

QIAsymphony® DSP DNA Midi Kit Instructions for Use (Protocol Sheet)

DNA_Blood_1000_V7_DSP protocol

Version 2

IVD

For In Vitro Diagnostic Use

For use with QIAsymphony DSP DNA Midi Kit (96)

CE

REF

937255



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R1

The protocol sheet available electronically and can be found under the resource tab of the product page on www.qiagen.com.

General information

The QIAAsymphony DSP DNA Kit is intended for in vitro diagnostic use.

This protocol is for purification of total genomic and mitochondrial DNA from fresh or frozen human whole blood using the QIAAsymphony SP and the QIAAsymphony DSP DNA Midi Kit.

Kit	QIAAsymphony DSP DNA Midi Kit (cat. no. 937255)
Sample material	Human whole blood (EDTA, citrate, or heparin anti-coagulated)
Protocol name	Blood_1000_V7_DSP
Default Assay Control Set	ACS_Blood_1000_V7_DSP
Editable	Elution volume: 200, 400, and 500 µl
Required software version	Version 4.0 or higher
Required software configuration for IVD use	Default Profile 1

“Sample” drawer

Sample type	Human whole blood (EDTA, citrate, or heparin anti-coagulated)
Sample volume	Depends on type of sample tube used; For more information, see the labware list which can be found under the resource tab of the product page on www.qiagen.com .
Primary sample tubes	For more information, see the labware list which can be found under the resource tab of the product page on www.qiagen.com .
Secondary sample tubes	For more information, see the labware list which can be found under the resource tab of the product page on www.qiagen.com .
Inserts	Depends on type of sample tube used; For more information, see the labware list which can be found under the resource tab of the product page on www.qiagen.com .

“Reagents and Consumables” drawer

Position A1 and/or A2	Reagent cartridge (RC)
Position B1	n/a
Tip rack holder 1–17	Disposable filter-tips, 200 or 1500 µl
Unit box holder 1–4	Unit boxes containing sample prep cartridges or 8-Rod Covers

n/a = not applicable.

“Waste” drawer

Unit box holder 1–4	Empty unit boxes
Waste bag holder	Waste bag
Liquid waste bottle holder	Empty liquid waste bottle

“Eluate” drawer

Elution rack (we recommend using slot 1, cooling position)

For more information, see the labware list that can be found under the resource tab of the product page on www.qiagen.com.

Required plasticware

Plasticware	One batch 24 samples*	Two batches 48 samples*	Three batches 72 samples*	Four batches 96 samples*
Disposable filter-tips, 200 µl†	4	4	8	8
Disposable filter-tips, 1500 µl†	114	220	334	440
Sample prep cartridges§	18	36	54	72
8-Rod Covers¶	3	6	9	12

* Use of less than 24 samples per batch decreases the number of disposable filter-tips required per run.

† There are 32 filter-tips/tip rack.

‡ Number of required filter-tips includes filter-tips for 1 inventory scan per RC.

§ There are 28 sample prep cartridges/unit box.

¶ There are twelve 8-Rod Covers/unit box.

Note: Numbers of filter-tips given may differ from the numbers displayed in the touchscreen depending on settings. We recommend loading the maximum possible number of tips.

Elution volume

The elution volume is selected in the touchscreen. Depending on the sample type and DNA content, the final eluate volume may vary by up to 15 µl less than the selected volume. Because the eluate volume may vary, we recommend checking the actual eluate volume when using an automated assay setup system that does not verify the eluate volume prior to transfer. Elution in lower volumes increases the final DNA concentration but slightly reduces the yield. We recommend using an elution volume appropriate for the intended downstream application.

Preparation of sample material

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, consult the appropriate safety data sheets (SDSs), available from the product supplier.

For general collection, transport and storage recommendations refer to the approved CLSI guideline MM13-A “Collection, Transport, Preparation, and Storage of Specimens for Molecular Methods”. Furthermore, the manufacturer’s instructions for the selected sample collection device shall be followed during sample preparation, storage, transport and general handling.

Independent of the blood collection tube manufacturer’s instructions ISO 20186-2:2019 (E) for automated gDNA extraction from venous whole blood should be considered.

Human whole blood

Whole blood samples treated with EDTA, citrate, or heparin can be used, and may be either fresh or frozen. If using fresh blood samples in primary tubes, mix the blood samples thoroughly (e.g., by inverting the tubes several times) before loading them onto the QIA Symphony SP. Frozen samples should be thawed quickly in a 37°C water bath with mild agitation to ensure thorough mixing and then equilibrated to room temperature (15–25°C) before beginning the procedure. To ensure reliable sample transfer, avoid generating foam in sample tubes. Try to avoid blood clots in the samples and, if necessary, transfer the sample without clots to a fresh tube.

Yield and quality of the purified DNA depend on the storage conditions of the blood. Fresher blood samples may yield better results. For short-term storage of up to 10 days, we recommend storage at 2–8°C. However, for applications requiring maximum fragment size, such as southern blotting, we recommend storage at 2–8°C for up to 3 days only, as low levels of DNA degradation will occur after this time. For long-term storage (over 10 days), collect blood in tubes containing a standard anticoagulant (preferably EDTA, if high-molecular-weight DNA is required), and store at –20°C or –80°C.

Note: Sample stability highly depends on various factors and relates to the specific downstream application. It has been established for the QIA Symphony DSP DNA Midi Kit in conjunction with exemplary downstream applications. It is the responsibility of the user to consult the instructions for use of the specific downstream application used in their laboratory and/or validate the whole workflow to establish appropriate storage conditions.

Storage of eluates

It is recommended to remove the eluate plate from the “Eluate” drawer immediately after the run has finished. Elution plates may be left in the QIA Symphony SP after the run is completed overnight (maximum 12 hours including run time; recommended environmental conditions: 18–26°C and 20–75% relative humidity). Depending on temperature and humidity, eluate may experience condensation or evaporation.

For short-term storage eluates might be stored at room temperature for up to 2 weeks. For long-term storage, we recommend storage at 2–8°C, –20°C, or –80°C. Frozen eluates must not be thawed more than three times.

Note: Eluate stability highly depends on various factors and relates to the specific downstream application. It has been established for the QIA Symphony DSP DNA Midi Kit in conjunction with exemplary downstream applications. It is the responsibility of the user to consult the instructions for use of the specific downstream application used in their laboratory and/or validate the whole workflow to establish appropriate storage conditions.

Important point before starting

- QIA Symphony magnetic particles may copurify RNA if it is present in the sample. To minimize RNA content in the sample, add RNase A to the sample before starting the procedure. The final RNase A concentration should be 2 mg/ml.

Limitations and interfering substances





Blood samples with high concentrations of triglycerides (>30 g/l) may lead to reduced gDNA yield.

Note: Testing was done using exemplary downstream applications for an assessment of the quality of the extracted nucleic acids. However, different downstream applications may have different requirements with respect to purity (i.e., absence of potential interfering substances), so the identification and testing of relevant substances also needs to be established as part of the downstream application development for any workflow involving the QIASymphony DSP DNA Midi Kit.

Note: Please note that during development of the QIASymphony DSP DNA Midi Kit no indications were observed that heparin has a negative impact on the performance. However, ISO 20186-2:2019(E) states that heparin from blood collection tubes may impact the purity of the isolated nucleic acids and possible carryover into eluates could cause inhibitions in some downstream applications. Therefore, it is the user's responsibility to validate if heparin has a negative influence on their workflow.

Symbols

The following symbols appear in this document. For a full list of symbols used in the instructions for use or on the packaging and labeling, please refer to the handbook.

Symbol	Symbol definition
	This product fulfills the requirements of the European Regulation 2017/746 for in vitro diagnostic medical devices.
	In vitro diagnostic medical device
	Catalog number
Rn	R is for revision of the Instructions for Use and n is the revision number
	Manufacturer

Revision history

Revision	Description
R1, June 2022	Version 2, Revision 1 <ul style="list-style-type: none"><li>• Update to version 2 for compliance to IVD<li>• Addition of Limitations and interfering substances section<li>• Addition of Storage of eluates section<li>• Addition of Symbols section<li>• Update of Preparation of sample material section

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