for purification of genomic DNA from up to 50 mg of plant material in 96-well format using a centrifuge
Instruction for Invisorb® DNA Plant HTS 96 Kit

For 96 DNA extractions from up to 50 mg of plant material for use on a centrifuge

The Invisorb® DNA Plant HTS 96 Kit has been designed to purify extremely fast genomic DNA from up to 50 mg plant material using a centrifuge. The patented technology for isolation of genomic DNA by binding onto the filter surface does not need any hazardous chaotropic buffer components. The isolation protocol as well as all buffers is optimized to provide high yield and purity of the extracted genomic DNA. The “hands-on time” necessary for the whole procedure is reduced to a minimum.

Trademarks: Invisorb®, Registered marks, trademarks, etc. used in this document, even when not specifically marked as such, are not to be considered unprotected by law.


Invisorb® is a registered trademark of Invitek Molecular GmbH.

The PCR process is covered by US Patents 4,683,195, and 4,683,202 and foreign equivalents owned by Hoffmann-La Roche AG.

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Contents

Kit content of Invisorb® DNA Plant HTS 96 Kit/ C 3
Symbols 4
Storage 4
Quality control and product warranty 4
Intended use 5
Product use limitation 5
Safety information 6
Product characteristic of the Invisorb® DNA Plant HTS 96 Kit/ C 7
Yield and quality of genomic DNA 7
Sampling and storage of starting material 7
Important points before starting a protocol 8
Preparing reagents and buffers 8
Reagents and equipment to be supplied by user 8
Scheme of Invisorb® DNA Plant HTS 96 Kit/ C 9
Principle and procedure 10
  Protocol: DNA extraction from plant material for use on a centrifuge 10
Troubleshooting 13
Appendix 14
Ordering information 15
## Kit contents of Invisorb® DNA Plant HTS 96 Kit/ C

<table>
<thead>
<tr>
<th></th>
<th>4 x 96 DNA extractions</th>
<th>24 x 96 DNA extractions</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Catalogue No.</strong></td>
<td>7037300300</td>
<td>7037300400</td>
</tr>
<tr>
<td><strong>Lysis Buffer P</strong></td>
<td>4 x 40 ml</td>
<td>24 x 40 ml</td>
</tr>
<tr>
<td><strong>Binding Buffer A</strong></td>
<td>2 x 24 ml</td>
<td>270 ml</td>
</tr>
<tr>
<td></td>
<td>(final volume 2 x 80 ml)</td>
<td>(final volume 900 ml)</td>
</tr>
<tr>
<td><strong>Elution Buffer</strong></td>
<td>60 ml</td>
<td>350 ml</td>
</tr>
<tr>
<td><strong>Proteinase K</strong></td>
<td>4 x 2 ml</td>
<td>24 x 2 ml</td>
</tr>
<tr>
<td><strong>Wash Buffer I</strong></td>
<td>2 x 80 ml</td>
<td>2 x 350 ml</td>
</tr>
<tr>
<td></td>
<td>(final volume 2 x 160 ml)</td>
<td>(final volume 700 ml)</td>
</tr>
<tr>
<td><strong>Wash Buffer II</strong></td>
<td>4 x 45 ml</td>
<td>4 x 270 ml</td>
</tr>
<tr>
<td></td>
<td>(final volume 4 x 150 ml)</td>
<td>(final volume 900 ml)</td>
</tr>
<tr>
<td><strong>2.0 ml Collection Plate</strong></td>
<td>2 x 4</td>
<td>12 x 4</td>
</tr>
<tr>
<td><strong>Elution Plate L</strong></td>
<td>4</td>
<td>24</td>
</tr>
<tr>
<td><strong>Prefilter Plate</strong></td>
<td>4</td>
<td>6 x 4</td>
</tr>
<tr>
<td><strong>DNA Binding Plate D</strong></td>
<td>4</td>
<td>6 x 4</td>
</tr>
<tr>
<td><strong>Plate Lid</strong></td>
<td>8</td>
<td>48</td>
</tr>
<tr>
<td><strong>Sealing Foils</strong></td>
<td>4</td>
<td>24</td>
</tr>
<tr>
<td><strong>Manual</strong></td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

### Initial steps

**Add 56 ml 99.7% Isopropanol to each Binding Buffer A. Mix by intensive shaking by inverting for 1 min. Shortly before use mix by inverting several times.**

**Add 80 ml 96-100% ethanol to each bottle Wash Buffer I**

**Add 105 ml of 96-100% ethanol to each bottle Wash Buffer II, mix shortly and keep the bottle always firmly closed!**

**Add 2 ml ddH₂O to each tube Proteinase K, mix thoroughly until completely dissolving**

**Add 630 ml 99.7% Isopropanol to the Binding Buffer A. Mix by intensive shaking by inverting for 1 min. Shortly before use mix by inverting several times.**

**Add 350 ml 96-100% ethanol to each bottle Wash Buffer I**

**Add 630 ml of 96-100% ethanol to each bottle Wash Buffer II, mix shortly and keep the bottle always firmly closed!**

**Add 2 ml ddH₂O to each tube Proteinase K, mix thoroughly until completely dissolving**
Symbols

Manufacturer
Lot number
Catalogue number
Expiry date
Consult operating instructions
Temperature limitation
Do not reuse
Humidity limitation

Storage

All buffers and kit contents of the Invisorb® DNA Plant HTS 96 Kit/ C, except dissolved Proteinase K should be stored at room temperature and are stable for at least 12 months.

Room temperature (RT) is defined as range from 15 - 30°C.

Before every use, make sure that all components have room temperature. If there are any precipitates within the provided solutions solve these precipitates by warming carefully (up to 30°C).

Proteinase K: Dissolved Proteinase K must be stored at 2 - 8 °C for up to two months. For longer storage -20 °C is recommended, freeze-thaw once only. Therefore, the dissolved Proteinase K is stable as indicated on the kit package.

Wash Buffers charged with ethanol should be appropriately sealed and stored at room temperature.

Binding Buffer charged with isopropanol should be appropriately sealed and stored at room temperature.

Quality control and product warranty

Invitek Molecular warrants the correct function of the Invisorb® DNA Plant HTS 96 Kit/ C for applications as described in this manual. Purchaser must determine the suitability of the Product for its particular use. Should any Product fail to perform the applications as described in the manual, Invitek Molecular will check the lot and if Invitek Molecular investigates a problem in the lot, Invitek Molecular will replace the Product free of charge.

Invitek Molecular reserves the right to change, alter, or modify any Product to enhance its performance and design at any time.

In accordance with Invitek Molecular’s EN ISO 13485 certified Quality Management System the performance of all components of the Invisorb® DNA Plant HTS 96 Kit/ C have been tested separately against predetermined specifications routinely on lot-to-lot to ensure consistent product quality.

If you have any questions or problems regarding any aspects of Invisorb® DNA Plant HTS 96 Kit/ C or other Invitek Molecular products, please do not hesitate to contact us. A copy of Invitek Molecular’s terms and conditions can be obtained upon request or are presented at the Invitek Molecular webpage www.invitek-molecular.com.

For technical support or further information please contact:
from Germany +49-(0)30-9489-2901/ 2910
from abroad +49-(0)30-9489-2907 or contact your local distributor.
Intended use

The Invisorb® DNA Plant HTS 96 Kit/ C is the ideal tools for a rapid and efficient isolation of high quality genomic DNA from up to 50 mg of a wide variety of plant species (fresh, frozen or dried plant material, for instance leaves, roots, fruits or seeds). The protocols for the isolation and all buffers are optimized for a high yield as well as a high purity. All hands on steps are reduced to a minimum.

For reproducible and high yields, appropriate sample storage is essential.

THE PRODUCT IS INTENDED FOR USE BY PROFESSIONALS ONLY, SUCH AS TECHNICIANS, PHYSICIANS AND BIOLOGISTS TRAINED IN MOLECULAR BIOLOGICAL TECHNIQUES. It is designed to be used with any downstream application employing enzymatic amplification or other enzymatic modification of DNA followed by signal detection or amplification. Any results generated using the sample preparation procedure in conjunction with any downstream assay should be interpreted with regard to other laboratory findings.

To minimize irregularities in your results, adequate controls for downstream applications should be used.

Product use limitation

The kit is neither suitable for isolation of RNA from plant material, as for DNA isolation from cultured or isolated cells, tissue samples or blood samples. The isolation of simultaneously isolation of bacterial DNA or DNA from fungi and parasites is not validated.

The included chemicals are only useable once.

Differing of starting material or flow trace may lead to inoperability; therefore, neither a warranty nor guarantee in this case will be given, neither implied nor express.

The user is responsible to validate the performance of the Invitek Molecular Product for any particular use. Invitek Molecular does not provide for validation of performance characteristics of the Product with respect to specific applications.

Invitek Molecular products may be used e.g. in clinical diagnostic laboratory systems under following conditions:

- If used in the US, based on the condition that the complete diagnostic system of the laboratory has been validated pursuant to CLIA’ 88 regulations.
- For other countries based on the condition that the laboratory has been validated pursuant to equivalents according to the respective legal basis.

All Products sold by Invitek Molecular are subject to extensive quality control procedures (according to EN ISO 13485) and are warranted to perform as described herein. Any problems, incidents or defects shall be reported to Invitek Molecular immediately upon detection thereof.

The chemicals and the plastic parts are for laboratory use only; they must be stored in the laboratory and must not be used for purposes other than intended.

The Product with its contents is unfit for consumption.
Safety Information

When and while working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles!

Avoid skin contact! Adhere to the legal requirements for working with biological material!

For more information, please consult the appropriate material safety data sheets (MSDS). These are available online in convenient and compact PDF format at www.invitek-molecular.com for each Invitek Molecular Product and its components. If buffer bottles are damaged or leaking, WEAR GLOVES, AND PROTECTIVE GOGGLES when discarding the bottles in order to avoid any injuries.

Invitek Molecular has not tested the liquid waste generated by the Invisorb® DNA Plant HTS 96 Kit procedures for residual infectious materials. Contamination of the liquid waste with residual infectious materials is highly unlikely, but cannot be excluded completely. Therefore, liquid waste has be considered infectious and be handled and discarded according to local safety regulations.

European Community risk and safety phrases for the components of the Invisorb® DNA Plant HTS 96 Kit to which they apply, are listed below as follows:

**Lysis Buffer P**  
![Warning]  
H319-H412-P280-P305+P351+P338-P273

**Proteinase K**  
![Danger]  
H315-319-334-335-P280-P305-P351-P338

**Wash Buffer I**  
![Warning]  
H302-H412-P280-P305-P351-P338-P273-EUH032

H302: Harmful if swallowed.
H315: Causes skin irritation.H317: May cause an allergic skin reaction.
H319: Causes serious eye irritation.
H334: May cause allergy or asthma symptoms or breathing difficulties if inhaled.
H335: May cause respiratory irritation.
H412: Harmful to aquatic life with long lasting effects.
P273: Avoid release to the environment.
P280: Wear protective gloves/protective clothing/eye protection/face protection.
P305+P351+P338: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
EUH032: Contact with acids liberates very toxic gas.

Emergency medical information can be obtained 24 hours a day from infotrac:

outside of USA: 1 – 352 – 323 – 3500
in USA: 1 – 800 – 535 – 5053
Product Characteristic of the Invisorb® DNA Plant HTS 96 Kit

<table>
<thead>
<tr>
<th>Starting material</th>
<th>Yield</th>
<th>Time</th>
<th>Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>up to 50 mg of a wide</td>
<td>up to 6 µg/ well</td>
<td>about 100 min (C) inclusive 60 min incubation</td>
<td>A$<em>{260}$/A$</em>{280}$ 1.6-2.0</td>
</tr>
<tr>
<td>variety of plant species</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The Invisorb® DNA Plant HTS 96 Kit has been designed for a semi-automated preparation of genomic DNA for up to 50 mg of plant sample in 96 well format.

Due to the high purity, the isolated genomic DNA is ready to use for a broad panel of downstream applications:

- PCR*-Reactions
- RFLP-Analysis
- Restriction Enzyme Digestion
- Hybridization

For the isolation of DNA from single plant sample Invitek Molecular offers the Invisorb® Spin Plant Mini Kit.

For further information please contact: Phone +49 (0) 30 9489 2901, 2910 in Germany and from foreign countries phone +49 (0) 30 9489 2907 or your local distributor.

Yield and quality of genomic DNA

The amount of purified DNA in the Invisorb® DNA Plant HTS 96 Kit procedure from plant material depends on sample source, transport conditions, storage and age of the sample. Yield and quality of isolated genomic DNA is suitable for any detection system.

Important indications for the Invisorb® DNA Plant HTS 96 Kit

The kit can also purify RNA besides DNA. For the elimination of RNA (if necessary) add RNase A according to the manufacturer’s instructions. You can find an example for preparing RNase A solution on page 11.

Sampling and Storage of starting material

Harvested plant samples can be stored at room temperature for 2 – 3 hours, for short time storage (up to one week) samples may be stored at 2 - 8°C. For long-term storage, we recommend freezing samples at –20°C or –80°C. Multiple thawing and freezing before isolating the DNA should be avoided.

*) The PCR method is covered by U.S. Patents 4,683,195 and 4,683,202 owned by Hoffmann-LaRoche Inc. The purchase of the Invisorb® DNA Plant HTS 96 Kit cannot be construed as an authorization or implicit licence to practice PCR under any patents held by Hoffmann-LaRoche Inc.
Important points before starting a protocol

Immediately upon receipt of the Product, inspect the Product and its components as well as the package for any apparent damages, correct quantities and quality. If there are any unconformities you have to notify Invitek Molecular in writing with immediate effect upon inspection thereof. If buffer bottles are damaged, contact the Invitek Molecular Technical Services or your local distributor. In case of liquid spillage, refer to “Safety Information” (see page 6). Do not use damaged kit components, since their use may lead to poor kit performance.

- Always change pipet tips between liquid transfers. To avoid cross-contamination, we recommend the use of aerosol-barrier pipet tips.
- All centrifugation steps are carried out at room temperature.
- When working with chemicals, always wear a suitable lab coat, disposable gloves and protective goggles.
- Discard gloves if they become contaminated.
- Do not combine components of different kits, unless the lot numbers are identical.
- Avoid microbial contamination of the kit reagents.
- This kit should only be used by trained personnel.

Preparing reagents and buffers

4 x 96 DNA-extractions:
Add 56 ml 99.7% Isopropanol to each Binding Buffer A. Mix by intensive shaking by inverting for 1 min. Shortly before use mix by inverting several times.
Add 80 ml 96-100% ethanol to each bottle Wash Buffer I
Add 105 ml of 96-100% ethanol to each bottle Wash Buffer II, mix shortly and keep the bottle always firmly closed!
Add 2 ml ddH2O to each tube Proteinase K, mix thoroughly until completely dissolving!

24 x 96 DNA-extractions:
Add 630 ml 99.7% Isopropanol to the Binding Buffer A. Mix by intensive shaking by inverting for 1 min. Shortly before use mix by inverting several times.
Add 350 ml 96-100% ethanol to each bottle Wash Buffer I
Add 630 ml of 96-100% ethanol to each bottle Wash Buffer II, mix shortly and keep the bottle always firmly closed!
Add 2 ml ddH2O to each tube Proteinase K, mix thoroughly until completely dissolving!

Reagents and equipment to be supplied by user

- Multichannel pipet with tips
- Reagents reservoirs for multichannel pipets
- Centrifuge: output ≥ 2,000 x g is necessary, for example Eppendorf Centrifuge 5804 / 5804 R / 5810 / 5810 R with Deepwell-Plate-Rotor (A-2-DWP)
- Ethanol (96-100%)
- Mixer Mill 300 and equipment from Qiagen
- Optionally RNAse A (Applichem, Product A3832,0xxx available as 50 mg, 250 mg and 500 mg, per kit with 4 x 96: 1.6 mg are required, for a kit with 24 x 96 preps: 9.5 mg)
- Isopropanol*

*The Invisorb® DNA Plant HTS 96 Kit/ C is validated with 2-Propanol; Rotipuran >99.7%, p.a., ACS, ISO (Order no. 6752) from Carl Roth.

* Possible suppliers for Isopropanol:

<table>
<thead>
<tr>
<th>Carl Roth</th>
<th>Applichem</th>
<th>Sigma</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-Propanol Rotipuran &gt;99.7%, p.a., ACS, ISO Order no. 6752</td>
<td>2-Propanol für die Molekularbiologie Order no. A3928</td>
<td>2-Propanol Order no. 59304-1L-F</td>
</tr>
</tbody>
</table>
Please read protocols prior to the start of the preparation

- 50 mg plant material into the QIAwell plate, homogenize with Mixer Mill MM 300 (Retsch) under liquid N
- Add 400 µl Lysis Buffer P Mix (inclusive Proteinase K and RNase A*) to each well of the Qiawell plate
- Incubate at 65 °C for 60 min.
- Centrifuge for 10 min at 1,700 x g (4,000 rpm)

*If you want to get your sample preparation free of RNA, use an amount of RNase A according to the manufacturer’s instructions.

- Transfer 350 µl of the lysate carefully from the QIAwell plate into the wells of the Prefilter Plate
- Close the Prefilter Plate with a Plate Lid
- Load the whole block (Prefilter Plate/ 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket
- Centrifuge at 1,700 x g (4,000 rpm) for 5 min
- Discard the Prefilter Plate

- Add 350 µl Binding Buffer A (follow preparing instructions) to each well of the 2 ml Collection Plate
- Place the DNA Binding Plate D on top of a new 2 ml Collection Plate.
- Transfer the suspension completely into each well of the DNA Binding Plate D.
- Close the DNA Binding Plate D with the Plate Lid and incubate for 1 min at RT
- Centrifuge at 1,700 x g (4,000 rpm) for 10 min at RT
- Remove the Plate Lid and discard the filtrate.

- Add 500 µl Wash Buffer I to each well of DNA Binding Plate D
- Centrifuge at 1,400 - 1,700 x g (3,700 – 4,000 rpm) for 5 min at RT
- Remove the Plate Lid and discard the filtrate

- Add 700 µl Wash Buffer II to each well of DNA Binding Plate D
- Centrifuge at 1,700 x g (4,000 rpm) for 5 min at RT
- Discard filtrate and place the DNA Binding Plate D back on the top of a 2 ml Collection Plate
- Repeat the wash step with Wash Buffer II

- Remove all waste from the waste tray
- Centrifuge at maximum speed for 15 min to dry the membrane
- Discard 2 ml Collection Plate

- Place the DNA Binding Plate D on the top of a Elution Plate L or of a Microtube Plate
- Add 60 µl Elution Buffer
- Incubate 5 min at RT.
- Centrifuge at 1,700 x g (4,000 rpm) for 5 min
- Repeat the elution step
- Take the DNA Binding Plate D and the Elution Plate L out of the centrifuge
- Discard the DNA Binding Plate D
Principle and procedure

The Invisorb® DNA Plant HTS 96 Kit procedure comprises following steps:

- lysis of sample material
- binding the genomic DNA to the membrane
- washing and elimination of ethanol
- elution of genomic DNA

After lysis the DNA binds to the membrane, contaminations and enzyme inhibitors are efficiently removed during the following three washing steps and highly purified DNA is eluted in Elution Buffer or water.

Protocol: DNA extraction from plant material for use on a centrifuge

Attention:

The complete disruption and homogenization of the plant samples is absolute essential for isolation of high yield of genomic DNA. The disruption procedure, the breakage of intercellular matrix and plasma membranes, is necessary to release the nucleic acids contained in the plant cell, thus inefficient disruption decreases the DNA yield.

The homogenization means the reduction of the viscosity of the lysate after disruption. Contaminating total RNA and other cellular components of high molecular weight are sheared to form a homogenous lysate. If the homogenization of the starting material is not done very carefully the yield of genomic DNA purified is reduced significantly.

It is possible to use a commercially available bead mill or rotor-stator homogenizer in combination with or without beads for the disruption and homogenization of the starting material. Alternatively, the starting material can reduced to a fine powder in liquid nitrogen using a mortar and pestle.

Rotor–stator homogenizers and bead mills simultaneously disrupt and homogenize the plant material, whereas plant tissue are only disrupted using a mortar and pestle, and a separate homogenization step should be performed.

Note: Elution with pre-warmed Elution Buffer (up to 80 °C) will also increase the final yield.

1. Disruption, Homogenization and Lysis of the plant material in a Mixer Mill*)

Note: It is recommended to use a mixer mill for the simultaneous homogenization of 96 plant samples. The Mixer Mill from Retsch works in a 96-well format, but only in combination with a special adapter from Qiagen. Therefore, the use of QIAwell plates (not provided) is necessary.

Transfer up to 50 mg plant material into the QIAwell plate (not provided) and homogenize with Mixer Mill MM 300 (Retsch) under liquid N2.

Mix one bottle of Lysis Buffer P with one tube of Proteinase K and add RNase A** according to the manufacturer’s instructions (optional, if you want to get your sample preparation free of RNA).

Add 400 µl Lysis Buffer P Mix (inclusive Proteinase K and optionally RNase A) to each well of the Qiawell plate.

Incubate at 65 °C for 60 min, centrifuge for 10 min at 1.700 x g (4.000 rpm).

*) If a Mixer Mill is not available, a disruption and homogenization of the plant material must be performed with other methods, like with mortar, pestle and N2 or within another kind of mill (like Savant) with beads, or on a Gyrator (UniEquip) or using a rotor–stator homogenizer.
**Example for RNase from Applichem (Ordering number A3832,0xxx)**

Make a stock solution of 1 mg / ml with TE – Buffer. Add 1 / 100-part V/V to the Lysis Buffer means 4 µl / Reaction; 10 µl / 1 ml; or 400 µl / 40 ml (1 plate). The RNase stock solution is stable at 4°C for about 8 weeks.

2. Prefiltration

Place the Prefilter Plate on top of a 2 ml Collection Plate. Transfer 350 µl of the lysate carefully from the QIAwell plate into the wells of the Prefilter Plate. Close the Prefilter Plate with a Plate Lid.

Load the whole block (Prefilter Plate/ 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket. Centrifuge at 1.700 x g (4.000 rpm) at RT.

Take the Prefilter Plate/ 2 ml Collection Plate out of the centrifuge. Remove the Plate Lid.

Discard the Prefilter Plate.

**Note:** You do not need to centrifuge, if the lysates have already run through the filter

3. Binding of the genomic DNA to the DNA Binding Plate D

Add 350 µl Binding Buffer A to each well of the 2 ml Collection Plate and mix the solution very well by pipetting several times up and down by using a multichannel pipet.

Place the DNA Binding Plate D on top of a new 2 ml Collection Plate. Transfer the suspension completely into each well of the DNA Binding Plate D. Close the DNA Binding Plate D with the Plate Lid and incubate for 1 min at RT.

Load the whole block (DNA Binding Plate D/ 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket. Centrifuge at 1.700 x g (4.000 rpm) for 10 min at RT.

Take the DNA Binding Plate D/ 2 ml Collection Plate out of the centrifuge. Remove the Plate Lid and discard the filtrate.

Place the DNA Binding Plate D back to the top of the 2 ml Collection Plate.

4. First Washing of the DNA Binding Plate D

Add 500 µl Wash Buffer I to each well of the DNA Binding Plate D. Close the DNA Binding Plate D with the Plate Lid. Load the whole block (DNA Binding Plate D/ 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket. Centrifuge at 1.700 x g (4.000 rpm) for 5 min at RT.

Remove the Plate Lid and discard the filtrate.

Place the DNA Binding Plate D back to the top of the 2 ml Collection Plate.

5. Second washing of the DNA Binding Plate D

Add 700 µl Wash Buffer II to each well of the DNA Binding Plate D. Close the DNA Binding Plate D with the Plate Lid. Load the whole block (DNA Binding Plate D / 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket. Centrifuge at 1.700 x g (4.000 rpm) for 5 min at RT.

Remove the Plate Lid and discard the filtrate.

Place the DNA Binding Plate D back to the top of the 2 ml Collection Plate.

Repeat this washing step (Wash Buffer II)
6. Removing of ethanol

Take the DNA Binding Plate D/ 2 ml Collection Plate out of the centrifuge. Remove the Plate Lid. Place the DNA Binding Plate D onto a clean surface (paper towel). Empty the 2 ml Collection Plate and dry its upper side with paper.

Place the DNA Binding Plate D on top of the 2 ml Collection Plate and close it with a Plate Lid. Load the whole block (DNA Binding Plate D/ 2 ml Collection Plate) into the holder and place the whole assembly in the rotor bucket of the centrifuge. Centrifuge at maximum speed for at least 15 min at RT to eliminate any traces of ethanol.

Take the DNA Binding Plate D/ 2 ml Collection Plate out of the centrifuge, discard the Plate Lid and place the plate on a clean paper towel.
Discard the 2 ml Collection Plate.

7. Elution of the genomic DNA

Place the DNA Binding Plate D on the top of a Elution Plate L.
Add 60 µl Elution Buffer directly onto the membrane in each well and incubate for 5 min.
Close the DNA Binding Plate D with a new Plate Lid and place the whole block (DNA Binding Plate D/ Elution Plate L) in the rotor bucket of the centrifuge.
Centrifuge for 5 min at 1.700 x g (4.000 rpm)
Remove the Plate Lid.

Add 60 µl Elution Buffer again directly onto the membrane in each well. Close the DNA Binding Plate D with the Plate Lid and place the whole block (DNA Binding Plate D/ Elution Plate L) in the rotor bucket of the centrifuge. Centrifuge for 5 min at 1.700 x g (4.000 rpm).

Take the DNA Binding Plate D and the Elution Plate L very carefully out of the centrifuge in order to avoid cross-contaminations with adherent fluid.
Discard the DNA Binding Plate D.

Take the Elution Plate L with the eluted DNA and cover the plate with sealing foil.
## Troubleshooting

<table>
<thead>
<tr>
<th>Problem/ probable cause</th>
<th>Comments and suggestions</th>
</tr>
</thead>
<tbody>
<tr>
<td>clogged DNA Binding Plate D</td>
<td>increase incubation time with Lysis Buffer P increase amount of starting material</td>
</tr>
<tr>
<td>incomplete lysis of starting material/viscous lysates</td>
<td>increase incubation time with Lysis Buffer P reduce amount of starting material</td>
</tr>
<tr>
<td>low amounts or none DNA</td>
<td>increase time for lysis reduce amount of starting material or use higher volume of Lysis Buffer P Increase Binding Buffer A to adequate volume. Lysis Buffer P/ Binding Buffer A ratio has to be always 2/1 make sure that the correct amount of ethanol is added to the Wash Buffers make sure that the Buffers are stored correctly Increase incubation time with prewarmed Elution Buffer to 10 min or repeat elution step once again prewarm Elution Buffer up to 80°C</td>
</tr>
<tr>
<td>insufficient lysis</td>
<td>make sure that the correct amount of ethanol is added to the Wash Buffers check up Wash Buffers for salt precipitates. If there are any precipitates solve these precipitates by careful warming</td>
</tr>
<tr>
<td>incomplete elution</td>
<td>make sure that the correct amount of ethanol is added to the Wash Buffers make sure that the Buffers are stored correctly Increase incubation time with prewarmed Elution Buffer to 10 min or repeat elution step once again prewarm Elution Buffer up to 80°C</td>
</tr>
<tr>
<td>problems with downstream-applications (e.g. PCR)</td>
<td>increase centrifugation time for removing of ethanol ensure that Wash Buffer I and Wash Buffer II are at room temperature. Check up Wash Buffers for salt precipitates. If there are any precipitates solve these precipitates by careful warming</td>
</tr>
<tr>
<td>ethanol carryover during elution</td>
<td>increase centrifugation time for removing of ethanol ensure that Wash Buffer I and Wash Buffer II are at room temperature. Check up Wash Buffers for salt precipitates. If there are any precipitates solve these precipitates by careful warming</td>
</tr>
<tr>
<td>salt carryover during elution</td>
<td>increase centrifugation time for removing of ethanol ensure that Wash Buffer I and Wash Buffer II are at room temperature. Check up Wash Buffers for salt precipitates. If there are any precipitates solve these precipitates by careful warming</td>
</tr>
</tbody>
</table>
Appendix

General notes on handling DNA

Nature of DNA
The length and delicate physical nature of DNA requires careful handling to avoid damage due to shearing and enzymatic degradation. Other conditions that affect the integrity and stability of DNA include acidic and alkaline environments, high temperature, and UV irradiation. Careful isolation and handling of high molecular weight DNA is necessary to ensure compatibility with various downstream applications. Damaged DNA could perform poorly in applications such as genomic Southern blotting, long-template PCR.

Storage of DNA
A working stock of DNA can be stored at 2–4 °C for several weeks. For long-term storage, DNA should be stored at –20 °C, but storing at –20 °C can cause shearing, particularly if the DNA is exposed to repeated freeze-thaw cycles. Note that the solution in which the nucleic acid is eluted in will affect its stability during storage. Pure water lacks buffering capacity and an acidic pH may lead to acid hydrolysis. Tris or Tris-EDTA buffer contains sufficient buffering capacity to prevent acid hydrolysis.

Drying, dissolving and pipetting DNA
Avoid overdrying genomic DNA after ethanol precipitation. Avoid vigorous pipeting. Pipeting genomic DNA through small tip openings causes shearing or nicking. One way to decrease shearing of genomic DNA is to use special tips that have wide openings designed for pipeting genomic DNA.

DNA Yield
The amount of purified DNA from the plant material depends on sample source, transport conditions, storage and age of the sample.
Ordering information

Invisorb® DNA Plant HTS 96 Kit/ C  7037300300  4 x 96 preps
Invisorb® DNA Plant HTS 96 Kit/ C  7037300400  24 x 96 preps

Related products

InviMag® Plant DNA Mini Kit/ KF96 w/o plastic  7437300250  5 x 96 preps
Invisorb® Spin Plant Mini Kit  1037100300  250 purifications

Possible suppliers for Isopropanol:

Carl Roth
2-Propanol
Rotipuran >99.7%, p.a., ACS, ISO
Order no. 6752

Applichem
2-Propanol für die
Molekularbiologie
Order no. A3928

Sigma
2-Propanol
Order no.
59304-1L-F