

# Certificate Of Analysis

(Analysenzertifikat)

# HOLTSCH

Medizinprodukte GmbH

**Product / Lot-No.** (Produkt / Ch.bez.): *Quickpad* *060321*  
**Date of Manufacturing** (Herstellungsdatum): *02.03.-09.03.2021*  
**Expiry Date** (Verfallsdatum): *02/2023*

	<b>Specification</b> (Spezifikation)	<b>Result</b> (Ergebnis)
<b>Description</b> (Beschreibung):	milky-white to yellowish transparent plastic container, filled with white and separately detachable non-woven pads and isopropanol 70% V/V Milchig-weiße bis leicht gelbliche, durchsichtige Kunststoffdose, befüllt mit weißen, einzeln abreißbaren Vliestupfern und Isopropanol 70% (V/V)	<i>complies (entspricht)</i>
<b>Leak tightness and intactness of container</b> (Dichtigkeit und Unversehrtheit des Behältnisses):	fully welded and intact vollständig verschweißt und unversehrt	<i>complies (entspricht)</i>
<b>Sterility</b> (Sterilität):	sterile (steril)	<i>complies (entspricht) <sup>1)</sup></i>

<sup>1)</sup> The sterility of product is ensured by the process of irradiation and is tested only at randoms intervalls.  
Die Sterilität des Produktes wird durch den Bestrahlungsprozess sichergestellt und nur stichprobenartig geprüft.

**Summary (Gesamtbeurteilung):**  
The above mentioned lot complies with the specification.  
Die o.g. Charge entspricht der Spezifikation.

**Comments** (Bemerkungen): *none (keine)*

**Date / Signature Head of Quality Control:**  
(Datum / Unterschrift Leiter Qualitätskontrolle)

*25.06.21 Wiebe*

Zertifikat  
Protokoll **BESTRAHLUNG**  
certificate  
protocol **IRRADIATION**

Auftragsnummer (AN) : 181184/2021  
order number (internal)  
Kundennummer : 728  
Customer number

DB Nr	181184
BP Nr	239.691

Synergy Health Radeberg GmbH; Juri-Gagarin-Str.15;01454 Radeberg

**Auftraggeber (Customer)**

**Auftragnehmer (performer)**

Holtsch Medizinprodukte GmbH  
  
Leipziger Str. 300  
**01139 Dresden**

Synergy Health Radeberg GmbH  
  
Juri-Gagarin-Str.15  
**01454 Radeberg**  
Tel.: +49 3528 / 4364 - 16  
Fax: +49 3528 / 4364 - 99

Auftragsnummer (AG) : 20.30.2021  
order number Customer

Auftragsdatum : 30.03.2021  
date of order

Name des Produktes : Quickpad (Kosmetik)  
name of product

Artikelcode :  
item code

Charge : 060321, Vref.: 02.2023  
batch number

Bestrahlungszyklus : A-14p/21  
irradiation cycle

Spezifikation : DD0728\_0440\_2020\_R34  
specification

Bestrahlungsdatum : 11.04.2021  
irradiation date

Identifikationsnummer : 845555 - 845558  
identification number

Bestrahlungseinheiten : EP-Palette  
irradiation unit

Anzahl : 4  
quantity

Verpackungseinheiten : Palette  
packaging unit

Anzahl : 2  
quantity

Besonderheiten :  
remarks

Die Bestrahlung erfolgt gemäß AAMI TIR33 / ISO 11137 und Produktspezifikation.  
The irradiation has been performed in accordance to AAMI TIR33/ ISO 11137 as well as to product specification.

Dosis  
dose

Energiedosis laut Auftrag  
dose in accordance with order

Dosimetrisch ermittelte Dosis  
calculated dose

Identnummer :  
identification number

**min. Dosis:**  
minimum dose 25,0 kGy

28,11 kGy

845558

**max. Dosis:**  
maximum dose 40,0 kGy

36,20 kGy

845557

Unterbrechungszeit >24 h : Nein  
interruption time over 24 hours

**Anlage** : GS3000  
(facility)

**Strahlungsart** : Co-60  
(kind of radiation)

Erstellung und Datenübernahme  
preparation and data transfer

13.04.2021

Freigabe des oben genannten Bestrahlungsgutes zur  
Auslieferung Release of specified product for delivery

Datum und Unterschrift Ersteller  
date and signature originator

13.04.2021

Protokollsignatur 83adfc25e4a38187c9e5292ff18c471f

Datum und Unterschrift Qualität  
date and signature quality



Zertifikat-Nr./Certificate no:  
DE\_SN\_01\_GMP\_2020\_0038

Aktenzeichen/Reference Number:  
26-5117/90

**BESTÄTIGUNG DER ÜBEREINSTIMMUNG EINES  
HERSTELLERS MIT GMP**

**Teil 1**

**Ausgestellt nach einer Inspektion gemäß**

- Art. 111 (5) der Richtlinie 2001/83/EG

Die zuständige deutsche Überwachungsbehörde bestätigt:

Der Hersteller  
**HOLTSCH Medizinprodukte GmbH**

Anschrift der Betriebsstätte  
**HOLTSCH Medizinprodukte GmbH  
Leipziger Straße 300  
01139 Dresden  
Deutschland**

- wurde im Rahmen der nationalen Arzneimittelüberwachung inspiziert in Verbindung mit der Herstellungserlaubnis Nr. DE\_SN\_01\_MIA\_2016\_0010 gemäß
  - Art. 40 der Richtlinie 2001/83/EGumgesetzt in deutsches Recht durch:  
§ 13 Abs. 1 und § 72 Arzneimittelgesetz

Aufgrund der aus der letzten Inspektion vom 24. Januar 2019 gewonnenen Erkenntnisse wird für die oben genannte Betriebsstätte des Herstellers die Übereinstimmung mit den Anforderungen der Guten Herstellungspraxis festgestellt, die sich aus

- den Grundsätzen und Leitlinien der Guten Herstellungspraxis gemäß
  - Richtlinie 2003/94/EG

ergeben.

**CERTIFICATE OF GMP COMPLIANCE OF A  
MANUFACTURER**

**Part 1**

**Issued following an inspection in accordance with**

- Art. 111 (5) of Directive 2001/83/EC

The competent authority of GERMANY confirms the following:

The manufacturer  
**HOLTSCH Medizinprodukte GmbH**

Site address  
**HOLTSCH Medizinprodukte GmbH  
Leipziger Straße 300  
01139 Dresden  
Germany**

- has been inspected under the national inspection programme in connection with manufacturing authorisation no. DE\_SN\_01\_MIA\_2016\_0010 in accordance with
  - Art. 40 of Directive 2001/83/ECtransposed in the following national legislation:  
Sect 13 para 1 and sect 72 Arzneimittelgesetz (German Drug Law)

From the knowledge gained during the inspection of this manufacturer, the latest of which was conducted on 24 January 2019, it is considered that it complies with the Good Manufacturing Practice requirements referred to in

- the principles and guidelines of Good Manufacturing Practice laid down in
  - Directive 2003/94/EC

Dieses Zertifikat bestätigt den Status der Betriebsstätte zum Zeitpunkt der oben genannten Inspektion. Es sollte nicht zur Bestätigung der Übereinstimmung herangezogen werden, wenn seit der genannten Inspektion mehr als drei Jahre vergangen sind. Nach Ablauf dieser Zeit sollte mit der zuständigen Behörde Kontakt aufgenommen werden. Das Zertifikat ist nur bei Vorlage sämtlicher Seiten inklusive der Teile 1 und 2 gültig. Die Echtheit dieses Zertifikates kann ggf. durch die ausstellende Behörde bestätigt werden.

This certificate reflects the status of the manufacturing site at the time of the inspection noted above and should not be relied upon to reflect the compliance status if more than three years have elapsed since the date of that inspection, after which time the issuing authority should be consulted. This certificate is valid only when presented with all pages and both parts 1 and 2. The authenticity of this certificate may be verified with the issuing authority.





## Teil 2

## Part 2

• Humanarzneimittel

• Human Medicinal Products

### 1 HERSTELLUNGSTÄTIGKEITEN

### 1 MANUFACTURING OPERATIONS

#### 1.1 Sterile Produkte

#### 1.1 Sterile Products

1.1.2 *Im Endbehältnis sterilisiert  
(Herstellungstätigkeiten für folgende  
Darreichungsformen)*

1.1.2 *Terminally sterilised (processing operations  
for the following dosage forms)*

1.1.2.5 Andere endsterilisierte Produkte  
Alkoholtupfer

1.1.2.5 Other terminally sterilised  
prepared products  
alcoholic pads

1.1.3 *Chargenfreigabe*

1.1.3 *Batch certification*

28. September 2020

28 September 2020



Name und Unterschrift des Bearbeiters der zuständigen  
Behörde

Name and signature of the authorised person of the  
Competent Authority

Klaus Hartmann  
Landesdirektion Sachsen  
Referat 26 Pharmazie, GMP-Inspektorat  
Braustraße 2  
04107 Leipzig  
Deutschland

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## To whom it may concern

The product line of our swab dispenser is ruled differently according to the claim which is posted on the product. In Germany both is possible.

It depends on the claim you choose. If the claim is for **disinfecting** the skin before an injection, etc, it is ruled by the **AMG** (German Drug Law). If we sell Quickpad just for **cleaning** the skin Quickpad is ruled as a **cosmetic** (for example like make up remover).

We produce and sell „Quickpad“ in Germany under the regulations of the German Drug Law (Arzneimittelgesetz **AMG**).

The European market and the European regulations (i.e. Guideline-for medical items-93/42/EWG), are converted in Germany into the Medizinproduktegesetz (**MPG**).

### **There is a difference between these two regulations.**

The **AMG** (strictly national) covers all products which have a pharmaceutical effect.

The **MPG** covers all products which are not drugs but support drugs or have a **physical** (not a pharmaceutical) effect, like for example our tourniquet. The MPG fulfils the guideline 93/42/EWG. MPG, items have to bear the CE signet.

As mentioned before it depends on the claim you choose. If the claim is for disinfecting the skin before an injection, etc., it is a drug and ruled by the AMG. For this reason we have the permission to produce and to market this product under AMG ( CE-signet is not possible in this case). The product has to be labeled strictly with the original HOLTSCHE label

If your claim is for **cleaning** the skin, it is considered as a cosmetic item and is ruled by the cosmetic act and has no CE signet and no special permission as per the AMG is necessary. This is also possible in Germany.

**Most of our customers do not declare Quickpad as a drug because they might run through some kind of registration with their ministry of health. They prefer to use Quickpad like a cosmetic and as explained above there is no possibility of a CE-signet on the product.**

**Maybe the regulations in other countries are different.**

**If so, please let us know.**

Taunusstein, September 07, 2021

  
Malte Hertzberg  
Qualified Person AMG

**Safety Data Sheet**

according to Regulation (EC) No 1907/2006

**Quickpads**

Revision date: 03.07.2020

Product code: 10692-0001

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**SECTION 1: Identification of the substance/mixture and of the company/undertaking****1.1. Product identifier**

Quickpads

**1.2. Relevant identified uses of the substance or mixture and uses advised against****Use of the substance/mixture**

Disposable alcohol swabs for cleansing the skin

**1.3. Details of the supplier of the safety data sheet**

Company name:	Holtsch Medizinprodukte GmbH	
Street:	In den Faltern 13	
Place:	D-65232 Taunusstein	
Telephone:	+49(0)6128 917 177	Telefax: +49(0)6128 447 42
e-mail:	info@holtsch-med.com	
Contact person:	Malte Hertzberg	
Internet:	www.holtsch-med.com	
Responsible Department:	Responsible for the safety data sheet: sds@gbk-ingelheim.de	

**1.4. Emergency telephone number:**

+49 (0) 172 6123 572  
 In England and Wales: NHS 111 In Scotland: NHS 24 - dial 111

**SECTION 2: Hazards identification****2.1. Classification of the substance or mixture****Regulation (EC) No. 1272/2008**

Hazard categories:

Flammable liquid: Flam. Liq. 2

Serious eye damage/eye irritation: Eye Irrit. 2

Specific target organ toxicity - single exposure: STOT SE 3

Hazard Statements:

Highly flammable liquid and vapour.

Causes serious eye irritation.

May cause drowsiness or dizziness.

**2.2. Label elements****Regulation (EC) No. 1272/2008****Signal word:** Danger**Pictograms:****Hazard statements**

H225 Highly flammable liquid and vapour.

H319 Causes serious eye irritation.

H336 May cause drowsiness or dizziness.

**Additional advice on labelling**

According to the EC hazardous substance regulations in compliance with the EC cosmetic product regulations this product must not be labelled as a hazardous substance.

**2.3. Other hazards**

According to Regulation (EC) No 1907/2006 (REACH) none of the substances, contained in this product are a PBT / vPvB substance.

If swallowed in higher quantities risk of intestinal passage obstruction.

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**SECTION 3: Composition/information on ingredients****3.2. Mixtures****Chemical characterization**

Disposable swab impregnated with 70%-Isopropanol (0,46 ml per swab).

**Hazardous components**

CAS No	Chemical name			Quantity
	EC No	Index No	REACH No	
	GHS Classification			
67-63-0	Propan-2-ol			0,46 ml %
	200-661-7	603-117-00-0	01-2119457558-25	
	Flam. Liq. 2, Eye Irrit. 2, STOT SE 3; H225 H319 H336			

Full text of H and EUH statements: see section 16.

**SECTION 4: First aid measures****4.1. Description of first aid measures****General information**

No specific precautions required.

**After inhalation**

Move to fresh air in case of accidental inhalation of vapours or decomposition products.  
In the event of symptoms refer for medical treatment.

**After contact with skin**

No specific precautions required.

**After contact with eyes**

Rinse immediately with plenty of water, also under the eyelids.  
If eye irritation persists, consult a specialist.

**After ingestion**

Consult a physician.

**4.2. Most important symptoms and effects, both acute and delayed**

Causes serious eye irritation.  
May cause drowsiness or dizziness.  
If swallowed in higher quantities risk of intestinal passage obstruction.

**4.3. Indication of any immediate medical attention and special treatment needed**

Treat symptoms.

**SECTION 5: Firefighting measures****5.1. Extinguishing media****Suitable extinguishing media**Alcohol-resistant foam, dry chemical, carbon dioxide (CO<sub>2</sub>), water-spray.**Unsuitable extinguishing media**

Full water jet

**5.2. Special hazards arising from the substance or mixture**

Fire may produce:  
carbon monoxide and carbon dioxide

**5.3. Advice for firefighters**

Use breathing apparatus with independent air supply.  
Protective suit.



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### Additional information

Vapours are heavier than air and spread along ground.  
The vapour/air mixture is explosive, even in empty, uncleaned receptacles.  
Cool containers at risk with water spray jet.  
Fire residues and contaminated firefighting water must be disposed of in accordance with the local regulations.

## SECTION 6: Accidental release measures

### 6.1. Personal precautions, protective equipment and emergency procedures

In case of vapour formation use respirator.  
Ensure adequate ventilation.  
Use personal protective clothing.  
Keep away sources of ignition.  
Avoid contact with skin, eyes and clothing.

### 6.2. Environmental precautions

Do not discharge into the drains/surface waters/ground water.

### 6.3. Methods and material for containment and cleaning up

Take up mechanically and collect in suitable container for disposal.

### 6.4. Reference to other sections

Observe protective instructions (see Sections 7 and 8).  
Information for disposal see section 13.

## SECTION 7: Handling and storage

### 7.1. Precautions for safe handling

#### Advice on safe handling

When using do not eat, drink or smoke.  
Avoid contact with eyes.

#### Advice on protection against fire and explosion

Keep away from heat and sources of ignition.  
Don't smoke.

### 7.2. Conditions for safe storage, including any incompatibilities

#### Requirements for storage rooms and vessels

Protect against direct sun radiation.  
Keep containers tightly closed in a cool, well-ventilated place.

#### Hints on joint storage

Incompatible with:  
Oxidizing agents  
Alkaline metals and earth alkaline metals.

#### Further information on storage conditions

Keep away from food, drink and animal feeding stuffs.

### 7.3. Specific end use(s)

Disposable alcohol swabs for cleansing the skin

## SECTION 8: Exposure controls/personal protection

### 8.1. Control parameters

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**Exposure limits (EH40)**

CAS No	Substance	ppm	mg/m <sup>3</sup>	fibres/ml	Category	Origin
67-63-0	Propan-2-ol	400	999		TWA (8 h)	WEL
		500	1250		STEL (15 min)	WEL

**8.2. Exposure controls****Appropriate engineering controls**

Ensure adequate ventilation, especially in confined areas.

**Protective and hygiene measures**

When using do not eat, drink or smoke.

Do not inhale vapours.

Avoid contact with eyes.

**Eye/face protection**

Not required under normal use.

**Hand protection**

Not required under normal use.

**Skin protection**

Not required under normal use.

**Respiratory protection**

No personal respiratory protective equipment normally required.

**SECTION 9: Physical and chemical properties****9.1. Information on basic physical and chemical properties**

Physical state: Tissue, tintured with clear disinfectant solution

Colour:

Odour: Alcoholic

**Test method**

pH-Value: n.d.

**Changes in the physical state**

Melting point: n.d.

Initial boiling point and boiling range: n.d.

Sublimation point: n.a.

Softening point: n.d.

Flash point: 12 °C \*)

**Flammability**

Solid: Combustible.

Gas: n.a.

**Explosive properties**

The product is considered non-explosive; nevertheless explosive vapour/air mixture can be generated.

Lower explosion limits: 2,0 vol. % \*)

Upper explosion limits: 12 vol. % \*)

Ignition temperature: 425 °C \*)

**Auto-ignition temperature**

Solid: The product is not self-igniting

Gas: n.a.

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Decomposition temperature:	n.d.
<b>Oxidizing properties</b>	
Not oxidising.	
Vapour pressure: (at 20 °C)	43 hPa *)
Density (at 20 °C):	0,87 g/cm <sup>3</sup> *)
Bulk density:	n.d.
Water solubility: (at 20 °C)	Miscible *)
<b>Solubility in other solvents</b>	
n.d.	
Viscosity / dynamic:	n.a.
Viscosity / kinematic:	n.a.
Flow time:	n.a.
Vapour density:	n.d.
Evaporation rate:	n.d.
Solvent separation test:	n.d.
Solvent content:	n.d.

**9.2. Other information**

\*) All data refer to the solution

**SECTION 10: Stability and reactivity****10.1. Reactivity**

No decomposition if stored and applied as directed.

**10.2. Chemical stability**

Stable under normal conditions.

**10.3. Possibility of hazardous reactions**

No hazardous reactions known.

**10.4. Conditions to avoid**

Heating can release vapours which can be ignited.

**10.5. Incompatible materials**Oxidizing agents  
Alkaline metals and earth alkaline metals.**10.6. Hazardous decomposition products**No hazardous decomposition products known.  
Fire may produce:  
Carbon monoxide and carbon dioxide**SECTION 11: Toxicological information****11.1. Information on toxicological effects****Acute toxicity**

Based on available data, the classification criteria are not met.

Propan-2-ol

LD50/oral/rat: 5045 mg/kg [RTECS]

LD50/dermal/rabbit: 12800 mg/kg

LC50/inhalation/rat: 46,5 mg/l/4h [RTECS]

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### **Irritation and corrosivity**

Causes serious eye irritation.

Skin corrosion/irritation: Based on available data, the classification criteria are not met.

### **Sensitising effects**

Based on available data, the classification criteria are not met.

### **Carcinogenic/mutagenic/toxic effects for reproduction**

Based on available data, the classification criteria are not met.

### **STOT-single exposure**

May cause drowsiness or dizziness.

### **STOT-repeated exposure**

Based on available data, the classification criteria are not met.

### **Aspiration hazard**

Based on available data, the classification criteria are not met.

### **Additional information on tests**

Classification in compliance with the assessment procedure specified in the Regulation (EC) no 1272/2008.

### **Practical experience**

#### **Other observations**

If appropriately handled and if in accordance with the general hygienic rules, no damages to health have become known.

Sometimes short-term reddening of skin may occur.

May cause irritation of the mucous membranes.

If swallowed in higher quantities risk of intestinal passage obstruction.

## SECTION 12: Ecological information

### **12.1. Toxicity**

Propan-2-ol

LC50/Lepomis macrochirus/96 h = 1400 mg/l [ECOTOX DATABASE]

EC50/Daphnia magna/48 h = 13299 mg/l [IUCLID]

EC50/Desmodesmus subspicatus/72 h > 1000 mg/l [IUCLID]

### **12.2. Persistence and degradability**

Propan-2-ol

Biodegradable (OECD): 95% [OECD 301 E]

### **12.3. Bioaccumulative potential**

Propan-2-ol

Product has a low bioaccumulating potential. Log Pow: 0,05 [OECD 107]

### **12.4. Mobility in soil**

No data available

### **12.5. Results of PBT and vPvB assessment**

According to Regulation (EC) No 1907/2006 (REACH) none of the substances, contained in this product are a PBT / vPvB substance.

### **12.6. Other adverse effects**

Low hazard to waters.

### **Further information**

Product is not allowed to be discharged into aquatic environment.

## SECTION 13: Disposal considerations

### **13.1. Waste treatment methods**

#### **Disposal recommendations**

Can be incinerated, when in compliance with local regulations.



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### List of Wastes Code - residues/unused products

180106 WASTES FROM HUMAN OR ANIMAL HEALTH CARE AND/OR RELATED RESEARCH (EXCEPT KITCHEN AND RESTAURANT WASTES NOT ARISING FROM IMMEDIATE HEALTH CARE); wastes from natal care, diagnosis, treatment or prevention of disease in humans; chemicals consisting of or containing hazardous substances; hazardous waste

### Contaminated packaging

Empty containers should be taken for local recycling, recovery or waste disposal.

Contaminated packaging should be emptied as far as possible and after appropriate cleansing may be taken for reuse.

Packaging that cannot be cleaned should be disposed of like the product.

## SECTION 14: Transport information

### Land transport (ADR/RID)

**14.1. UN number:** UN 3175  
**14.2. UN proper shipping name:** SOLIDS CONTAINING FLAMMABLE LIQUID, N.O.S. (Propan-2-ol)  
**14.3. Transport hazard class(es):** 4.1  
**14.4. Packing group:** II  
 Hazard label: 4.1



Classification code: F1  
 Limited quantity: 1 kg / 30 kg  
 Excepted quantity: E2  
 Transport category: 2  
 Hazard No: 40  
 Tunnel restriction code: E

### Inland waterways transport (ADN)

**14.1. UN number:** UN 3175  
**14.2. UN proper shipping name:** SOLIDS CONTAINING FLAMMABLE LIQUID, N.O.S. (Propan-2-ol)  
**14.3. Transport hazard class(es):** 4.1  
**14.4. Packing group:** II  
 Hazard label: 4.1



Classification code: F1  
 Limited quantity: 1 kg / 30 kg  
 Excepted quantity: E2

### Marine transport (IMDG)

**14.1. UN number:** UN 3175  
**14.2. UN proper shipping name:** SOLIDS CONTAINING FLAMMABLE LIQUID, N.O.S. (Propan-2-ol)  
**14.3. Transport hazard class(es):** 4.1  
**14.4. Packing group:** II  
 Hazard label: 4.1

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Marine pollutant:	No
Limited quantity:	1 kg / 30 kg
Excepted quantity:	E2
EmS:	F-A, S-I

**Air transport (ICAO-TI/IATA-DGR)**

<b>14.1. UN number:</b>	UN 3175
<b>14.2. UN proper shipping name:</b>	SOLIDS CONTAINING FLAMMABLE LIQUID, N.O.S. (Propan-2-ol)
<b>14.3. Transport hazard class(es):</b>	4.1
<b>14.4. Packing group:</b>	II
Hazard label:	4.1



Limited quantity Passenger:	5 kg
Passenger LQ:	Y441
Excepted quantity:	E2
IATA-packing instructions - Passenger:	445
IATA-max. quantity - Passenger:	15 kg
IATA-packing instructions - Cargo:	448
IATA-max. quantity - Cargo:	50 kg

**14.5. Environmental hazards**

ENVIRONMENTALLY HAZARDOUS: no

**14.6. Special precautions for user**

Take the usual precautions when handling with chemicals.

**14.7. Transport in bulk according to Annex II of Marpol and the IBC Code**

The transport takes place only in approved and appropriate packaging.

**SECTION 15: Regulatory information****15.1. Safety, health and environmental regulations/legislation specific for the substance or mixture****EU regulatory information**

2004/42/EC (VOC):	0,46 ml
Information according to 2012/18/EU (SEVESO III):	P5c FLAMMABLE LIQUIDS

**National regulatory information**

Employment restrictions:	Observe restrictions to employment for juveniles according to the 'juvenile work protection guideline' (94/33/EC). Observe employment restrictions under the Maternity Protection Directive (92/85/EEC) for expectant or nursing mothers.
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**15.2. Chemical safety assessment**

For this substance a chemical safety assessment has not been carried out.

**SECTION 16: Other information**

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**Abbreviations and acronyms**

ADR = Accord européen relatif au transport international des marchandises Dangereuses par Route  
 RID = Règlement concernant le transport international ferroviaire de marchandises dangereuses  
 ADN = Accord européen relatif au transport international des marchandises dangereuses par voie de navigation int er  
 IMDG = International Maritime Code for Dangerous Goods  
 IATA/ICAO = International Air Transport Association / International Civil Aviation Organization  
 MARPOL = International Convention for the Prevention of Pollution from Ships  
 IBC-Code = International Code for the Construction and Equipment of Ships Carrying Dangerous Chemicals in Bulk  
 GHS = Globally Harmonized System of Classification and Labelling of Chemicals  
 REACH = Registration, Evaluation, Authorization and Restriction of Chemicals  
 CAS = Chemical Abstract Service  
 EN = European norm  
 ISO = International Organization for Standardization  
 DIN = Deutsche Industrie Norm  
 PBT = Persistent Bioaccumulative and Toxic  
 vPvB = Very Persistent and very Bio-accumulative  
 LD = Lethal dose  
 LC = Lethal concentration  
 EC = Effect concentration  
 IC = Median immobilisation concentration or median inhibitory concentration

**Relevant H and EUH statements (number and full text)**

H225 Highly flammable liquid and vapour.  
 H319 Causes serious eye irritation.  
 H336 May cause drowsiness or dizziness.

**Further Information**

Data of items 4 to 8, as well as 10 to 12, do partly not refer to the use and the regular employing of the product (in this sense consult information on use and on product), but to liberation of major amounts in case of accidents and irregularities.

The information describes exclusively the safety requirements for the product(s) and is based on the present level of our knowledge.

The delivery specifications are contained in the corresponding product sheet.

This data does not constitute a guarantee for the characteristics of the product(s) as defined by the legal warranty regulations.

(n.a. = not applicable; n.d. = not determined)

*'The data for the hazardous ingredients were taken respectively from the last version of the sub-contractor's safety data sheet.)*

HOLTSCH MED

Quality. Safety. Trust.

## Quickpad

The Quickpad alcohol swab dispenser, being sterile and physiologically verified as harmless, is ideal for cleaning and disinfecting the skin.

The active agent 2 propanol acts as an effective, "mild on skin" disinfecting agent. The "swab tear-off" ready to use system allows for the single use of the swab. This makes Quickpad not only economical but also efficient in its use.

The lid of the Quickpad alcohol swab dispenser container seals air-tight, keeping the alcohol swabs moist and sterile. As a result, the swab dispenser has a particularly long shelf life of 24 months.

Quickpad is supplied ready for use and with its simple operation, can be used by specialists as well as patients for cleaning/disinfecting the skin. An ideal product for diabetics and other users of self-injection syringes. Additionally, it is always possible through the transparent container to see the fill level and it is suitable for self-application by the patient.

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**Quickpad**

Wirkstoff: 2-Propanol (Ph. Eur.) 70% (V/V),  
Getränke-Typ für Anwendung auf der Haut  
für Säuglinge, Kinder und Erwachsene

**Stoff- oder Indikationsgruppe:** Desinfektionsmittel / Antiseptikum  
**Anwendungsgebiete:** Zur Desinfektion der Haut vor Injektionen  
**Gegenanzeigen:** Nicht anwenden bei Überempfindlichkeit gegen 2-Propanol (Isopropylalkohol) • zur Hautdesinfektion vor Eingriffen, die eine größere Eröffnung erfordern als eine Injektion • zur Antiseptik offener Wunden, • zur Schweißhautentzündung, insbesondere am Auge • zur Hautdesinfektion bei Fräugeborenen  
**Warnhinweise und Vorsichtsmaßnahmen für die Anwendung:** Keine Anwendung auf Schleimhäuten, insbesondere dem Auge. Waschen Sie in Schwangerschaft und Stillzeit beachten! Bei sachgemäßem Gebrauch kann QUICKPAD® während der Schwangerschaft und Stillzeit verwendet werden. Wechselwirkungen mit anderen Mitteln: Keine bekannt.  
**Wirkungen und Art der Anwendung:** Zur Desinfektion der betroffenen Hautfläche mit QUICKPAD® streichen und trocknen lassen. Als Munddesinfektionsmittel sind 15 s einzuhalten. Dauer der Anwendung: QUICKPAD® kann ohne zeitliche Begrenzung angewendet werden.  
**Nebenwirkungen:** Bei Hautirritationen mit 2-Propanol (Ph. Eur.) 70% (V/V) können in

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