Efficient desalting of DNA with the removal of modifying enzymes,<br>RNA polymerases, ligases, kinases, nucleases, phosphatases,<br>endonucleases, <i>etc</i> .  |
| Plasmid DNA Clean-up                     | Efficiently purifies plasmid DNA from "home-made" preparations<br>of cell free lysates or from commercial kits. Plasmid DNA purified<br>and concentrated using the <b>Genomic DCC®</b> has proven an<br>excellent substrate for high quality DNA sequencing.             |
| Isotope and Dye Removal                  | Efficiently removes unincorporated fluorescent ( <i>i.e.</i> , AMCA, FITC, BIO, DIG, Cy3, Cy5, FAM, <i>etc.</i> ) and radiolabeled dNTP derivatives from DNA following <i>in vitro</i> labeling reactions.   |
| Genomic DNA Clean-Up                     | Efficiently purifies genomic DNA from "home-made" preparations<br>of cell free lysates or from commercial kits. Genomic DNA purified<br>and concentrated using the <b>Genomic DCC</b> <sup>®</sup> has proven an<br>excellent substrate for high quality DNA sequencing. |

- ✓ For purification of DNA from 50 bp to 23 kb, use the DNA Clean & Concentrator (D4003 & D4013).
- ✓ For purification of short DNA or RNA oligonucleotides ≥ 16 nt, use the Oligo Clean & Concentrator (D4060, D4061).
- ✓ For ChIP (Chromatin Immunoprecipitation) sample cleanup, use the ChIP DNA Clean & Concentrator<sup>®</sup> (D5201, D5205) for high quality DNA from any step in a standard ChIP protocol.
- ✓ For post-cycle sequencing samples, use the ZR Sequencing DNA Clean-up Kit™ (D4050, D4051) for dye blob elimination.
- ✓ For samples containing PCR inhibitors, use the OneStep<sup>™</sup> PCR Inhibitor Removal Kit (D6030, D6035).

## Protocol

#### **Buffer Preparation**

✓ <u>Before starting</u>: Add 24 ml 100% ethanol (26 ml 95% ethanol) to the 6 ml DNA Wash Buffer concentrate. Add 96 ml 100% ethanol (104 ml 95% ethanol) to the 24 ml DNA Wash Buffer concentrate.

### Sample Processing

All centrifugation steps should be performed between 10,000 - 16,000 x g.

 In a 1.5 ml microcentrifuge tube, add 2-5 volumes of ChIP DNA Binding Buffer to each volume of DNA sample<sup>1</sup> (see table below). Mix thoroughly.

| Application                  | DNA Binding Buffer : Sample | Example         |
|------------------------------|-----------------------------|-----------------|
| Plasmid, genomic DNA (>2 kb) | 2 : 1                       | 200 µl : 100 µl |
| PCR product, DNA fragment    | 5 : 1                       | 500 µl : 100 µl |

- 2. Transfer mixture to a provided **Zymo-Spin™ IIC-XLR Column**<sup>2</sup> in a **Collection Tube**.
- 3. Centrifuge for 30 seconds. Discard the flow-through.
- 4. Add 400 µl **DNA Wash Buffer** to the column. Centrifuge for 1 minute. Empty the **Collection Tube**. Repeat the wash step.
- 5. Transfer the column to a 1.5 ml microcentrifuge tube. Add  $\geq$  50 µl<sup>3</sup> **DNA Elution Buffer**<sup>4</sup> or water<sup>5</sup> directly to the column matrix and incubate at room temperature for five minutes and centrifuge for 30 seconds to elute the DNA.

Ultra-pure DNA is now ready for use.

<sup>&</sup>lt;sup>1</sup> It may be necessary to add RNase A to cell lysates <u>prior</u> to performing the procedure to ensure RNA-free DNA will be recovered in Step 5.

<sup>&</sup>lt;sup>2</sup>The sample capacity of the column is 900 ul. It may be necessary to load and spin a column multiple times if a sample has a volume larger than 900 ul.

<sup>&</sup>lt;sup>3</sup> To increase concentrations of first elution, use  $\geq$  35 µl **DNA Elution Buffer**. To increase yield, load the eluate directly on the matrix a second time, incubate for 3 minutes at room temperature, and centrifuge again.

<sup>&</sup>lt;sup>4</sup> DNA Elution Buffer: 10mM Tris-HCl, pH 8.5, 0.1mM EDTA

<sup>&</sup>lt;sup>5</sup> Elution of DNA from the column is dependent on pH and temperature. If water is used, make sure the pH is >6.0. The total yield may be improved by eluting the DNA with 60-70°C DNA Elution Buffer.

## Appendix

### Isolated DNA stored in DNA/RNA Shield

For previously isolated/purified DNA stored in **DNA/RNA Shield**, use the following protocol to recover ultra-pure DNA, ready for downstream applications.

- 1. If frozen, thaw samples<sup>1</sup> at room temperature (20-30°C).
- 2. Add an equal volume of ethanol (95-100%) to the sample and mix well.
- 3. Continue with Step 2 of the Sample Processing Protocol on page 5.

### **RNase A Treatment**

Dissolve RNase A (E1008-30), sold separately, in DNase/RNase-free water or TE to a stock concentration of 10 mg/ml.

- 1. Add enough 10 mg/ml RNase A to the sample for a final concentration of 10-100  $\mu$ g/mL and mix well.
- 2. Incubate at room temperature for 15 minutes.
- 3. Continue with step 1 of the Sample Processing protocol on page 5.

<sup>&</sup>lt;sup>1</sup>Adjust the sample volume to 50 µl (minimum) with DNA/RNA Shield.

# Troubleshooting

| Problem  | Possible Causes and Suggested Solutions  |
|--|--|
|  | <b>Improperly Prepared/Stored DNA Wash Buffer.</b><br>Make sure ethanol has been added to the <b>DNA Wash Buffer</b> concentrate. Cap the bottle tightly to prevent evaporation over time.   |
| Low Recovery   | Addition of DNA Elution Buffer. Add elution buffer directly to the column matrix, not to the walls of the column. Elution buffer requires contact with the matrix for at least 1 minute for large DNA $\geq$ 10 kb.  |
|  | <b>Incomplete Elution.</b> DNA elution is dependent on pH, temperature, and time. For large genomic DNA ( $\geq$ 50 kb), apply heated elution buffer (60-70 °C) to the column and incubate for several minutes prior to elution. Sequential elutions may be performed for quantitatively higher recovery but lower final DNA concentration. This is recommended for DNA $\geq$ 10 kb.  |
| Low A <sub>260</sub> /A <sub>230</sub> ratio                                   | <b>Column tip contaminated.</b> When removing the column from the collection tube, be careful that the tip of the column does not come into contact with the flowthrough. Trace amounts of salt from the flowthrough can contaminate a sample resulting in a low $A_{260}/A_{230}$ ratio. Ethanol contamination from the flowthrough can also interfere with DNA elution. Zymo-Spin <sup>TM</sup> columns are designed for complete elution with no buffer retention or carryover. |
| Following Clean-up<br>with DCC®, Multiple<br>Bands Appear in an<br>Agarose Gel | Acidification of DNA Loading Dye. Most loading dyes do not contain EDTA and will acidify ( $pH \le 4$ ) over time due to some microbial growth. This low pH is enough to cause DNA degradation. Therefore, if water is used to elute the DNA, 6X Loading Dye containing 1 mM EDTA is recommended.  |

# **Ordering Information**

| Product Description   | Catalog No. | Size          |
|---|-------------|---------------|
| Genomic DNA Clean & Concentrator®-10                          | D4010       | 25 Preps.     |
| (for purification of up to 10 µg genomic DNA per prep.)       | D4011       | 100 Preps.    |
| Genomic DNA Clean & Concentrator®₋25                          | D4064       | 25 Preps.     |
| (for purification of up to 25 µg genomic DNA per prep.)       | D4065       | 100 Preps.    |
| ZR-96 Genomic DNA Clean & Concentrator®-5                     | D4066       | 2 x 96 Preps. |
| (for 96-well purification of up to 5 µg genomic DNA per well) | D4067       | 4 x 96 Preps. |

| Individual Kit Components     | Catalog No.                         | Amount                           |
|-------------------------------|-------------------------------------|----------------------------------|
| Chip DNA Binding Buffer       | D5201-1-50                          | 50 ml                            |
| DNA Wash Buffer (concentrate) | D4003-2-6<br>D4003-2-24             | 6 ml<br>24 ml                    |
| DNA Elution Buffer            | D3004-4-1<br>D3004-4-4              | 1 ml<br>4 ml                     |
| Collection Tubes              | C1001-50<br>C1001-500<br>C1001-1000 | 50 Pack<br>500 Pack<br>1000 Pack |
| Zymo-Spin™ IIC-XLR Columns    | C1104-25                            | 25 Pack                          |

## **Complete Your DNA Methylation Workflow**

#### ✓ Rapid Method for Complete Bisulfite Conversion of DNA

| EZ DNA Methylation Kits                     | Size  | Catalog No.    |
|---|---|----------------|
| EZ DNA Methylation-Lightning Kit            | 50 Rxns.<br>200 Rxns.                               | D5030<br>D5031 |
| EZ-96 DNA Methylation-Lightning Kit         | 2x96 Rxns. (Deep-Well)<br>2x96 Rxns. (Shallow-Well) | D5032<br>D5033 |
| EZ DNA Methylation-Lightning Automation Kit | 96 Rxns.  | D5049          |
| EZ-96 DNA Methylation Lightning MagPrep     | 4 X 96 Rxns.<br>8 X 96 Rxns.                        | D5046<br>D5047 |

#### ✓ Innovative Solutions for Next Generation Sequencing

| Library Prep Kits                | Size                   | Catalog No.    |
|----------------------------------|------------------------|----------------|
| Zymo-Seq WGBS Library Kit        | 24 Preps.              | D5465          |
| Pico Methyl-Seq Library Prep Kit | 10 Preps.<br>25 Preps. | D5455<br>D5456 |
| Zymo-Seq RRBS Library Kit        | 24 Preps.<br>48 Preps. | D5460<br>D5461 |

#### ✓ Optimal Amplification of Bisulfite-Treated DNA

| ZymoTaq Polymerase     | Size                  | Catalog No.    |
|------------------------|-----------------------|----------------|
| ZymoTaq Premix         | 50 Rxns.<br>200 Rxns. | E2003<br>E2004 |
| ZymoTaq DNA Polymerase | 50 Rxns.<br>200 Rxns. | E2001<br>E2002 |
| ZymoTaq qPCR Premix    | 50 Rxns.<br>200 Rxns. | E2054<br>E2055 |

#### ✓ Industry Leading Tools for Assessing Your DNA Methylation Workflow

| DNA Methylation Standards                                      | Size           | Catalog No.    |
|--|----------------|----------------|
| Human Methylated & Non-methylated DNA Set                      | 5 µg/20 µl     | D5014          |
| Universal Methylated DNA Standard                              | Human<br>Mouse | D5011<br>D5012 |
| Bisulfite-Converted Universal Methylated<br>Human DNA Standard | 1 µg/50 µl     | D5015          |
| Human Methylated & Non-Methyated (WGA) DNA Set                 | 5 µg/20 µl     | D5013          |

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Integrity of kit components is guaranteed for up to one year from date of purchase. Reagents are routinely tested on a lot-to-lot basis to ensure they provide the highest performance and reliability.

This product is for research use only and should only be used by trained professionals. It is not for use in diagnostic procedures. Some reagents included with this kit are irritants. Wear protective gloves and eye protection. Follow the safety guidelines and rules enacted by your research institution or facility.

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## The BEAUTY of SCIENCE is to Make Things $\textbf{SIMPLE}^{\text{\tiny 8}}$





