Oxidative damage to DNA is a result of the interaction of DNA with reactive oxygen species (ROS), in particular, the hydroxy radical which is converted from superoxide and hydrogen peroxide by the Fenton reaction. Hydroxy radicals produce a multiplicity of modifications in DNA. Oxidative attack by hydroxy radical on the deoxyribose moiety will lead to the release of free bases from DNA, generating strand breaks with various sugar modifications and simple abasic sites (AP sites). In fact, AP sites are one of the major types of damage generated by ROS. It has been estimated that endogenous ROS can result in about 2x10^5 base lesions per cell per day.

Aldehyde Reactive Probe (ARP) reagent (N'-aminooxymethylcarbonylhydrazino-D-biotin, Fig. 1) reacts specifically with an aldehyde group which is the open ring form of the AP sites. This reaction makes it possible to detect DNA modifications that result in the formation of an aldehyde group. After treating DNA containing AP sites with ARP reagent, AP sites are tagged with biotin residues. By using an excess amount of ARP, all AP sites can be converted to biotin-tagged AP sites. Therefore, AP sites can be quantified using avidin-biotin assay followed by a colorimetric detection of peroxidase or alkaline phosphatase conjugated to the avidin (Fig. 2).

DNA Damage Quantification Kit contains all the necessary solutions, enabling the determination of 1 to 40 AP sites per 1x10^5 bp.

**Kit Contents**

<table>
<thead>
<tr>
<th>Item</th>
<th>5 samples</th>
<th>20 samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>• ARP-DNA Standard Solution</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 ARP-DNA Standard Soln.* (0 ARP/100,000 bp)</td>
<td>250 µl x 1</td>
<td>250 µl x 1</td>
</tr>
<tr>
<td>2.5 ARP-DNA Standard Soln.* (2.5 ARP/100,000 bp)</td>
<td>250 µl x 1</td>
<td>250 µl x 1</td>
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<tr>
<td>5 ARP-DNA Standard Soln.* (5 ARP/100,000 bp)</td>
<td>250 µl x 1</td>
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<td>10 ARP-DNA Standard Soln.* (10 ARP/100,000 bp)</td>
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<tr>
<td>20 ARP-DNA Standard Soln.* (20 ARP/100,000 bp)</td>
<td>250 µl x 1</td>
<td>250 µl x 1</td>
</tr>
<tr>
<td>40 ARP-DNA Standard Soln.* (40 ARP/100,000 bp)</td>
<td>250 µl x 1</td>
<td>250 µl x 1</td>
</tr>
<tr>
<td>• ARP Solution (10 mmol/l ARP)</td>
<td>100 µl x 1</td>
<td>250 µl x 1</td>
</tr>
<tr>
<td>• DNA Binding Solution</td>
<td>10 ml x 1</td>
<td>10 ml x 1</td>
</tr>
<tr>
<td>• Substrate Solution</td>
<td>10 ml x 1</td>
<td>10 ml x 1</td>
</tr>
<tr>
<td>• TE Buffer</td>
<td>15 ml x 1</td>
<td>40 ml x 1</td>
</tr>
<tr>
<td>• HRP-Streptavidin</td>
<td>25 µl x 1</td>
<td>25 µl x 1</td>
</tr>
<tr>
<td>• Washing Buffer (powder for 1 L)</td>
<td>x 1</td>
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<tr>
<td>• Filtration Tube</td>
<td>x 5</td>
<td>x 20</td>
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<tr>
<td>• 96-well Microplate/ U bottom</td>
<td>x 1</td>
<td>x 1</td>
</tr>
<tr>
<td>• Manual</td>
<td>x 1</td>
<td>x 1</td>
</tr>
</tbody>
</table>

* ARP-DNA Standard Solutions: 0.5 µg DNA / ml
1. Required Equipments and Materials for Abasic Sites Determination Using This Kit
- Microplate reader with 650 nm filter
- 0.5 ml and 1.5 ml tube
- 10 µl and 100-200 µl pipettes
- 50-250 µl multi-channel pipette

- Centrifuge: for ARP-labeled DNA purification using filtration tubes
- Incubator: for incubation at 37°C

2. Protocol

Purification of genomic DNA
Several different methods and products are available for the isolation of genomic DNA from samples such as membrane binding method, guanidine/detergent lysis method, and polyelectrolyte precipitation method. Among these methods, the guanidine/detergent lysis method is simple, and it gives highly purified genomic DNA for the ARP-based abasic sites detection. During the purification process, avoid heating of the DNA solution. Determine the concentration and purity of the purified genomic DNA using the spectrophotometer* and agarose gel electrophoresis. Dissolve the genomic DNA in TE at the concentration of 100 µg/ml. It is important for an accurate assay that the DNA concentration is adjusted exactly to 100 µg/ml.

For more information, please contact us at info@dojindo.com or 1-877-987-2667.

* 1 O.D. at 280 nm = 50 µg/ml. The ratio of O.D. at 260 nm/O.D. at 280 nm of highly purified DNA solution is 1.8 or higher. Protein contamination in the sample solution may cause a positive error.

ARP reaction (Preparation of ARP-labeled DNA)
1) Mix 10 µl of purified genomic DNA solution (100 µg/ml) and 10 µl of ARP Solution in a 0.5 ml tube, and incubate at 37°C for 1 hour.
2) Wash the inside of the Filtration Tube cup with 100 µl of TE twice (see Fig. 3).
3) Add 380 µl of TE to the reaction solution, and transfer the solution to the Filtration Tube (see Fig. 3).^1
4) Centrifuge the Filtration Tube at 2,500 x g for 15 min, and discard the filtrate solution.
5) Add 400 µl of TE to the Filtration Tube and resuspend the DNA on the filter with a pipette.
6) Centrifuge the Filtration Tube at 2,500 x g for 15 min. 2)
7) Add 200 µl of TE to the Filtration Tube to resuspend the DNA on the filter with a pipette.
8) Transfer the DNA solution to the 1.5 ml tube, and add 200 µl of TE again to the Filtration Tube to transfer the ARP-labeled DNA on the filter completely to the 1.5 ml tube.\(^3\)
9) Store the ARP-labeled genomic DNA solution at 0-5°C.
\(^1\) Ethanol precipitation can be used for the purification of the ARP-labeled DNA instead of purification by Filtration Tube. After ethanol precipitation, dissolve the DNA pellet in 100 µl of TE, and determine the DNA concentration.
\(^2\) If the DNA solution still remains on the filter after the centrifugation, spin for another 5 min, then proceed to step 7.
\(^3\) Recovery rate of DNA using the filtration tube is 90%, so the concentration of the ARP-labeled DNA is 2.25 µg/ml.
For more accurate determination of the number of abasic sites in the sample DNA, we recommend measuring the DNA concentration.

Determination of the number of abasic sites in DNA
Day 1:
1) Dilute 90 µl of the ARP-labeled genomic DNA with 310 µl of TE.
2) Add 60 µl of Standard ARP-DNA Solution per well. Use three wells per 1 standard solution (please see “DNA Solution Arrangement on the Plate” on page 3).
3) Add 60 µl of the diluted ARP-labeled genomic DNA solution per well. Use at least three wells per 1 sample.
4) Add 100 µl of the DNA Binding Solution to each well\(^4\) and mix by pipetting several times, then allow the plate to remain at room temperature overnight.

\(^4\) Alternatively, mix 200 µl of DNA solution and 333 µl of DNA Binding Solution in a tube, and put 160 µl of the mixture to each well (see notes 4).

Day 2:
Preparation of the solutions
Washing Buffer: Dissolve the contents of the Washing Buffer packet in 1 L of deionized or distilled water. Store this Washing Buffer solution at room temperature.

HRP-Streptavidin solution: Dilute HRP-Streptavidin with Washing Buffer to prepare 1/4,000 diluted working solution.*

1/4,000 dilution preparation: Add 10 µl of HRP-Streptavidin into 40 ml of Washing Buffer solution, and mix well.

* Since this working solution is not stable, always use freshly prepared solution.

5) Discard the DNA Binding Solution in the wells, and wash the wells with 250 µl Washing Buffer 5 times.\(^5\)
6) Add 150 µl of diluted HRP-Streptavidin solution to each well, and incubate the plate at 37°C for 1 hour.
7) Discard the solution in the well, and wash the well with 250 µl Washing Buffer 5 times.\(^5\)
8) Add 100 µl of Substrate Solution to each well, and incubate at 37°C for 1 hour.
9) Measure the O.D. at 650 nm within 1 hour after the incubation is finished, and prepare a calibration curve using the data obtained by standard ARP-DNA solutions.
10) Determine the number of abasic sites in the genomic DNA using the calibration curve.

\(^5\) After discarding the solution, invert the plate and tap it on a paper towel several times to remove the solution completely.

* Typical standard curve is shown in the Certificate of Analysis of this kit.
DNA Solution Arrangement on the Plate

Standard ARP-DNA (60 μl/well) or sample DNA (60 μl/well) + DNA binding solution (100 μl/well)

### Notes

1. Please store the kit at 0-5 °C. Do not freeze. Store Washing Buffer solution at room temperature.
2. AP-DNA is not stable. Please treat it with ARP and purify with Filtration Tube after the isolation of genomic DNA from a sample. Purified ARP-DNA solution in TE Buffer is stable over one year at 0-5 °C storage.
3. After the spinning of Filtration Tube for ARP-labeled DNA purification, add 200 µl TE immediately. If the DNA stays in Filtration Tube for more than 30 minutes after the spinning, the DNA recovery ratio may decline.
4. γ-Ray-sterilized tubes may cause DNA binding on the surface of the tube during the mixing of the DNA solution with DNA Binding Solution. If you prefer to mix ARP-DNA solution with DNA Binding Solution in a tube rather than mixing them in a well, please avoid using γ-ray-sterilized tubes.
5. For accurate determination of the number of abasic sites in sample DNA, multiple measurements are recommended.
6. If the volume of the sample DNA solution is smaller than 10 µl, use an equal volume of ARP Solution, and determine the concentration of DNA after the purification by Filtration Tubes.
7. If the 650 nm filter is not available for the measurement of O.D. after the color development, transfer 50 µl of the solution in each well to a well of a new plate (not provided). Then, add 50 µl of 1 mol/l sulfuric acid, and measure the O.D. at 450 nm.
8. Remaining solution in a well may cause error, so please remove the solution thoroughly by tapping the plate on a paper towel in each step.

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**Fig. 4** Typical calibration curve for AP-site determination.
A. No coloring or very low O.D. in all wells
1. Did you add DNA Binding Solution to the DNA solution? - Insufficient amount of DNA Binding Solution may cause no DNA binding on a plate.
2. Did you add 1/4,000 diluted HRP-Streptavidin solution?
3. Did you add Substrate Solution?
4. Is the O.D. measurement in the right wavelength?
5. Did you prepare 1/4,000 diluted HRP-Streptavidin solution just prior to use?
6. Is the HRP-Streptavidin solution properly stored at 0-5 °C? - Storage at room temperature or at freezing temperature may cause significant decrease in the enzyme activity.

B. No coloring in sample DNA
1. Did you adjust the original DNA concentration of your sample exactly to 100 µg/ml?
2. Did you add ARP solution to DNA solution?
3. Did you purify the DNA solution after the ARP reaction with a Filtration Tube or ethanol precipitation?
4. The number of AP sites in the sample DNA may be smaller than 1/100,000 bp (the detection limit).

C. High coloring in all wells
1. Did you dilute HRP-Streptavidin to 1/4,000 with Washing Buffer solution? - High concentration of HRP-Streptavidin causes high background.
2. Did you wash each well 5 times with Washing Buffer solution?

D. Calibration curve is not linear and/or high CV (over 20 %)
1. Did you take the same volume of ARP-DNA Standard Solution from each vial and mix them with DNA Binding Solution?
2. Did you wash each well 5 times with Washing Buffer solution?
3. Did you remove Washing Buffer solution thoroughly? - After discarding the solution, invert the plate and tap it on a paper towel several times to remove the solution completely.
4. Did you use the Washing Buffer in this kit and dissolved it in 1 liter of deionized or distilled water? - Lower or higher concentration of Washing Buffer causes serious error.
5. Did you leave the plate overnight at room temperature in order to allow the DNA to bind on the plate surface? - At least 4 hours of incubation time is necessary to bind DNA to the plate surface in the right consistency.
6. Did you follow the direction of the DNA purification protocol using Filtration Tube after the ARP reaction? - ARP contamination in the ARP-labeled DNA solution causes serious error.

E. Signal of the sample DNA is out of range
1. Dilute the sample DNA with 0 ARP-DNA Standard Solution, and determine the number of AP sites in the diluted sample DNA. Then, multiply the number by the dilution ratio. Zero ARP-DNA Standard Solution for a sample dilution is sold separately, please contact us for further information.
2. Did you dilute the DNA solution with TE after the purification by Filtration Tube?
3. Is your sample DNA pure enough? - Please measure the ratio of O.D.\text{260nm} / O.D.\text{280nm}.
   The ratio should be 1.8 or higher.

References
1. T. Lindahl and B. Nyberg, Biochemistry, 1972, 1f, 3610.
Day 1

Step 1) Add 10 μl of DNA solution and 10 μl of ARP solution, and incubate at 37°C for 1 hour.

a) If the volume of the DNA solution is smaller than 10 μl, add the same volume of ARP solution.

Step 2) Purify the ARP-labeled DNA with a Filtration Tube.

First, add 380 μl of TE to the ARP-labeled DNA solution (20 μl) from Step 1.

Step i) Wash the cup with 100 μl TE (x2). Put the 400 μl of ARP-labeled DNA solution into a Filtration Tube and spin at 2500 g for 15 min.

Step ii) Discard the filtrate, add 400 μl of TE and spin at 2500 g for 15 min.

Step iii) Add 200 μl of TE and rinse the filter several times to dissolve the DNA on the filter. Transfer the ARP-labeled DNA solution to a 1.5 ml tube. Repeat this step one more time.

Step 3) Take 90 μl of the ARP-labeled DNA solution, and dilute it with 310 μl of TE. Add 60 μl of the diluted solution or ARP-DNA Standard Solution to each well.

b) Follow "DNA Solution Arrangement on the plate" diagram on page 3.

Step 4) Add 100 μl of DNA Binding Solution to each well and mix by pipetting several times. Incubate the plate at room temperature overnight.
Day 2

Read Technical information carefully prior to using this General Protocol

Step 5)
Discard the solution from each well, and wash the well with 250 μl of Washing Buffer solution 5 times. Tap the plate on a paper towel several times to remove the Washing Buffer solution thoroughly from each well.

Step 6)
Add 150 μl of the diluted HRP-Streptavidin solution c) to each well, and incubate at 37°C for 1 hour.

c) Use 1/4,000 diluted HRP-Streptavidin solution. Add 10 μl HRP-Streptavidin in 40 ml of Washing Buffer solution, and mix well.

Step 7)
Discard the solution from each well, and wash the well with 250 μl of Washing Buffer solution 5 times. Tap the plate on a paper towel several times to remove the Washing Buffer solution thoroughly from each well.

Step 8)
Add 100 μl of Substrate Solution to each well, and incubate at 37°C for 1 hour.

Step 9)
Read the O.D. of each well at 650 nm d), and determine the number of abasic sites in the sample DNA using a calibration curve.

d) If 650 nm filter is not available, transfer 50 μl from each well to a well of a new plate, and add 50 μl of 1 mol/l sulfuric acid. Then, measure the O.D. at 450 nm.