

## INSTRUCTION FOR USE

# chemagic™ BBS DNA Kit H24

**Product number:** IVD-1074

Reagents for 240 extractions.

**UDI-DI:** 4260543364175

**Version:** V240503 EN

**Manufacturer:** Revvity chemagen Technologie GmbH  
Arnold-Sommerfeld-Ring 2  
52499 Baesweiler, Germany  
[www.revvity.com](http://www.revvity.com)

CE

FOR *IN VITRO* DIAGNOSTIC USE.

## 1. TABLE OF CONTENTS










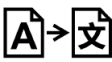



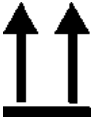
1. Table of contents .....	1
2. Explanation of the signal words in this IFU .....	3
3. Symbols used in the IFU and on labels .....	3
4. Intended purpose .....	5
5. Summary and principle .....	5
6. Reporting of incidents .....	6
7. General and storage information .....	7
8. Electronic instructions for use .....	8
9. Warnings and precaution .....	8
10. Kit reagents and safety information .....	10
10.1 Magnetic Beads .....	10
10.2 Lysis Buffer 1 .....	10
10.3 Binding Buffer 2 .....	11
10.4 Wash Buffer 3 .....	12
10.5 Wash Buffer 4 .....	13
10.6 Wash Buffer 5 .....	14
10.7 Wash Buffer 6 .....	14
10.8 Elution Buffer 7 .....	15
10.9 Proteinase K .....	15
10.10 Further kit components .....	16
11. Required Protocols Files .....	17
12. Material required but not supplied with the kit .....	17
12.1 Items from Revvity chemagen Technologie GmbH .....	17
12.2 Additional required items .....	18
12.3 Additional optional items from Revvity chemagen Technologie GmbH .....	18
12.4 Other additional optional items .....	18
13. Specimen collection and handling .....	19
14. Preparation of buffy coat samples .....	20
15. Detailed protocol description .....	21
15.1 Protocol procedure .....	21
15.2 Processing steps .....	22
15.3 Short description/ Quick guide .....	25
16. Performance characteristics .....	28
16.1 Linearity and recovery with spiked DNA sample .....	28
16.2 DNA yields with blood and buffy coat samples .....	29
17. Cleaning and maintenance .....	30






18. Downstream applications.....	31
19. Further questions.....	32
20. Limitations of the procedure .....	32
21. Influence of interfering substances .....	32
22. Warranty .....	33

## 2. EXPLANATION OF THE SIGNAL WORDS IN THIS IFU

Signal word	Description
<b>CAUTION!</b>	Potential hazard that could lead to slight or medium harm.
<b>ATTENTION!</b>	Improper handling can damage the instrument.
<b>NOTE:</b>	Errors committed by the operator can cause that the optimal performance of the kit cannot be guaranteed.

## 3. SYMBOLS USED IN THE IFU AND ON LABELS

Symbol	Symbol Title	Symbol	Symbol Title
	CE mark European conformity		Temperature limit
	<i>In vitro</i> medical device		Contains sufficient for <n> tests
	Consult instructions for use or electronic instructions for use		Quantity
	Manufacturer		Do not re-use
	Batch code		Translation
	Catalogue number		Use-by date
	Do not use if package is damaged and consult instructions for use		This way up

Symbol	Symbol Title	Symbol	Symbol Title
	GHS02		Dangerous goods: Class 3 Flammable liquid
	GHS07		Dangerous goods: Class 8 Corrosive substances
	GHS08	-	-

chemagic™ is a trademark of Revvity chemagen Technologie GmbH.

## 4. INTENDED PURPOSE

The chemagic™ BBS DNA Kit H24 (IVD-1074) is a kit for the automated isolation and purification of DNA from human blood, buffy coats (blood with reduced amount of plasma) and stabilized saliva for *in vitro* diagnostic purposes.

Other sample materials such as swabs or tissue lysates may be compatible but have not yet been validated. For such materials, a validation must be performed by the user.

The product is used on the chemagic™ 360-D instrument and is intended for laboratory personnel trained for the chemagic 360-D instrument in combination with chemagic nucleic acid purification kits. The kit is designed to be used with IVD downstream applications employing enzymatic amplification and detection of DNA (e.g. PCR, RT-PCR, NGS).

For further information please refer to the sections “KIT REAGENTS AND SAFETY INFORMATION“ and “WARNINGS AND PRECAUTION” in this document.

## 5. SUMMARY AND PRINCIPLE

The chemagic BBS DNA Kit H24 is based on a magnetic bead technology platform proprietary to Revvity chemagen Technologie GmbH. White blood cells or other source of DNA present in blood, buffy coat and saliva are lysed during the extraction process. The released nucleic acids bind to small magnetizable particles which are then magnetically separated from the sample material. During subsequent steps contaminants are removed and the purified nucleic acids are transferred into an elution buffer. The automated sample processing is performed using the chemagic 360-D instrument with a chemagic 24 Rod Head Set or equivalent instrument.

To minimize irregularities in diagnostic results, the product is intended to be used with appropriate controls throughout the process of sample preparation, sample amplification and detection according to the downstream assay used.

## 6. REPORTING OF INCIDENTS

For a user/ third party in the European Union and in countries with an identical regulatory regime (IVDR (EU) 2017/746); if, during the use of this device or as a result of its use, a serious incident has occurred, please report it to your national authority and to the manufacturer Revvity chemagen Technologie GmbH, +49 (0) 2401805500 or [support.chemagen@revvity.com](mailto:support.chemagen@revvity.com) or it's legal representatives.

The competent authority in Germany is the Federal Institute for Drugs and Medical Devices (Bundesinstitut für Arzneimittel und Medizinprodukte, BfArM). Current contact information can be found on the BfArM website: <https://www.bfarm.de>.

## 7. GENERAL AND STORAGE INFORMATION

The kit contains reagents sufficient to perform 240 extractions.

The expiry date of the unopened kit is stated on the outer label. Do not use any component beyond the expiry date. Store at +2 to +25 °C.

Once opened, the kit components have a limited stability. The stability after opening is stated for each component separately in the reagent listing below (section “KIT REAGENTS AND SAFETY INFORMATION”).

**NOTE: Recap the bottles tightly immediately after use to prevent evaporation.**

The bottles may discolor during storage. The discoloration of the bottles has no effect on the functionality of the assay.

In some cases, traces of Magnetic Beads may be left in the eluate. Though such particles will usually not interfere with PCR or most downstream applications, an additional separation step either using centrifugation or a magnetic separator (chemagic Stand 12, provided with the chemagic 360 24 Rod Head Set) is recommended, in order to separate any traces of particles.

Extracted DNA should be used immediately after extraction in the desired *in vitro* diagnostic test.

In this IFU we refer to the chemagic 360-D User Manual. This manual will be provided with the chemagic 360-D instrument.

Kit-related protocol files are available on the webpage or will be provided by customer support (see section “REQUIRED PROTOCOLS FILES”).



## 8. ELECTRONIC INSTRUCTIONS FOR USE

Electronic Instructions for Use (eIFU) in different languages are available on our webpage.

To download these electronic Instructions for Use please visit

<https://chemagen.com/products/ce-ivd-chemagic-kits/ivd-1074-chemagic-bbs-dna-kit-h24/>.

The eIFU are provided in at least English (EN), French (FR), Spanish (ES) and Italian (IT) and upon request also in other required languages.

In case of any questions regarding download or the electronic Instructions for Use please contact us: [support.chemagen@revvity.com](mailto:support.chemagen@revvity.com), [info.chemagen@revvity.com](mailto:info.chemagen@revvity.com) or +49 (0) 2401805500.

## 9. WARNINGS AND PRECAUTION

For *in vitro* diagnostic use.

The product is intended for laboratory personnel trained for the chemagic 360-D instrument in combination with chemagic nucleic acid purification kits.

A thorough understanding of this IFU and the chemagic 360-D User Manual is a prerequisite and necessary for successful use of the chemagic BBS DNA Kit H24.

The reagents supplied with this kit are intended for use as an integral unit. Do not mix identical reagents from kits with different lot numbers.

Do not use kit reagents after the expiry date printed on the kit label. Once opened, the reagents can be used for the time period stated in the reagent listing of this IFU.

Any deviation from the protocol may affect the results.

The reagents are automatically dispensed in whole rows and therefore the chemagic Tips XL and the chemagic Elution Tubes 13 mL should be used also in whole rows on each rod in contact with any reagent solution.

It should also be noted that if partial plates are run, the solutions may not be sufficient for 240 extractions.

Check all kit components for integrity. In case of damage, contact your supplier.

Handle all specimens as potentially infectious. Potentially infectious samples shall be inactivated. Please refer to the U.S. Department of Health and Human Services publication "Biosafety in Microbiological and Biomedical Laboratories" or any other

local or national regulation.

Lysis Buffer 1 contains guanidinium chloride and is harmful if swallowed, in contact with skin or if inhaled. Binding Buffer 2, Wash Buffer 3 and Wash Buffer 4 contain sodium perchlorate and ethanol and are flammable liquids and vapors and are harmful if swallowed. Wash Buffer 5 contains ethanol and is a flammable liquid and vapor. Proteinase K contains Tritirachium album serine Proteinase and causes skin irritation and serious eye irritation. It may cause allergy or asthma symptoms or breathing difficulties or respiratory irritation if inhaled. See specific precautions for all components in the section "KIT REAGENTS AND SAFETY INFORMATION".

To avoid injuries when working with the kit components, always wear safety glasses, disposable gloves, and protective clothing. For detailed information, please refer to the corresponding safety data sheets (SDS) available on our webpage.

Follow local regulations for handling of ethanolic solutions.

Disposal of all waste should be in accordance with local regulations.

## 10. KIT REAGENTS AND SAFETY INFORMATION

The chemagic BBS DNA Kit H24 contains the following reagents.

### 10.1 MAGNETIC BEADS

Component	Quantity	Shelf life and storage
Magnetic Beads	1 bottle (volume see label)	+2 to +25 °C until expiry date stated on the bottle label.  Once opened, stable for 60 days at +2 to +25 °C.

Suspension of particles containing nanoparticulate iron oxide encapsulated in a matrix of polyvinyl alcohol. Magnetic Beads bind the DNA/ RNA during the extraction process.

### 10.2 LYSIS BUFFER 1

Component	Quantity	Shelf life and storage
Lysis Buffer 1	1 canister (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.



WARNING

Ready-for-use aqueous buffer (pH 6.7-7.2) solution containing guanidinium chloride (30–50 %) and isotridecyl alcohol (1-1.5 %). Lysis Buffer 1 is used to lyse the cells or other DNA source present in the sample to release DNA in solution.

**CAUTION! Lysis Buffer 1 contains guanidinium chloride and isotridecyl alcohol.**

#### Hazard, precautionary and EUH phrases

H302	Harmful if swallowed.
H315	Causes skin irritation.
H319	Causes serious eye irritation.
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection/ hearing protection.

---

**Hazard, precautionary and EUH phrases**



---

P301+P312	IF SWALLOWED: Call a POISON/ doctor if you feel unwell. P330 Rinse mouth.
P330	Rinse mouth.
P305+P351+P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
P332+P313	If skin irritation occurs: Get medical advice/ attention.
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

---

**10.3 BINDING BUFFER 2**


---

Component	Quantity	Shelf life and storage
Binding Buffer 2 	2 canisters (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.
DANGER		

Ready-for-use Tris-HCl-buffered (pH 5.2–5.9) solution with sodium perchlorate (20–30 %), ethanol (30–50 %) and acetic acid (0.75–1.5 %). Binding Buffer 2 is used to create the appropriate conditions to get the DNA bound to the Magnetic Beads.

**CAUTION! Binding Buffer 2 contains ethanol and sodium perchlorate.**

---

**Hazard, precautionary and EUH phrases**


---

H226	Flammable liquid and vapor.
P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P240	Ground and bond container and receiving equipment.
P241	Use explosion-proof [electrical/ ventilating/ lighting] equipment.
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection/ hearing protection.

---

**Hazard, precautionary and EUH phrases**



---

P303+P361+P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower].
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

---

**10.4 WASH BUFFER 3**


---

Component	Quantity	Shelf life and storage
Wash Buffer 3 	1 canister (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.
<b>DANGER</b>		

---

Ready-for-use Tris-HCl-buffered (pH 5.0–5.6) solution with sodium perchlorate (10–20 %) and ethanol (10–30 %). Used for removing non-DNA contaminants during washing step.

**CAUTION! Wash Buffer 3 contains ethanol and sodium perchlorate.**

---


**Hazard, precautionary and EUH phrases**


---

H226	Flammable liquid and vapor.
P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P240	Ground and bond container and receiving equipment.
P241	Use explosion-proof [electrical/ ventilating/ lighting] equipment.
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection/ hearing protection.
P303+P361+P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower].
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

---

## 10.5 WASH BUFFER 4

Component	Quantity	Shelf life and storage
Wash Buffer 4 	1 canister (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.
DANGER		


Ready-for-use Tris-HCl-buffered (pH 5.0–5.6) solution with sodium perchlorate (10–20 %) and ethanol (10–30 %). Used for removing non-DNA contaminants during washing step.

**CAUTION! Wash Buffer 4 contains ethanol and sodium perchlorate.**

### Hazard, precautionary and EUH phrases

H226	Flammable liquid and vapor.
P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P240	Ground and bond container and receiving equipment.
P241	Use explosion-proof [electrical/ ventilating/ lighting] equipment.
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection/ hearing protection.
P303+P361+P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower].
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

## 10.6 WASH BUFFER 5

Component	Quantity	Shelf life and storage
Wash Buffer 5  DANGER	1 canister (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.

Ready-for-use solution contains ethanol (50–70 %). Used for removing last traces of non- DNA contaminants during washing step.

**CAUTION! Wash Buffer 5 contains ethanol.**

### Hazard, precautionary and EUH phrases

H225	Highly flammable liquid and vapor.
P210	Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking.
P240	Ground and bond container and receiving equipment.
P241	Use explosion-proof [electrical/ ventilating/ lighting] equipment.
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection/ hearing protection.
P303+P361+P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water [or shower].
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

## 10.7 WASH BUFFER 6

Component	Quantity	Shelf life and storage
Wash Buffer 6	1 canister (volume see label)	+2 to +25 °C until expiry date stated on the canister label.  Once opened, stable for 60 days at +2 to +25 °C.



Ready-for-use ultra-filtered water solution. Used for removing possible residuals of ethanol.

## 10.8 ELUTION BUFFER 7

Component	Quantity	Shelf life and storage
Elution Buffer 7	1 bottle (volume see label)	+2 to +25 °C until expiry date stated on the bottle label.  Once opened, stable for 60 days at +2 to +25 °C.

Ready-for-use 10 mM Tris-HCl-buffered (pH 7.8–8.4) solution.

## 10.9 PROTEINASE K

Component	Quantity	Shelf life and storage
Proteinase K  	4 glass vials (lyophilized)	+2 to +25 °C until expiry date stated on the vial label.  Once reconstituted, stable for 28 days at +2 to +8 °C.
DANGER		

The Proteinase K is reconstituted by adding 1.25 mL of purified water. Proteinase K is added to enhance the efficiency of the lysis step.

**CAUTION! Proteinase K contains Proteinase, Tritirachium album serine and calcium acetate hydrate.**

### Hazard, precautionary and EUH phrases

H315	Causes skin irritation.
H319	Causes serious eye irritation.
H334	May cause allergy or asthma symptoms or breathing difficulties if inhaled.
H335	May cause respiratory irritation.
P261	Avoid breathing dust/ fume/ gas/ mist/ vapors/ spray.
P280	Wear protective gloves/ eye protection/ face protection.
P284	[In case of inadequate ventilation] wear respiratory protection.



---

**Hazard, precautionary and EUH phrases**


---

	IF IN EYES: Rinse cautiously with water for several minutes. P305+P351+P338 Remove contact lenses, if present and easy to do. Continue rinsing.
P405	Store locked up.
P501	Dispose of contents/ container in accordance with local/ regional/ national/ international regulations.

---

**10.10 FURTHER KIT COMPONENTS**

The chemagic BBS DNA Kit H24 contains the following plastic material.

<b>Component</b>	<b>Quantity</b>	<b>Storage</b>
chemagic Tips XL	250	+2 to +25 °C
chemagic Deep Well Plate 24 XL	50	+2 to +25 °C
chemagic Elution Tube 13 mL	288	+2 to +25 °C

---

## 11. REQUIRED PROTOCOLS FILES

The following protocol files will be provided by Revvity chemagen Technologie GmbH and are available on the webpage or will be provided by customer support.

<b>Protocol (.che file)</b>	<b>Protocol type/ purpose</b>
chemagic BBS DNA 360 H24 EB 13ml Tubes prefilling VD191213.che	Kit-related extraction file for the chemagic 360-D instrument
prime manifolds H24 all 360 V150116.che	Filling and priming the chemagic 360-D instrument tubing with reagents
check manifolds H24 all 360 V150116.che	Checking the functionality of the pumps
regular cleaning procedure 24 dispenser 360 V150116.che	Regular cleaning of the chemagic 360-D instrument (once per week)
intensive cleaning procedure H24 dispenser 360 V150116.che	Intensive cleaning of the chemagic 360-D instrument (once per month)

## 12. MATERIAL REQUIRED BUT NOT SUPPLIED WITH THE KIT

The chemagic BBS DNA Kit H24 requires the following items.

### 12.1 ITEMS FROM REVVITY CHEMAGEN TECHNOLOGIE GMBH

<b>Item</b>	<b>Product no.</b>
chemagic 360-D instrument	2024-0010
chemagic 24 Rod Head Set	CMG-376
Racks for chemagic Elution Tubes 13 mL (supplied with chemagic 24 Rod Head Set)	-

**12.2 ADDITIONAL REQUIRED ITEMS**

<b>Item</b>	<b>Purpose</b>
Pipettes and pipette tips with aerosol barriers	Prefilling of Magnetic Beads, Elution Buffer 7 and Proteinase K
Molecular biology grade water	Reconstitution of the Proteinase K
70% Ethanol	Cleaning of the chemagic 360-D instrument

**12.3 ADDITIONAL OPTIONAL ITEMS FROM REVVITY CHEMAGEN TECHNOLOGIE GMBH**

<b>Product</b>	<b>Product no.</b>
chemagic Stand 12 (supplied with the chemagic 24 Rod Head Set)	CMG-308
Red Cell Lysis Buffer	CMG-848

**12.4 OTHER ADDITIONAL OPTIONAL ITEMS**

<b>Product</b>	<b>Purpose</b>
Isotonic saline solution, sterile	Resuspension of buffy coats

### 13. SPECIMEN COLLECTION AND HANDLING

The chemagic BBS DNA Kit H24 is usable with human blood, buffy coat and saliva in aliquots of 2 – 4 mL per isolation.

Human whole blood (2 – 4 mL) or buffy coat samples (up to 4 mL) that are fresh, frozen or stored typically for a maximum of 10 days at +2 to +8 °C should be used. For long-term storage, freezing at –20 °C or –80 °C in aliquots is recommended. The recommended blood stabilizers are EDTA or citrate.

**NOTE: The use of heparin-stabilized blood samples can cause inhibition in downstream applications and is therefore not recommended.**

The white blood cell count in the whole blood sample decreases during storage. Prolonged storage of the samples may cause a poor yield of the DNA after extraction.

Human saliva (4 mL) stored according to the collection tube provider instructions should be used. The recommended stabilized collection tubes are from DNAgenotek<sup>®</sup>, Isohelix<sup>™</sup> and Spectrum Solutions. Incubation of the collection tubes prior to the extraction for > 2 h at 50 °C will result in higher DNA yields and is therefore recommended.

Buffy coat samples stored for a maximum of one week at +2 to +8 °C should be used. For long-term storage, freezing at –20°C or –80°C in aliquots is recommended. The buffy coats should be derived from stabilized blood tubes (follow tube recommendation above for blood samples). Prior to the extraction, buffy coats should be thawed at 37 °C. We recommend the following procedure for buffy coat preparation.

## 14. PREPARATION OF BUFFY COAT SAMPLES

- Add up to 5 mL of fresh whole blood into a sterile 50 mL tube.
- Add 40 mL Red Cell Lysis Buffer (RCLB) to the blood and invert the tube 4 times.
- Incubate for 5 minutes or until the suspension becomes translucent.
- Centrifuge at 4,000 rpm for 10 minutes to collect the white blood cells.
- Decant the supernatant and carefully aspirate the remaining supernatant from the top of the sample by pipetting.

**NOTE: Be careful not to disturb the cell pellet. Exercise caution in pipetting to avoid loss of the white blood cell pellet.**

- Add 20 mL Red Cell Lysis Buffer and carefully wash and remove remaining red cells on top of the white pellet without disturbing the white pellet.
- Decant the supernatant and carefully aspirate the remaining supernatant from the top of the sample by pipetting.
- Resuspend the buffy coat in up to 2 mL isotonic saline solution (0.9 % NaCl).
- If the starting volume of blood is less or more than 6 mL, change the volume of Red Cell Lysis Buffer used proportionately.
- The buffy coats can be frozen, stored at +2 to +8 °C for one week, or directly used for extraction.

## 15. DETAILED PROTOCOL DESCRIPTION

### 15.1 PROTOCOL PROCEDURE

The following procedure describes the preparation and the execution of the extraction protocol using the chemagic 360-D instrument.

The duration of the automated extraction protocol is approximately 78 minutes.

The protocol is suitable for processing up to 24 samples in parallel (see “PROCESSING STEPS” below). For detailed instructions on the use of the chemagic 360-D instrument, please refer to the chemagic 360-D User Manual.

**NOTE: Samples and reagents must be brought to room temperature (+19 to +25 °C) before use.**

Connect the reagent bottles to the chemagic 360-D instrument as follows:

Pump	Buffer	Minimum filling volume
Pump 1	Lysis Buffer 1	150 mL
Pump 2	Binding Buffer 2	350 mL
Pump 3	Wash Buffer 3	200 mL
Pump 4	Wash Buffer 4	200 mL
Pump 5	Wash Buffer 5	200 mL
Pump 6	Wash Buffer 6	200 mL

**NOTE: Recap the bottles tightly immediately after use or keep the bottles connected tightly to the chemagic 360-D instrument. Binding Buffer 2, Wash Buffer 3, Wash Buffer 4 and Wash Buffer 5 contain ethanol. If ethanol evaporates, the optimal yield or detection sensitivity cannot be guaranteed.**

## 15.2 PROCESSING STEPS

1. Check all kit components for integrity. In case of damage, contact your supplier.
2. Before prefilling the plates and the chemagic Elution Tubes 13 mL in the rack, mark them with material in position (samples, Magnetic Beads and buffers).

The reagents are automatically dispensed in whole rows and therefore the chemagic Elution Tubes 13 mL should be used also in whole rows on each rod in contact with any reagent solution.

3. Reconstitute the Proteinase K:

Component	Reconstitution
Proteinase K	Add 1.25 mL molecular biology grade water to Proteinase K vial and mix gently until dissolved.

4. Fill and prime the chemagic 360-D tubing with reagents by choosing the protocol “**prime manifolds H24 all 360 V150116.che**”. Press [Insert IDs], follow the instructions given in the chemagic QA software and start priming by pressing [OK]. If functions enabling the ID data input are deactivated, start priming directly by pressing [Start].

**NOTE: Priming needs to be done when reagent bottles are connected to the chemagic 360-D instrument for the first time or when the instrument’s tubing is not already filled with the above mentioned reagents.**

5. If priming is not needed, select the protocol “**check manifolds H24 all 360 V150116.che**” and press [Insert IDs] or - if the enhanced functions are deactivated - [Start]. A small volume of buffer will be dispensed by each pump sequentially starting with the first pump used for this application. If one of the pumps does not show dispensing of buffer through all nozzles, please use the corresponding priming protocol for this pump. Performing several runs a day it is only necessary to check the pumps once at the beginning of the day.
6. Select the protocol “**chemagic BBS DNA 360 H24 EB 13ml prefilling VD191213.che**” and press [Insert IDs] and follow the instructions given in the chemagic QA software.
7. Ensure chemagic Tips XL are sufficient and aligned with the positions of the samples and place the Tip Rack in position 1 on the tracking system. The reagents are automatically dispensed in whole rows and therefore the chemagic Tips XL should be used also in whole rows on each rod in contact with any

reagent solution.

8. Check the volumes in the buffer supply containers and confirm by pressing [OK]. See above “PROTOCOL PROCEDURE” minimum filling heights.

**NOTE: Take care that all buffer supply bottles contain enough buffer. Only if the liquid level for all buffers is sufficient 24 isolations can be performed.**

9. Select the number of samples for prefilling by using the drop down menu. The scheme for positioning the samples will be shown after selecting. Take care to use the given positions. Confirm by pressing [OK].
10. Prefill the selected wells of the sample plate with up to 4 mL sample. To ensure the homogeneity of the samples, mix the samples gently prior to pipetting in the wells of the sample plate.
11. Load two Elution Racks with chemagic Elution Tubes 13 mL.
12. Prefill Elution Buffer 7 and the thoroughly resuspended Magnetic Beads by pipetting manually according to each corresponding well or tube in use.

Component	Plate position on chemagic 360-D instrument	Volume/ well or tube
Magnetic Beads	4	400 µL
Elution Buffer 7	7	300–500 µL

**NOTE: The Magnetic Bead suspension should be mixed vigorously before dispensing; otherwise the suspension is not homogenous and the DNA yield could be low.**

13. Add 15 µL Proteinase K to the wells containing sample.
14. Place the chemagic Deep Well Plates 24 XL and the two racks with chemagic Elution Tubes 13 mL (one rack with chemagic Elution Tubes 13 mL prefilled with Elution Buffer 7 and the other rack with empty chemagic Elution Tubes 13 mL for the disposal of the chemagic Tips XL) on the tracking system according to the instructions given by the chemagic QA software.
15. Place the sample plate in position 2 on the tracking system.
16. Check all plates and racks for accurate orientation and fitting.
17. Close the front door and start the process by pressing [Start].
18. The automated DNA extraction process is initiated.



19. After the isolation procedure has finished use the [Turn Table] button to unload the tracking system. Each click on [Turn Table] moves the tracking system (table) clockwise by one position.

**ATTENTION! Never move the tracking system (table) manually. This might damage the instrument. All movements must be performed with the [Turn Table] function.**

**NOTE: Opening the chemagic 360-D instrument door while the automated extraction run is ongoing, will terminate the run and the samples in process may be lost.**

For information on cleaning the instrument see section "CLEANING AND MAINTENANCE".

### 15.3 SHORT DESCRIPTION/ QUICK GUIDE

#### Automated DNA extraction run on chemagic 360-D instrument:

- Select the protocol “**check manifolds H24 all 360 V150116.che**” to flush the tubing prior to starting the automated extraction run.
- Press [Insert IDs], follow the instructions given in the chemagic QA software and start flushing by pressing [OK].
- When using the functions enabling the ID data input, select the protocol “**chemagic BBS DNA 360 H24 EB 13 ml Tubes prefilling VD191213.che**” and press [Insert IDs]. Follow the instructions given in the chemagic QA software to fill in the required data.
- The reagents are automatically dispensed in whole rows and therefore the chemagic Tips XL and the chemagic Elution Tubes 13 mL should be used also in whole rows on each rod in contact with any reagent solution.
- Load the plates and the racks with chemagic Elution Tubes 13 mL on the tracking system positions 1-8 as follows.  
(Numbers on tracking system refer to the positioning of the plate on the chemagic 360-D instrument.)

Position on tracking system	Material in position	Protocol step in detail
1	Rack with chemagic Tips XL	Use chemagic Tips XL according to the positions of the samples and place the rack.  <b>NOTE: Tips need to be present in the rack in full rows.</b>
2	Sample plate (chemagic Deep Well Plate 24 XL)	Place the plate with prepared samples (up to 4 mL sample) and 15 $\mu$ L Proteinase K. Lysis Buffer 1 and Binding Buffer 2 are dispensed in the plate automatically.
3	chemagic Deep Well Plate 24 XL with 400 $\mu$ L Magnetic Beads	Pipette 400 $\mu$ L thoroughly resuspended Magnetic Beads in each well of the plate according to sample positions and place the plate. Wash Buffer 3 is dispensed in the plate automatically.
4	chemagic Deep Well Plate 24 XL	Place empty plate. Wash Buffer 4 is dispensed in the plate automatically.
5	chemagic Deep Well Plate 24 XL	Place empty plate. Wash Buffer 5 is dispensed in the plate automatically.
6	chemagic Deep Well Plate 24 XL	Place empty plate. Wash Buffer 6 is dispensed in the plate automatically.
7	Rack with chemagic Elution Tubes 13 mL prefilled with 300 – 500 $\mu$ L Elution Buffer 7	Pipette 300 – 500 $\mu$ L Elution Buffer 7 to each chemagic Elution Tubes 13 mL according to the sample positions and place the rack.

Position on tracking system	Material in position	Protocol step in detail
8	Rack with empty chemagic Elution Tubes 13 mL	Place the rack.

- Check all plates and racks for accurate orientation and fitting.
- After all plates and racks are in place, press [OK].
- Close the front door and start the DNA extraction process immediately by pressing [Start]. Subsequently the sample lysate will be mixed automatically.
- If the functions enabling the ID data input are deactivated, load the plates on the tracking system positions 1-8.
- After all plates are in place, select the protocol “**chemagic BBS DNA 360 H24 EB 13 ml Tubes prefilling VD191213.che**”, mark the columns in use on the plate map in the dialog and start the extraction run directly by pressing [Start].
- After the isolation procedure has finished use the [Turn Table] button to unload the tracking system. Each click on [Turn Table] moves the tracking system (table) clockwise by one position.

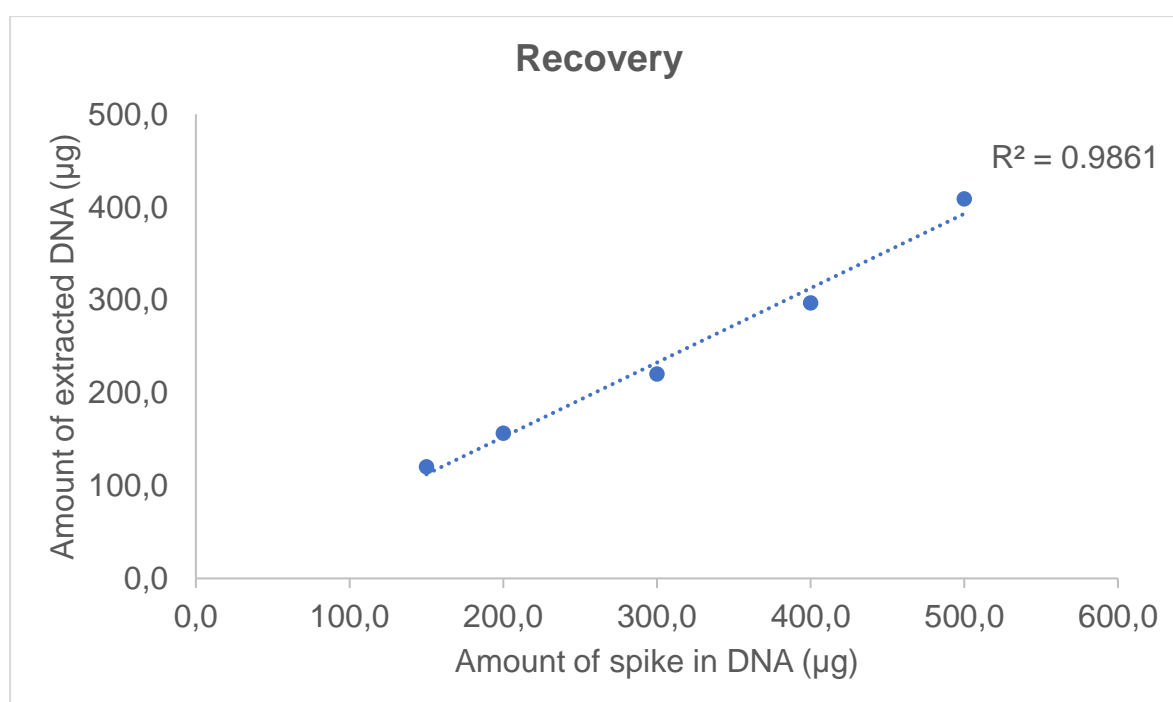
**ATTENTION: Never move the tracking system (table) manually. This might damage the instrument. All movements must be performed with the [Turn Table] function.**

**NOTE: Opening the chemagic 360-D instrument door while the automated extraction run is ongoing, will terminate the run and the samples in process may be lost.**

## 16. PERFORMANCE CHARACTERISTICS

### 16.1 LINEARITY AND RECOVERY WITH SPIKED DNA SAMPLE

To show linearity of the extraction with IVD-1074 kit, 0.9 % NaCl solution was spiked with five concentrations of human genomic DNA; 150 µg, 200 µg, 300 µg, 400 µg and 500 µg. Extractions were performed using 4 mL of the various DNA amounts as sample with the extraction protocol “**chemagic BBS DNA 360 H24 prefilling VD190913.che**”. Four replicates of each DNA amount spiked in 0.9 % NaCl were extracted.



**Figure 1:** Extraction recovery of spike in DNA samples. Extracted with the chemagic 360-D using the IVD-1074 kit.

The recovery shows a good linearity in the range of 150 – 500 µg of DNA as sample input. The recovery ranges between 69.8 and 98.9 % depending on the input amount of human genomic DNA.

## 16.2 DNA YIELDS WITH BLOOD AND BUFFY COAT SAMPLES

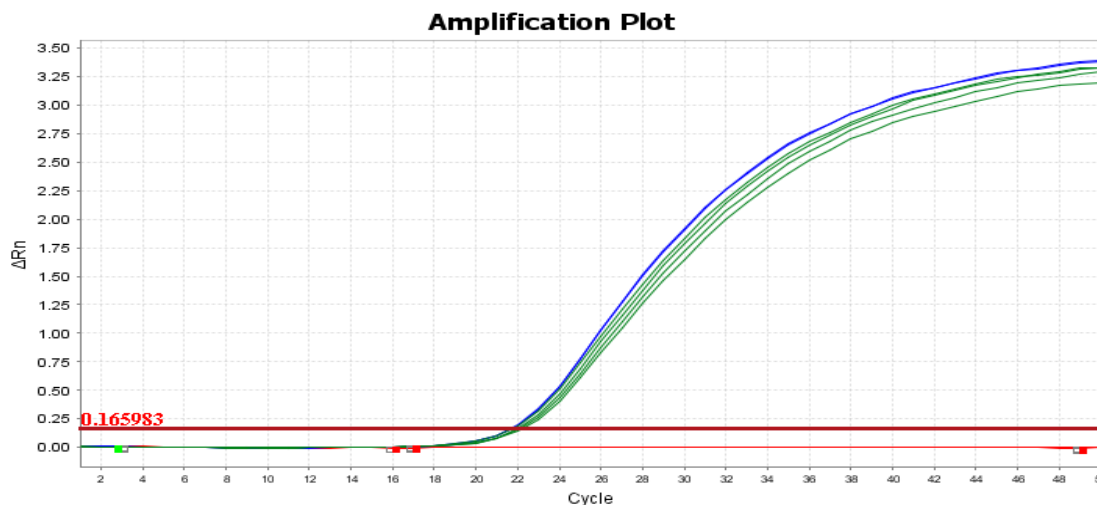
The expected DNA yields for the extraction from human blood and buffy coat is depending on the number of white blood cells. The number of the extracted white blood cells is determined by the input volume and the white blood cell count (WBC). For most of the samples the white blood cell count will not be known, but for healthy individuals it is in the range of 4 – 10 mio. white blood cells per mL of blood. The IVD-1074 kit using the “**chemagic BBS DNA 360 H24 prefilling VD190913.che**” protocol extracts in average 4.38 pg per white blood cell. Using 4 mL of blood with a white blood cell count of 8.2 mio. white blood cells per mL of blood as input is expected to result in 150 µg DNA.

**Table 1:** Average yield, %CV (standard deviation) and purity of extracted DNA using the IVD-1074 kit extracted with the chemagic 360-D instrument.

Sample Material / Storage Condition	Volume [mL]	WBC [mio. cells/ mL Blood]	Average Yield [µg]	CV [%]	Average Purity [260/280]
Blood 1 / 4 °C	4	8.2	150.0	9.2	1.84
Blood 1 / -20 °C	4	8.2	141.1	8.3	1.79
Blood 3 / 4 °C	2	6.2	73.3	2.4	1.85
Blood 4 / 4 °C	2	5.7	61.7	1.9	1.86
Blood 4 / 4 °C	4	5.7	98.7	12.0	1.86
Blood 4 / -20 °C	4	5.7	94.2	8.2	1.81
Buffy Coat*	4	5.7	125.5	10.0	1.72
Saliva / 4 °C	4	n.a.	42.3	20.9	1.78

\* Buffy Coat was generated from 5 mL of Blood 4

From all extractions including different sample materials, storage conditions and input volumes, specific eluates were used in a human albumin qPCR to prove the suitability of the extracted DNA for enzymatic reactions. All eluates performed without any issues - see example qPCR plot below.



**Figure 2:** qPCR curves of DNA eluates extracted with the chemagic 360-D using the IVD-1074 kit. Red - negative control, blue – positive control, green extracted DNA from Blood 1.

## 17. CLEANING AND MAINTENANCE

Cleaning and maintenance of the system is described in detail in the chemagic 360-D User Manual. The system cleaning is performed once per week. Clean the chemagic Dispenser as follows.

- Select the protocol “**regular cleaning procedure 24 dispenser 360 V150116.che**” and press [Insert IDs] or [Start] if the enhanced functions are deactivated. Follow the instructions as given in the software.
- Prior to the next use of the chemagic Dispenser perform the appropriate priming protocol.
- The cleaning of the chemagic Dispenser with 70 % ethanol is recommended once per month. Simply use the “**intensive cleaning procedure H24 dispenser 360 V150116.che**” instead of the regular one for this purpose.
- If the chemagic Dispenser will not be used for longer period of time, it is mandatory to perform the "regular cleaning procedure" to maintain the performance of the instrument when bringing it back into service.

## 18. DOWNSTREAM APPLICATIONS

The following downstream applications were successfully performed and described in literature after isolation of genomic DNA.

**Table 2:** Peer reviewed and published downstream applications.

Sample Material	Downstream Application	Title	Reference
Blood	Genotyping, DNA methylation assay, PCR, Sequencing	Identification of dynamic glucocorticoid-induced methylation changes at the FKBP5 locus	Wiechmann et al. Clinical Epigenetics 11 Article Number 83 (2019) <a href="https://clinicalepigeneticsjournal.biomedcentral.com/articles/10.1186/s13148-019-0682-5#Abs1">https://clinicalepigeneticsjournal.biomedcentral.com/articles/10.1186/s13148-019-0682-5#Abs1</a>
Whole Blood	Methylation Analysis	A new molecular classification to drive precision treatment strategies in primary Sjögren's syndrome	Soret P. et al. Nature Communications 12, Article number: 3523 (2021) <a href="https://www.nature.com/articles/s41467-021-23472-7">https://www.nature.com/articles/s41467-021-23472-7</a>
Lymphocytes	Long-range PCR, NGS, Sanger Sequencing, MLPA	Advances in genetic technologies result in improved diagnosis of mismatch repair deficiency in colorectal and endometrial cancers	Evans D.G. et al. Med Genet: first published as 10.1136/jmedgenet-2020-107542 (2021) <a href="https://pubmed.ncbi.nlm.nih.gov/33452216/">https://pubmed.ncbi.nlm.nih.gov/33452216/</a>
Saliva	Genome sequencing, qPCR	Genetic Effect of Chemotherapy Exposure in Children of Testicular Cancer Survivors	Gregory V. et al. Clinical Cancer Research Volume 22, Issue 9 (2016) <a href="https://aacrjournals.org/clinicalres/article/22/9/2183/79766/Genetic-Effect-of-Chemotherapy-Exposure-in">https://aacrjournals.org/clinicalres/article/22/9/2183/79766/Genetic-Effect-of-Chemotherapy-Exposure-in</a>



## 19. FURTHER QUESTIONS

For further application, technical questions, or more information on how the data was generated please contact [support.chemagen@revvity.com](mailto:support.chemagen@revvity.com) or +49 (0) 2401805500.

## 20. LIMITATIONS OF THE PROCEDURE

The IVD-1074 kit is validated for the extraction of DNA from blood, buffy coat and saliva. Other sample materials such as tissue, cell or buccal swab lysates may be compatible but have not been validated. For such materials, a validation must be performed by the user.

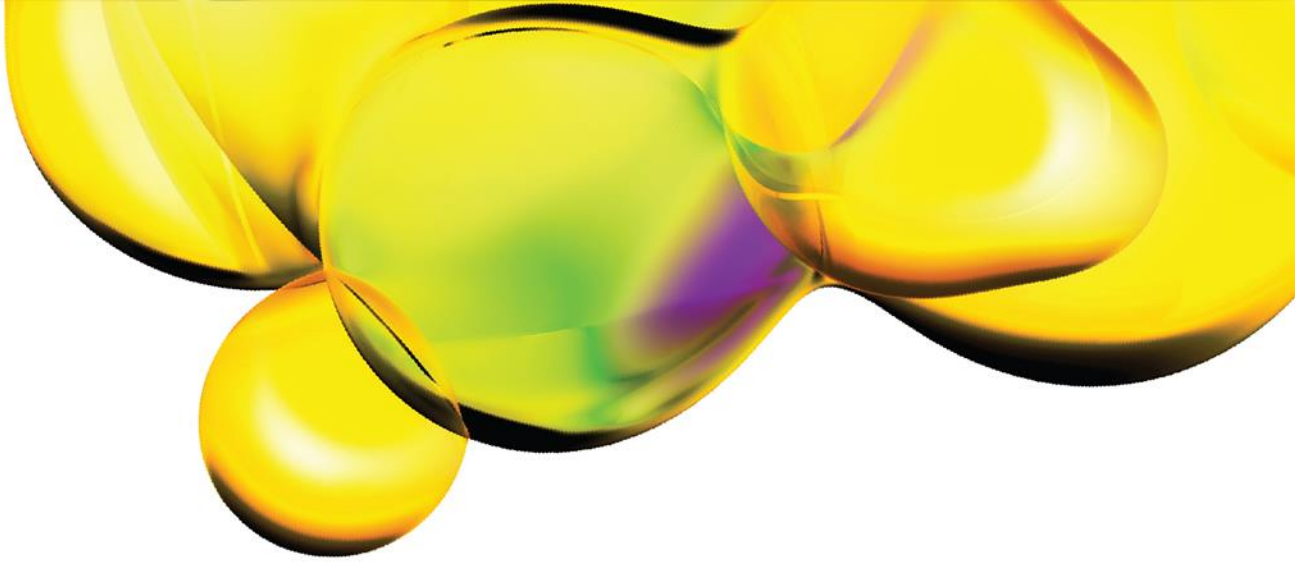
The use of heparin-stabilized blood samples can cause inhibition in downstream applications and is therefore not recommended.

## 21. INFLUENCE OF INTERFERING SUBSTANCES

The effect of interfering substances contained in human whole blood possibly interfering with DNA extraction were tested in whole blood. The tested substances and concentrations are presented in table below. Based on the results, it was concluded that the tested substances do not interfere with the DNA extraction.

**Table 3:** Influence of interfering substances.

<b>Interfering substances</b>	<b>Concentration [µg/mL]</b>	<b>Interference</b>
Bilirubin conjugated	332	No
Bilirubin unconjugated	200	No
Triglycerides	30	No
Human Serum Albumin	30	No



## 22. WARRANTY

Any change or modification of the procedure not recommended by the manufacturer may affect the results, in which event Revvity chemagen Technologie GmbH and its affiliates disclaim all warranties expressed, implied or statutory including the implied warranty of merchantability and fitness for use.

Revvity chemagen Technologie GmbH, its affiliates and its authorized distributors, in such an event, shall not be liable for damages indirect or consequential.

May 2024

[www.revvity.com](http://www.revvity.com)

revvity