CLH report

Proposal for Harmonised Classification and Labelling

Based on Regulation (EC) No 1272/2008 (CLP Regulation), Annex VI, Part 2

International Chemical Identification:

2,2'-[[3-methyl-4-[(4-nitrophenyl)azo]phenyl]imino]bisethanol

EC Number:	221-665-5
CAS Number:	3179-89-3
Index Number:	-

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1 IDENTITY OF THE SUBSTANCE

1.1 Name and other identifiers of the substance

Table 1: Substance identity and information related to molecular and structural formula of the substance

Name(s) in the IUPAC nomenclature or other international chemical name(s)	2,2'-[[3-methyl-4-[(E)-(4-nitrophenyl)azo]phe- nyl]imino]diethanol
Other names (usual name, trade name, abbreviation)	C.I. 11210, Disperse Red 17
ISO common name (if available and appropriate)	
EC number (if available and appropriate)	221-665-5
EC name (if available and appropriate)	2,2'-[[3-methyl-4-[(4-nitrophenyl)azo]phenyl]imino]bi- sethanol
CAS number (if available)	3179-89-3
Other identity code (if available)	
Molecular formula	$C_{17}H_{20}N_4O_4$
Structural formula	
SMILES notation (if available)	Cc1cc(ccc1N=Nc2ccc(cc2)[N+]([O-])=O)N(CCO)CCO
Molecular weight or molecular weight range	344.37
Information on optical activity and typical ratio of (stereo) isomers (if applicable and appropriate)	-
Description of the manufacturing process and identity of the source (for UVCB substances only)	-
Degree of purity (%) (if relevant for the entry in Annex VI)	-

1.2 Composition of the substance

Table 2: Constituents (non-confidential information)

Constituent (Name and numeri- cal identifier)	Concentration range (% w/w mini- mum and maximum in multi-con- stituent substances)	Current CLH in Annex VI Table 3 (CLP)	Current self- classification and labelling (CLP)
2,2'-[[3-methyl-4- [(E)-(4-nitro- phenyl)azo]phe- nyl]imino]diethanol CAS: 3179-89-3 EC: 221-665-5	-	-	Acute Tox. 4; H301 (34) Skin Sens. 1; H317 (3) STOT RE 2; H373 (30) Aquatic Chronic 2; H411 (42)

Table 3: Impurities (non-confidential information) if relevant for the classification of the substance

Impurity (Name and numeri- cal identifier)	Concentration range (% w/w minimum and maximum)	Current CLH in Annex VI Table 3.1 (CLP)	Current self- classi- fication and label- ling (CLP)	The impurity con- tributes to the clas- sification and label- ling
-				

Table 4: Additives (non-confidential information) if relevant for the classification of the substance

Additive (Name and nu- merical identi- fier)	Function	Concentration range (% w/w mini- mum and maxi- mum)	Current CLH in Annex VI Table 3.1 (CLP)	Current self- classification and labelling (CLP)	The additive contributes to the classifica- tion and label- ling
-					

Table 5: Test substances (non-confidential information) (this table is optional)

Identification of test sub- stance	Purity	Impurities and additives (iden- tity, %, classification if availa- ble)	Other information	The study(ies) in which the test sub- stance is used
-				

2 PROPOSED HARMONISED CLASSIFICATION AND LABELLING

2.1 Proposed harmonised classification and labelling according to the CLP criteria

Table 6: Proposed harmonised classification and labelling according to the CLP criteria

					Classificat	tion		Labelling			
	Index No	International Chemical Identifi- cation	EC No	CAS No	Hazard Class and Category Code(s)	Hazard statement Code(s)	Pictogram, Signal Word Code(s)	Hazard statement Code(s)	Suppl. Haz- ard state- ment Code(s)	Specific Conc. Limits, M-factors	Notes
Current An- nex VI entry	No existing entry in Annex VI of CLP										
Dossier sub- mitters pro- posal	tba	2,2'-[[3-methyl-4-[(4- nitrophenyl)azo]phe- nyl]imino]bisethanol	221-665-5	3179-89-3	Skin Sens 1	H317	GHS07 Wng	H317			
Resulting An- nex VI entry if agreed by RAC and COM	tba	2,2'-[[3-methyl-4-[(4- nitrophenyl)azo]phe- nyl]imino]bisethanol	221-665-5	3179-89-3	Skin Sens 1	H317	GHS07 Wng	H317			

Hazard class	Reason for no classification	Within the scope of public consultation		
Explosives				
Flammable gases (including chemically unstable gases)				
Oxidising gases				
Gases under pressure				
Flammable liquids				
Flammable solids				
Self-reactive substances				
Pyrophoric liquids				
Pyrophoric solids				
Self-heating substances				
Substances which in contact with water emit flammable gases	Not assessed in this dossier	No		
Oxidising liquids				
Oxidising solids				
Organic peroxides				
Corrosive to metals				
Acute toxicity via oral route				
Acute toxicity via dermal route				
Acute toxicity via inhalation route				
Skin corrosion/irritation				
Serious eye damage/eye irrita- tion				
Respiratory sensitisation				
Skin sensitisation	Harmonised classification proposed	Yes		
Germ cell mutagenicity				
Carcinogenicity				
Reproductive toxicity				
Specific target organ toxicity- single exposure				
Specific target organ toxicity- repeated exposure	Not assessed in this dossier	No		
Aspiration hazard				
Hazardous to the aquatic envi- ronment				
Hazardous to the ozone layer				

3 HISTORY OF THE PREVIOUS CLASSIFICATION AND LABELLING

Disperse Red 17 is listed as a pre-registered substance under REACH (substances indicated, in 2009, as being intended to be registered by at least one company in the EEA). It does not have a harmonised classification and labelling in Annex VI to the CLP regulation (ECHA, 2020).

Furthermore, Disperse Red 17 is a substance likely to meet the criteria of Annex III to the REACH Regulation, based on an analysis of publicly available databases with experimental data and by using (Q)SAR model results. According to this analysis, Disperse Red 17 is indicated as "suspected carcinogen", "suspected mutagen", "suspected persistent in the environment", and "suspected toxic for reproduction".

4 JUSTIFICATION THAT ACTION IS NEEDED AT COMMUNITY LEVEL

There is evidence from the literature that Disperse Red 17 elicits skin sensitisation in human as shown in studies from a high number of dermatological clinics. Harmonised classification is proposed because there are differences between the self-classifications notified in the classification and labelling (C&L) inventory (ECHA (2014), section 4.1.1). Furthermore, the DS disagrees with most of the current self-classifications. In fact, the vast majority of the notifiers did not self-classify Disperse Red 17 as a skins sensitiser (Table 8). Harmonised classification of Disperse Red 17 would ensure an adequate perception of the skin sensitisation hazard by setting the concentration limit for the classification as Skin Sens. 1 could improve consumer safety in the context of restriction proposals on the use of the substance referring to harmonised classifications as skin sensitiser. In fact, Disperse Red 17 is listed on the restriction proposal for the placing on the market of textile, leather, hide and fur articles containing skin sensitising substances (ECHA, 2019b) and the restriction proposal for substances in tattoo inks and permanent make up (ECHA, 2019a).

Hazard Class and Cate- gory Code(s)	Hazard Statement Code(s)	Pictograms, Signal Word Code(s)	Number of Noti- fiers	
Acute Tox. 4	H302	GHS07 Wng	48	
STOT RE 2	H373 (blood)	GHS09		
Aquatic Chronic 2	H411	GHS08 Wng	36	
Aquatic Chronic 2	H411	GHS09 Wng	21	
Skin Sens. 1	H317	GHS07 Wng	3	
Not Classified	-	-	3	
Acute Tox. 4	H302	GHS09		
STOT RE 2	H373 (blood)	GHS08 GHS07	1	
Aquatic Chronic 2	H411	Wng		

Table 8: Notified classification and labelling according to CLP criteria (ECHA, 2020)

4.1 Identified uses

Based on data from a literature research (sources: cf. section 6), Disperse Red 17 is used to dye fabrics made of synthetic fibres such as polyester fibres (Clauss and Weiss, 1992; Svedman et al., 2019; Varma et al., 1980; Veena et al., 1979). These fibres are used in turn to produce garments that are mostly worn directly on the skin (Hausen and Schulz, 1984; Suter, 1965). Balato and colleagues reported Disperse Red 17 to be among the most frequently isolated dyes in 51 stockings and panty hoses investigated (Balato et al., 1990a). Based on data from newer studies available to the DS, analysing a limited number of textiles from a very large market, Disperse

Red 17 was not detected in the analysed fabrics (BVL, 2010; Malinauskiene et al., 2012; Wu et al., 2019; Zhou et al., 2014). However, with respect to the frequency of positive patch test reactions to Disperse Red 17 in humans, as shown in several clinical settings from newer investigations, it cannot be excluded that Disperse Red 17 is still used in dyeing processes for clothes or in other application areas (Foley et al., 2019; Heratizadeh et al., 2017; Isaksson et al., 2015; Ortiz-Salvador et al., 2017; Toholka et al., 2015). The ÖkoTex Standard 100 lists Disperse Red 17 as an allergenic dye, defining a limit value in textiles produced according to this Standard (OEKO-TEX, 2020). For labelling of textiles with the EU Ecolabel, Disperse Red 17 shall not be used for dyeing polyester, acrylic, polyamide, elasticated or stretchable skin contact garments, or underwear (EU Ecolabel, 2015).

Disperse Red 17 was found as ingredient of haircare products (Katugampola and Statham, 2005). It is used as a non-reactive hair colouring agent in oxidative hair dye formulations (maximum on-head concentration of 2 %, including dispersant) and as a non-reactive hair colouring agent (direct dye) in semi-permanent hair dye formulations (maximum on-head concentration of 0.2 %, including dispersant, SCCS (2013)). Furthermore, Disperse Red 17 sensitisation was identified in patients that had had a temporary "black henna tattoo" (de Groot, 2013; Kind et al., 2012; Le Coz and Tromp, 2002; Moro et al., 2016; Saunders et al., 2004). Disperse Red 17 is suspected to be used as colorant in tattoo inks and it may also be used for dyeing spectacle frames (Walsh and Wilkinson, 2006).

5 DATA SOURCES

Data for Disperse Red 17 were obtained from the public ECHA dissemination site and from a thorough search of the published literature in bibliographic databases (Web of Science, Embase, PubMed, Scopus, Wiley Online library, and Google Scholar). Furthermore, data were taken from the Scientific Committee on Consumer Safety (SCCS) Opinion on Disperse Red 17 (SCCS, 2013).

6 PHYSICOCHEMICAL PROPERTIES

Property	Value	Reference	Comment (e.g. measured or estimated)
Physical state at 20°C and 101,3 kPa	No value available		
Melting/freezing point	No value available		
Boiling point	No value available		
Relative density	No value available		
Vapour pressure	No value available		
Surface tension	No value available		
Water solubility	No value available		
Partition coefficient n-oc- tanol/water	No value available		
Flash point	No value available		
Flammability			
Explosive properties			
Self-ignition temperature			
Oxidising properties			
Granulometry	No value available		

 Table 9: Summary of physicochemical properties

Property	Value	Reference	Comment (e.g. measured or estimated)
Stability in organic solvents and identity of relevant degra- dation products	No value available		
Dissociation constant	No value available		
Viscosity	No value available		

7 EVALUATION OF PHYSICAL HAZARDS

Not assessed in this dossier

8 TOXICOKINETICS (ABSORPTION, METABOLISM, DISTRIBUTION AND ELIMI-NATION)

Table 10: Summary table of toxicokinetic studies

Method	Test sub- stance	Results		Reference
OECD TG 428 GLP compliance Reliability 4: Not assignable Adopted from (SCCS, 2013) Human (female) dermatomed abdominal skin (3 different donors, 9 skin prepara- tions) Test item: formulation with 0.2 % (w/w) test substance (non-oxidative condi- tions), formulation with 2.0 % (w/w) test substance (oxidative conditions); Doses: non-oxidative: 16.0 mg (25 mg/cm ²), oxidative: 13.1 mg (20 mg/cm ²) Exposure: 60 min Method of Analysis: HPLC	Disperse Red 17 Dye content: 31 % Batch No.: 40T60N4520	Mean total absorption (= amount present in the receptor fluid, receptor compart- ment wash and the skin, excluding tape strips): 0.41 µg/cm ² (0.89 % of the applied dose, non-oxidative conditions), 0.50 µg/cm ² (0.11 % of the applied dose, oxidative conditions)		(SCCS, 2013)
In vitro absorption through human and pig epidermis No guideline study GLP: No information Reliability 3: Not reliable Disperse Red 17, applied at a rate of 200 μ /cm ² (200 μ g/cm ²), Prepared suspensions, 1000 μ g/ml in TWEEN 80 (0.5 % in distilled water), Epidermis was prepared from whole skin samples of both species and mounted on glass diffusion cells; Absorption was measured under occlu- sion, Exposure: 55 h Samples from receptor chamber were an- alysed by HPLC	Disperse Red 17 Purity: No in- formation	Time period (h) Human epidermis 1-10 4-30 Pig epidermis 1-10 31-55	$\begin{array}{c} 0.07 \pm 0.01 \\ \hline 0.11 \pm 0.01 \\ \hline 0.16 \pm 0.01 \\ \hline 1.80 \pm 0.28 \\ \hline \text{vas absorbed by human} \end{array}$	(ZENECA, 1997)

Toxicological data giving adequate information on the absorption, distribution, biotransformation, and excretion of Disperse Red 17 are lacking. However, two studies available to the DS investigated the absorption of Disperse Red 17 through human epidermis. The absorption of Disperse Red 17, showing a dye content of 31 %, was analysed in a study performed according to OECD TG 428 with GLP compliance. This study was not available to the DS, but the study summary was taken from the Scientific Committee and Consumer Safety's opinion on Disperse Red 17 (SCCS, 2013). The authors applied the dye to human dermatomed abdominal skin mounted in flow-through diffusion cells. Disperse Red 17 was tested in two formulations under non-oxidative and oxidative conditions, representing in use-conditions as semi-permanent hair dye. The mean total absorption (amount present in the receptor fluid, receptor compartment wash and the skin, excluding tape strips) was 0.41 μ g/cm² (0.89 % of the applied dose) under non-oxidative conditions and 0.50 μ g/cm² (0.11 % of the applied dose) under oxidative conditions.

In a second non-guideline study (ZENECA, 1997), the epidermis was prepared from human whole skin samples and pig whole skin (prepared from the cartilage of pig ears) and mounted in glass diffusion cells. Disperse Red 17 (no information on purity) was applied with a solution of 0.5% TWEEN 80 in distilled water to give a dye concentration of 1 000 μ g/mL using an application rate of 200 μ L/cm (corresponding to 200 μ g/mL). Receptor fluids were taken at certain intervals over an exposure period of 55 hours. Fluids were analysed for dye concentrations using HPLC. Mean absorption rates for Disperse Red 17 were 0.07 ± 0.01 μ g/cm² per hour analysed from samples taken one to 10 hours after dye application, and 0.11 ± 0.01 μ g/cm² per hour analysed from samples taken four to 30 hours after dye application. The authors concluded that Disperse Red 17 is absorbed by human skin with a mean absorption rate of 0.1 μ g/cm²/hr, which corresponds to 0.2 % of the applied dose.

8.1 Short summary and overall relevance of the provided toxicokinetic information on the proposed classification(s)

Two studies investigated the absorption of Disperse Red 17 through human skin *in vitro*. Studies were of low reliability or not assignable to the DS. Mean absorption rates of Disperse Red 17 through human skin were between 0.1 % (oxidative conditions) and 0.9 % (non-oxidative conditions) of the applied dose.

9 EVALUATION OF HEALTH HAZARDS

9.1 Acute toxicity - oral route

Not assessed in this dossier

9.2 Acute toxicity - dermal route

Not assessed in this dossier

9.3 Acute toxicity - inhalation route

Not assessed in this dossier

9.4 Skin corrosion/irritation

Not assessed in this dossier

9.5 Serious eye damage/eye irritation

Not assessed in this dossier

9.6 Respiratory sensitisation

Not assessed in this dossier

9.7 Skin sensitisation

9.7.1 Animal data

Table 11: Summary table of animal studies on skin sensitisation

Method, guideline, deviations if	Species,	Test sub-	Dose levels	Results	Reference
any	strain, sex, no/group	stance,	duration of exposure		
"Sensitive mouse lymph node as- say" (SLNA) Non-guideline study No information on GLP Study reliability 3: Not reliable <u>Deviations to OECD TG 429:</u> Intradermal injection: Day 1, adju- vant was used Topical application: Day 6-8, in- stead of monophasic application; Endpoint analysis: Day 9, instead of two days without treatment; Analysis of lymph node cell num- ber (SI _n) after excision of lymph nodes, using automated cell coun- ter; Determination of ³ HTdR incorpo- ration in lymphocytes after 24 h of cell culture (SI _p) was analysed; Individual body weights at start of dosing and at scheduled sacrifice not reported;	Mouse, BALB/c, female n=3/dose	Disperse Red 17 (CI 11210) <u>Purity:</u> No in- formation p-Phenylene- diamine (PPD) (CI 76060) <u>Purity:</u> No in- formation; purchased from Wako Pure Chemi- cal Industries, Ltd. (Osaka, Japan)	Intradermal injection: 2 % in saline/Freund's complete adjuvant (FCA) (1:1) Topical application: 10 % in DMF Results of stimulation index (SI), defined by authors: SI _n (Disperse Red 17): 0.9 SI _p (Disperse Red 17): 0.9 SI _{total} (SI _n x SI _p) = total LN response: 0.8 Vehicle control: intradermal injection of vehicle-FCA emulsion, topical applica- tion of vehicle alone PPD, intradermal injection: 0.2 % in DMSO/FCA Topical application: 10 % in DMSO SI _n (PPD): 4.1 SI _p (PPD): 7.3 SI _{total} (SI _n x SI _p): 29.6	Negative	(Ikarashi et al., 1996)
GPMT OECD TG 406 GLP-compliant Study reliability 4: Not assignable Cited from secondary reference (SCCS, 2013)	Guinea pig, Dun- kin-Hart- ley, fe- male N=10/ dose N=5/con- trol group	Disperse Red 17, dispersed in water <u>Dye content:</u> 41.2% Batch No.: 928017/02	Intradermal induction (3x): 0.1 mL 5 % (w/v) test sub- stance/FCA ¹ ; 0.1 mL 50 % FCA; 0.1 mL 5 % (w/v) test substance Day 6, induction of irrita- tion: 10 % sodium lauryl sulphate Day 8, topical induction: 0.5 mL 2.5 % test substance for 48 h (occluded) Two weeks later, challenge: 2.5 % test substance for 24 h (occluded) Excessive staining due to test substance precluded ac- curate assessment in 6/10 animals	Negative	(Karunarat ne, 1995)

Two animal studies investigated the sensitising potential of Disperse Red 17. In a non-guideline "Sensitive mouse lymph node assay (SLNA)" Disperse Red 17 of unspecified purity was intradermally injected in a 2 %

¹ FCA - Freund's Complete Adjuvant

test chemical-FCA emulsion into two sites of the abdominal skin at both sides of the ventral midline. After five days, topical application on the ears followed with 10 % test substance for three consecutive days (day 6 to 8). The following day, excised auricular lymph nodes were pooled for each experimental group. A single cell suspension of a defined number of local lymph node cells was cultured with [³H] methyl thymidine (³HTdR). After 24 hours, ³HTdR incorporation was determined using liquid scintillation counting. The increase in local lymph node cell number and ³HTdR incorporation compared to controls were expressed as stimulation index n (SI_n - calculated from local lymph node cell number after excision), and SI_p (calculated from local lymph node cell proliferation in cell culture), respectively. According to the authors, a chemical was regarded as a sensitiser, if SI_{total} (SI_n x SI_p, which indicates the total lymph node activation induced by the test chemicals) showed a value of 3 or more. SI values for Disperse Red 17 were SI_n= 0.9, SI_p= 0.9, resulting in a SI_{total} of 0.8. The authors concluded that Disperse Red 17 was not a sensitiser in this test. However, each test substance was tested without varying concentrations and concentrations higher than 2 % were not applied, "to prevent systemic toxicity" (Ikarashi et al., 1996). However, there are no further information on which (pre-) study this concentration is based on.

Furthermore, the skin sensitising potential of Disperse Red 17 was investigated in a guinea pig maximisation test according to OECD testing guidelines (OECD TG 406) and in compliance with GLP. This study was not available to the DS, but was already assessed by the Scientific Committee on Consumer Safety (SCCS). According to the SCCS report, a preliminary intradermal study showed that a concentration of 5 % test substance did not induce an irritant response. For induction in the main study, guinea pigs received three intradermal injections of Disperse Red 17 (dye content: 41.2 %), using 5 % test substance in Freund's complete adjuvant (FCA), followed by a single epidermal induction on day 8 using 2.5 % of the test material under occlusive patch for 48 hours. Two weeks after completed induction, animals were challenged by a single application of 2.5 % test substance under occlusive conditions for 24 hours. Skin examination followed 24 and 48 hours after removal of the challenge patches. In its opinion on Disperse Red 17 the SCCS stated, that "skin staining was observed due to the test substance and was reported to preclude accurate assessment of erythema after the induction and the challenge application in 6/10 animals. No adverse reaction was observed in any of the treated guinea pigs. The author concluded that the test substance was not a sensitiser to guinea pig skin" (SCCS, 2013).

Altogether, none of the two available animal studies revealed a relevant skin sensitising potential of Disperse Red 17. However, one study is of low reliability due to the insufficient characterisation of the test material and deviations from OECD test guideline procedures, e.g. by not testing a dose series of the test material. A second study comprising a GPMT was performed with Disperse Red 17 of low dye content (41.2 %). This study was not assignable to the DS. According to the SCCS opinion on Disperse Red 17 this GPMT was of low reliability as well, because "excessive staining due to test substance preclude accurate assessment in 6/10 animals".

In conclusion, none of the available animal studies is sufficiently reliable to conclude on the skin sensitising potential of Disperse Red 17.

9.7.2 Human data

Table 12: Summary table of human patch test data and published cases on skin sensitisation²

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
	Dermatitis patients (un	selected, consecutive)			
1	Retrospective analy- sis from dermatologi- cal departments to identify the most rele- vant allergens	01/2001 - 12/2010: 5 521 patients presented, of whom 5 281 were generally patch-tested with an extended European stand- ard series and additional allergens or series based on the der- matologist's assessment (806 patients were patch-tested with Disperse Red 17, 1 % in pet.). Patch test data were presented for positive and relevant reactions of test substances in the whole patch test population (N= 5281). This assumed that sub- jects not patch tested to the allergen would not have tested pos- itive and that the test substance was available for the whole 10- year patch test study. Data were calculated for the most accu- rate comparison of allergens in the whole patch test population; however, patch test results for substances not tested in all pa- tients are considerably underestimated.	Disperse Red 17: 1.5 % (77/5 281) positive reactions Disperse Red 17: 0.2 % (10/5 281) relevant reactions	Positive Low/moderate frequency of relevant reactions Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Toholka et al., 2015)
2	Patch tests from der- matological clinic	Consecutive patients with eczema (n = 327) and healthy stu- dent volunteers (n = 205, non-patient population, recruited by advertisement; confirmed or suspected textile allergy was nei- ther an inclusion nor an exclusion criterion) were patch-tested with the modified European baseline series and textile dye al- lergens (incl. Disperse Red 17, 1 % in pet.). No time window reported; self-selected volunteers, sensitisa- tion rate may be over-represented; volunteers aged 20-27 years	Disperse Red 17: Consecutive eczema patients: 0.9 % positive reactions Healthy volunteers: 2.0 % posi- tive reactions	Positive Low/moderate frequency among consecutive eczema patients Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Li, 2010)
3	Patch tests/consumer tests from two derma- tological departments	02 - 12/2005 (first department) and $08/2004 - 11/2005$ (second department): 982 dermatitis patients were consecutively patch- tested with baseline patch test series of respective departments, temporarily including a textile dye mix and its eight separate components (incl. Disperse Red 17, 0.5 % in pet.); 858 patients answered a questionnaire.	20/982 positive reactions to textile dye mix Disperse Red 17: 0.3 % (3/982) positive reactions	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Ryberg et al., 2009)

² Available patch test readings according to International Contact Dermatitis Research Group criteria: (+) weak positive (erythema, infiltration, possibly papules), (++) strong positive (erythema, infiltration, papules, vesicles), (++) extreme positive reaction (intense erythema, infiltrate, coalescing vesicles) (Johansen et al., 2015).

³ Frequency and exposure are rated as relatively high or low/moderate in line with Tables 3.2 and 3.3 of the ECHA "Guidance on the Applicability of the CLP criteria", where possible.

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
			(++) reaction in 1 patient, (+) in 2 patients		
4	Patch test from two dermatological clinics	286 consecutive patients were patch-tested over a period of one year with the TRUE Test standard series and a textile colour and finish series (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 0.3 % (1/286) positive reactions	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Lazarov and Cor- doba, 2000)
5	Patch test from der- matological clinic	78 unselected patients were patch-tested with the Portuguese standard series and textile dye mixes. Mixes with three (Dis- perse Blue 35, Disperse Blue 106, and Disperse Orange 3, each at 1 %), five (Disperse Red 1, Disperse Red 17, Disperse Yel- low 3, Disperse Blue 35, and Disperse Blue 124), and eight (five-component mix, plus Disperse Orange 3, Disperse Blue 3, and Disperse Orange 37) textile dye components were tested. Time window not reported.	Two positive reactions with five- and eight-component mixes (5 mix and 8 mix), each gave ++ reactions at 2 and 4 days Disperse Red 17: 1.3 % (1/78) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Sousa-Ba- sto and Azenha, 1994)
6	Patch test from der- matological depart- ment	593 cadets (18 to 28 years) without a present or previous his- tory of dermatitis were patch-tested with the modified GIRDCA ⁴ standard series (time window not reported). The later part of the group, comprising 336 soldiers, was also patch- tested with textile dyes, finishes, and mordant (incl. Disperse Red 17, 1 % in pet.).	In total, 74 subjects (12.5 %) were sensitised. Disperse Red 17: 1.2 % (4/336) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Seidenari et al., 1990)
	Selected dermatitis path	ients			
7	Patch test from der- matological depart- ment	01/2013 - 12/ 2015: Among 753 patients attending for cutane- ous allergy testing, 99 subjects presented with anogenital symptoms. Among patients with symptoms, 36 subjects were patch-tested with a textile and leather series (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 0 % (0/36) positive reactions	Negative	(Foley et al., 2019)
8	Retrospective analy- sis including 56 der- matological depart- ments (IVDK ⁵)	2007 - 2014: Among 98 417 patients in total, 3 207 patients with suspected textile allergy (study group) and 95 210 patients as control group were patch-tested with textile and leather dye series. Among subjects of study group, 1 594 patients were patch-tested with Disperse Red 17, 1 % in pet.	Disperse Red 17: 1.1 % (18/1 594) positive reactions (++) reaction in 5 patients, (+) in 13 patients	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Heratizadeh et al., 2017)

 ⁴ GIRDCA - Italian Research Group on Contact and Environmental Dermatitis
 ⁵ IVDK - Information Network of Departments of Dermatology

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
9	Retrospective study of dermatological de- partment	1996 - 2015: 389 children were patch-tested; reactions of 52 children with dermatitis to the feet exclusively were compared to children with dermatitis of other locations than the feet (n = 337). Patch tests were performed with GEIDAC ⁶ standard series and specific shoe series (28 patients), with additional series and the child's daily footwear, where indicated (incl. Disperse Red 17, 1 %, no further information).	Disperse Red 17: 1.9 % (1/52) positive reactions Relevance: 100 %, shoes iden- tified as source	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Ortiz-Sal- vador et al., 2017)
10	Outcome of patch tests with textile dye mix (TDM) at nine clinics from nine countries representing ICDRG ⁷	03 - 12/2013: 2 493 consecutive dermatitis patients were patch- tested with a TDM, consisting of six disperse dyes (6.6 % in pet.). 83 patients allergic to the TDM were patch-tested with eight separate dyes of the mix (incl. Disperse Red 17, 1 % in pet.). Patch test reactions to single separate dyes are presented for pa- tients allergic to the textile dye mix.	3.6 % (1.3 - 18.2 %; 90/2493) positive reactions to TDM; Patch testing with separate tex- tile dye Disperse Red 17: 16.9 % (14/83) positive reac- tions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Isaksson et al., 2015)
11	Retrospective analy- sis from dermatologi- cal departments to identify the most rele- vant allergens	01/2001 - 12/2010: 5 521 patients presented, of whom 5 281 were generally patch-tested with an extended European stand- ard series and additional allergens or series based on the der- matologist's assessment (806 patients were patch-tested with Disperse Red 17, 1 % in pet.).	Disperse Red 17: 10 % (77/806) positive reactions Disperse Red 17: 1 % (10/806) relevant reactions	Positive Low/moderate frequency of relevant reactions Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Toholka et al., 2015)
12	Patch test evaluation of clinical features and epidemiology of textile contact derma- titis	277 selected textile dermatitis patients were patch-tested, in- cluding 264 patients that were affected by allergic textile con- tact dermatitis (time window not reported). The SIDAPA ⁸ baseline series, textile series, and suspected garment samples, when available, were used for patch testing (incl. Disperse Red 17, 1 % in pet.). Only strong positive reactions (++ and +++) were considered.	154/277 positive reactions to textile allergens, (132 non-oc- cupational and 22 occupa- tional) Disperse Red 17: 3.9 % (6/154) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Lisi et al., 2014)
13	Investigations of the patch testing outcome of 12 EECDRG ⁹ clin-	01 - 06/2011: 2 907 consecutive dermatitis patients were patch- tested to TDM, 6.6 % in pet. (Comprising six disperse dyes, incl. Disperse Red 17, 1% in pet.). Ninety-four mix-positive patients were tested with single dyes.	3.7 % (108/2 907) positive re- actions to TDM; Disperse Red 17: 5.3 % (5/94) positive reactions	Positive High frequency	(Ryberg et al., 2014)

⁶ GEIDAC - Spanish Group for the Study of Contact Dermatitis and Cutaneous Allergy
⁷ ICDRG - International Contact Dermatitis Research Group
⁸ SIDAPA - Societa Italiana di Dermatologia Allergologica Professionale e Ambientale
⁹ EECDRG - European Environmental Contact Dermatitis Research Group

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
	ics from nine coun- tries to textile dye mix (TDM).			Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	
14	logical department perse dyes, and an empirical selection of problematic tattoo ink stock products, which were selected based on observations and experience in the clinic (incl. Disperse Red 17, assumed 1 % in Previous of Red 17 not		Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Serup and Hutton Carl- sen, 2014)	
15	Patch test from De- partment of Dermato- Allergology	f Dermato- baseline series supplemented with allegenes identified in a step-		Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no	(Friis et al., 2013)
16	Retrospective review of patch tests from department of derma- tology	01/2000 - 09/2011: A total of 671 patients with suspected aller- gic contact dermatitis to textile dyes and resins were patch- tested with the standard series (no further information) and ad- ditional textile dye series (containing 42 dyes and resins; 664 patients tested to Disperse Red 17, 1% in pet.).	Disperse Red 17: 5.3 % posi- tive reactions (97.1 % total rel- evant reactions) 2.3 % irritant reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Wentworth et al., 2012)
17	Patch tests from gen- eral and occupational contact dermatitis clinics at the Skin and Cancer Foundation	1993 - 2006: 2 069 patients with suspected textile allergy were patch-tested with an extended European baseline series and textile series. One hundred and fifty-seven patients reacted to any of the textile-related allergens (incl. Disperse Red 17, 1 % in pet.)	Disperse Red 17: 0.1 % (3/2 069) positive patch test re- actions	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Slodownik et al., 2011)
18	Patch test analysis from contact sensiti- sation research net- work IVDK ¹⁰			(Uter et al., 2008)	

¹⁰ IVDK - Information Network of Departments of Dermatology¹¹ DKG - German Contact Allergy Group

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
19	Patch test results from Department of Occupational and En- vironmental Derma- tology and Depart- ment of Dermatology	01/1999 - 12/2003: 3 325 patients were consecutively patch- tested with standard series of the departments including a tex- tile dye mix (TDM, incl. Disperse Red 17, 0.5 % in pet). Pa- tients who reacted positively to a patch test with the mix were tested with the eight components separately (47 subjects).	50/3 325 patients patch test re- acted positively to TDM Disperse Red 17: 11 % (5/47) positive reactions among TDM-positively tested patients (+) reaction in 4 patients, (++) in 1 patient	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Ryberg et al., 2006)
20	Patch test from BCDS ¹² dermatologi- cal departments	Over a period of 2 to 6 years, data on footwear allergens from nine dermatology centres were investigated. Patients were patch-tested to BCDS Standard series (Disperse Red 17 tested in one centre, no further information).	Disperse Red 17: 0 % (0/486) reactions	Negative	(Katugam- pola and Statham, 2005)
21	Patch test analysis from 37 IVDK ¹⁰ der- matological clinics	1998 - 2002: 696 patients with suspected textile dermatitis were patch-tested with the DKG ¹¹ textile dye series; 680 test subjects were patch-tested with Disperse Red 17 (1 % in pet.).	Disperse Red 17: 1.9 % (13/680) positive reactions (+) reaction in 9 patients, (++) in 3 patients, (+++) in 1 patient	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Bauer et al., 2004)
22	Patch test from der- matological clinic	01/1999 - 12/2002: 644 patients with suspected textile allergic contact dermatitis were patch-tested with a standard series (TRUE Tests, no further information), textile colour and finish series (TCFS) and additional series, as well as clothing extracts in 21 cases (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 0.6 % (4/644) positive reactions	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Lazarov, 2004)
23	Patch test analysis from dermatological clinic	128 patients patch-tested positively to PPD, were also patch- tested to textile dyes; the clinical presentation was dermatitis where textiles were suspected as the cause. Patch testing to tex- tile dye allergens (incl. Disperse Red 17, assumed 1 % in pet.) was performed (time window not reported).	Disperse Red 17: 11.5 % (6/61) positive reactions among PPD positively patch-tested patients	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Goon et al., 2003)
24	Patch test analysis of patients with textile dye allergic contact dermatitis from 10 clinics or physicians	09/2000: 20 patients with suspected dyed fabric allergic contact dermatitis were identified from reports of 10 clinics. Results of 16 patients, patch-tested with 12 commercial disperse dyes from the Textile Colour & Finishes series (incl. Disperse Red 17, assumed 1 % in pet.) are presented. Disperse dyes in 32	Disperse Red 17: 25 % (4/16) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Hatch et al., 2003)

¹² BCD - British Contact Dermatitis Society

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
	representing five countries	garments submitted by the patients were identified using HPLC and confirmed by LC/MS analysis.35 different disperse dyes were identified in 22/32 garments. However, Disperse Red 17 was not detected in textiles.Small number of patients investigated			
25	Retrospective analy- sis from a dermato- logical department	01/1996 - 12/1999, data of all patients patch-tested with the European standard series and showing positive patch test reactions to para-phenylenediamine (PPD) were included; 154 patients were patch-tested with para compounds and 577 patients were patch-tested with disperse (azo) dyes (incl. Disperse Red 17, assumed 1 % in pet.).	Disperse Red 17: 0.5 % (3/577) positive reactions (0/3 reactions to PPD)	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Koopmans and Bruynzeel, 2003)
26	Patch test from der- matological clinic	08/1997 - 04/2001: 203 patients with eyelid dermatitis were patch-tested with a diagnostic "standard" series and cosmetic ingredients (incl. Disperse Red 17, no further information).	Disperse Red 17: 1 % (2/203) positive reactions (++) reaction in 1 patient, (+) in 1 patient	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Guin, 2002)
27	Patch test from der- matological clinic	06/1996 - 02/2000: allergic contact dermatitis to textile aller- gens (disperse dyes), was seen in 28 (1.7 %) of 1 638 patients; 18 patients had been patch-tested to a modified British Contact Dermatitis Group standard series, and a series consisting of 18 dyes and four textile chemicals (incl. Disperse Red 17, 1% in pet.). Small number of patients	Disperse Red 17: 16.7 % (3/18) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Smith and Gawkrodger, 2002a; Smith and Gawkrodger, 2002b)
28	Patch test from der- matological clinic	During five years: 18 out of 1 400 patients with suspected con- tact dermatitis due to textile fibres were patch-tested with the GRDCI ¹³ standard battery and a battery of textile allergens (incl. Disperse Red 17, assumed 1% in pet.)	Disperse Red 17: 16.7 % (3/18) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Fuentes Cuesta et al., 2000)
29	Patch test analysis from dermatological clinic	During 1998, 103 patients with suspected allergic contact der- matitis to clothing were clinically evaluated and patch-tested with standard series (TRUE Tests) and Textile Colour & Finish series (incl. Disperse Red 17; concentration and vehicle as- sumed 1 % in pet.)	Disperse Red 17: 1.0 % (1/103) positive reactions	Positive Low/moderate frequency	(Lazarov and Cor- doba, 2000)

¹³ GRDCI - Grupo Europeo de Investigación de Dermatitis de Contacto

No.	Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
			Disperse Red 17 evokes devel- opment of purpuric allergic contact dermatitis ¹⁴	Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	
30	A retrospective study on textile dermatitis from three dermato- logical clinics	Data of 55 patients, patch-tested from 1991 to 1997 and with positive reactions to allergens from the Textile Colours and Finish series in three contact dermatitis clinics were reviewed (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 15 % positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Lazarov et al., 2000)
31	Patch test from dermatological clinic	01/1997 - 06/1999: 788 patients with textile dye allergy were patch-tested to standard series (NACDG ¹⁵ or European). A textile series was utilised in 271 patients (incl. Disperse Red 17, 1 % pet.) because of suspected textile dermatitis. Forty patients reacted positively to one or more textile dyes.	Disperse Red 17: 3.7 % (10/271) positive reactions (+++) reaction in 3 patients, (++) in 1 patient, (+) in 6 patients	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Pratt and Taraska, 2000)
32	Patch test from dermatological department	1990 - 1995: 6 203 patients were consecutively patch-tested with textile dyes included in standard series (Department of Dermatology in Modena, Italy); 236 were sensitised to at least 1 of 6 azo dyes. Thirty-three patients out of 236 azo-dye- positive subjects were patch-tested with an additional textile dye series (Disperse Red 17 included; no further information).	Disperse Red 17: 9.1 % (3/33) positive reactions among azo dye-positive patients	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Seidenari et al., 1997)
33	Patch test from dermatological clinic	04/1992 - 04/1994: 1 236 patients were patch-tested in total, among the test subjects 26 patients were identified with sus- pected contact dermatitis to textiles. Patch tests were performed with DKG ¹⁶ standard series, textile dyes, and finishing sub- stances (incl. Disperse Red 17, 1% in pet.). Small number of patients	Disperse Red 17: 11.5 % (3/26) positive reactions (++) reaction in 1 patient, (+) in 1 patient, one reading not reported	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Maurer et al., 1995)
34	Patch test analysis from dermatological department	1987 - 1991: 3 336 patients were investigated for contact dermatitis and patch-tested with the European standard series; 159 patients were also tested with 15 textile dyes (incl. Disperse Red 17, assumed 1 % in pet.).	Disperse Red 17: 3.8 % (6/159) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Dooms- Goossens, 1992)
35	Patch test from	1990 - 1991: 32 patients with presumable allergic contact	Disperse Red 17: 9.4 % (3/32)	Positive	(Thierbach

 ¹⁴ Purpuric lesions have been described as an uncommon manifestation of allergic contact dermatitis (Lazarov and Cordoba, 2000).
 ¹⁵ NACDG - North American Contact Dermatitis Group

¹⁶ DKG - German contact allergy group

Type of data/report	Relevant information about the study (as applicable)	Observations	Results³, classification	Reference
dermatological clinic	dermatitis and all with a positive patch test reaction to p- aminoazobenzene were additionally patch-tested with a series of textile azo dyes (incl. Disperse Red 17, 1 % in pet.) and one food azo dye.	positive reactions among patients positively patch-tested to p-aminoazobenzene	High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	et al., 1992)
Patch test from dermatological department	Small number of subjects 10/1987 - 04/1990: 100 subjects were found to be sensitised to textile dyes. They were identified from 2 752 consecutive patients patch-tested with the GIRDCA ⁴ standard series supplemented with disperse dyes and specifically patch testing with textile dyes. Among patients, 98 were also patch-tested with 12 further textile dyes (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 20.4 % (20/98) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Seidenari et al., 1991)
Patch test from dermatological clinic	During two years, 145 patients, suspected of having allergic contact dermatitis from textile chemicals, were patch-tested with a textile series (Disperse Red 17, 1 % pet.). In all cases, readings were ++ or +++.	Disperse Red 17: 4.4 % (3/145) positive reactions	Positive High frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Balato et al., 1990b)
Allergological study of selected workers with known exposure or dermatitis	1986 - 1987: 161 subjects with suspected occupational contact dermatitis were examined; 104 subjects were garment industry and 57 textile industry workers. Patch testing was performed using a battery of haptens prepared according to recommendations in the literature and experience using the Rapid Patch Test technique (incl. Disperse Red 17, 1 % in pet.).	Disperse Red 17: 0.6 % (1/161) positive reactions among workers 1.7 % (1/57) positive reactions among textile industry workers 0 % (0/104) reactions among garment industry workers	Positive Low/moderate frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Gasperini et al., 1989)
	Case report			
Case report	1981 - 1984: Ten patients with suspected textile dye allergy from stockings and other textiles, including black blouse, blue trousers, or grey pantsuit presented. Most subjects reported itching, and erythema at the inner thighs, shortly after wearing fabrics. Patch testing with piece of stockings and textile, dye extracts, and single dye components was performed (incl. Disperse Red 17, 1 % in pet.). Dyes were extracted from 27 commercial stockings of different colours and analysed using preparative thin-layer	2/7 women reacted positively to Disperse Red 17; (Case 8) Disperse Red 17 patch test re- actions (-/++) and dye extract (-/+) after 24 and 72h; (Case 9) Disperse Red 17 positive patch test reactions (++/+++) after 24 and 72 h, dye extract not tested Disperse Red 17 was identified	Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible	(Hausen and Schulz, 1984)
	dermatological clinic Patch test from dermatological department Patch test from dermatological clinic Allergological study of selected workers with known exposure or dermatitis	dermatological clinic dermatitis and all with a positive patch test reaction to p- aminoazobenzene were additionally patch-tested with a series of textile azo dyes (incl. Disperse Red 17, 1 % in pet.) and one food azo dye. Small number of subjects Small number of subjects Patch test from dermatological department 10/1987 - 04/1990: 100 subjects were found to be sensitised to textile dyes. They were identified from 2 752 consecutive patients patch-tested with the GIRDCA ⁴ standard series supplemented with disperse dyes and specifically patch-tested with textile dyes. Among patients, 98 were also patch-tested with 12 further textile dyes (incl. Disperse Red 17, 1 % in pet.). Patch test from dermatological clinic During two years, 145 patients, suspected of having allergic contact dermatitis from textile chemicals, were patch-tested with a textile series (Disperse Red 17, 1 % pet.). In all cases, readings were ++ or +++. Allergological study of selected workers with known exposure or dermatitis 1986 - 1987: 161 subjects with suspected occupational contact dermations in the literature and experience using the Rapid Patch Test technique (incl. Disperse Red 17, 1 % in pet.). Case report 1981 - 1984: Ten patients with suspected textile dye allergy from stockings and other textiles, including black blouse, blue trousers, or grey pantsuit presented. Most subjects reported itching, and erythema at the inner thighs, shortly after wearing fabrics. Patch testing with piece of stockings and textile, dye extracts, and single dye components was performed (incl. Disperse Red 17, 1 % in pet.). Dyes were extracted from 27 commercial stockings of different	dermatological clinic dermatitis and all with a positive patch test reaction to p- aminoazobenzene were additionally patch-tested with a series of textile azo dyes, (incl. Disperse Red 17, 1 % in pet.) and one food azo dye. positive reactions among patients positively patch-tested to p-aminoazobenzene Patch test from dermatological department 10/1987 - 04/1990: 100 subjects were found to be sensitised to textile dyes. They were identified from 2 752 consecutive patients patch-tested with the GIRDCA ⁴ standard series supplemented with disperse dyes and specifically patch testing with textile dyes. Among patients, 98 were also patch-tested with 12 further textile dyes (incl. Disperse Red 17, 1 % in pet.). Disperse Red 17: 20.4 % (20/98) positive reactions Patch test from dermatological clinic During two years, 145 patients, suspected of having allergic contact dermatitis from textile chemicals, were patch-tested with a textile series (Disperse Red 17, 1 % pet.). In all cases, readings were ++ or +++. Disperse Red 17: 0.6 % (1/161) positive reactions among workers Allergological study of selected workers with known exposure or dermatitis 1986 - 1987: 161 subjects with suspected occupational contact dermations in the literature and experience using the Rapid Patch Test technique (incl. Disperse Red 17, 1 % in pet.). Disperse Red 17: 0.6 % (1/161) positive reactions among workers Case report 1981 - 1984: Ten patients with suspected textile dye allergy from stockings and other textiles, including black blouse, blue trousers, or grey pantsuit presented. Most subjects reported itching, and erythema at the inner thighs, shortly after wearing fabrics. Patch testing with piece of stockings and textile, dye extracts, and sing	dermatological clinic antinoazobenzene were additionally patch-tested with a positive patch-tested with a positive reactions among patients positively patch-tested to p-aminoazobenzene were additionally patch-tested to p-aminoazobenzene Figh Frequency Previous exposure to Disperse Red 17 not documented, no sub-categorisation possible Patch test from dermatological department 10/1987 - 04/1990: 100 subjects were found to be sensitised to p-aminoazobenzene were identified from 2 752 consecutive patients patch-tested with disperse dyes and specifically patch testing with textile dyes. Among patients, 98 were also patch-tested with 12 further textile dyes (incl. Disperse Red 17, 1 % in pet.). Disperse Red 17: 20.4 % (20/98) positive reactions with extile dyes, Among patients, 98 were also patch-tested with textile dyes. Among patients, 98 were patch-tested with a textile series (Disperse Red 17, 1 % pet.). In all cases, readings were ++ or +++. Disperse Red 17: 4.4 % (3/145) positive reactions workers Positive Allergological study of selected worker with know reposure or dermatitis a battery of haptens prepared according to readings a battery of haptens prepared according to reading a dattery of selected workers with know recks. Parch testing a

There is strong evidence from human data that Disperse Red 17 consistently and repetitively evokes skin sensitisation as indicated in diagnostic patch tests from individual clinics or collated clinic data. Patch test studies considered as reliable (reliable with restriction) and relevant are summarised in Table 12, while studies of low reliability (not reliable or not assignable to the DS) were precluded from further assessment (Chromej et al., 2008; Cunha et al., 2003; Garcia-Bravo et al., 2004; Gee and Powell, 2001; Su et al., 2014; Thomas et al., 2013).

Diagnostic patch tests comprise studies with unselected (consecutive) or selected dermatitis patients analysing the number of patients sensitised to Disperse Red 17 compared to all patients tested in a certain time period. In studies with unselected, consecutive dermatitis patients, patch testing is generally more standardised. In contrast, for selected (specific) groups of patients or workers, usually targeted patch testing with special test series is performed.

Consecutive patients patch-tested with Disperse Red 17 show frequencies between 0.2 % and 1.3 % positive reactions. According to ECHA's Guidance on the application of CLP criteria, most of the studies revealed a relatively low/moderate frequency of positive patch test reactions (4/6 studies, frequency < 1 % for consecutive, unselected dermatitis patients, Section 3.4.2.2.3.1, Table 3.2 (ECHA, 2017)). Less studies revealed a relatively high frequency for Disperse Red 17 positive reactions among consecutive dermatitis patients (2/6 studies, frequency ≥ 1 % for consecutive, unselected dermatitis patients, Section 3.4.2.2.3.1, Table 3.2 (ECHA, 2017)).

Aimed testing with Disperse Red 17 in selected dermatitis patients identified between 0 % and 25 % positive patch test reactions. The majority of the patch tests identified high frequencies of Disperse Red 17 positively patch-tested patients (17/32 studies, frequency ≥ 2 % for selected dermatitis patients (ECHA, 2017)). A low/moderate frequency of Disperse Red 17 reactions in selected dermatitis patients was seen in numerous publications (13/32 studies, frequency < 2 % for selected dermatitis patients). Two available studies on patch testing in selected dermatitis patients revealed negative results for Disperse Red 17.

Furthermore, a high number of case reports are available, which describe the patients' clinical history with dermatitis due to wearing coloured textiles or contact to hair dyes and report positive patch test results for Disperse Red 17 in these subjects. Hausen and Schulz investigated ten women with suspected textile dye allergy from stockings and other dyed textiles, including a black blouse, blue trousers, or grey pantsuit (Hausen and Schulz, 1984). Most subjects reported itching and erythema on the inner thighs, shortly after wearing fabrics. The authors analysed the dyes of 27 commercial stockings using chromatography, and among other dyes identified Disperse Red 17. Positive patch test reactions to Disperse Red 17 were shown in two patients with dermatitis from stockings. Further case reports with Disperse Red 17 positive patch test reaction were considered not relevant. In these reports, a detection of Disperse Red 17 in the suspected source of dermatitis (e.g. textiles) is missing (Alberta et al., 2005; Ameur et al., 2019; Batchelor and Wilkinson, 2006; Crichlow and Warin, 2004; Dejobert et al., 1995; Fuentes Cuesta et al., 2000; Goldminz and Scheinman, 2018; Hausen, 2006; Hausen, 1993; Kind et al., 2012; Kuuliala et al., 2006; Lisboa et al., 1994; Mohamoud and Andersen, 2017; Mota et al., 2000; Nakagawa et al., 1996; Narganes et al., 2013; Patrizi et al., 1990; Pousa-Martínez et al., 2018; Pousa-Martinez et al., 2016; Pratt and Taraska, 2000; Raffi et al., 2019; Ramírez et al., 2007; Saunders et al., 2004; Seidenari et al., 1995; Shehade and Beck, 1990; Su and Horton, 1998; Warren and Marren, 1997; Wilkinson and Thomson, 2000).

Human data do not give information on previous exposure levels to Disperse Red 17. Furthermore, human induction studies such as a Human Repeated Insult Patch Test (HRIPT) or Human Maximisation Test (HMT) performed with Disperse Red 17 are not available to the DS.

Analytical investigations using thin-layer chromatography revealed that Disperse Red 17 patch test preparations showed not only one main spot, but also one (or two) additional weaker spots in the chromatograms (Foussereau and Dallara, 1986; Ryberg et al., 2008). However, there was no characterisation of the additional spots with regard to cleavage products or impurities. Patch testing with impure preparations may result in false positive test results and complicate the diagnosis of the patients and prevention of contact allergy. Furthermore, analysis revealed that the mean concentrations of several commercial Disperse Red 17 patch test preparations were much lower than labelled (0.35 % (0.3-0.5 %) instead of 1.0 %, 14 preparations (Ryberg et al., 2008)). In such cases an overestimation of the dye concentration causing sensitisation (and therefore, an underestimation of its potency) and/or a smaller than expected number of sensitised subjects may result. However, the

occurrence of positive diagnostic patch test reactions from a large number of dermatological clinics, representing numerous different countries leaves no doubt that Disperse Red 17 elicits skin sensitisation in humans.

9.7.3 Short summary and overall relevance of the provided information on skin sensitisation

Two animal studies investigated the skin sensitising potential of Disperse Red 17, namely a none-guideline "Sensitive mouse lymph node assay" and a GPMT, performed according to OECD testing guideline 406. None of these studies on skin sensitisation obtained any positive test result for Disperse Red 17. However, these studies were considered too unreliable to allow for a conclusion on the skin sensitising properties of Disperse Red 17 (for details cf. section 9.7.1).

There is strong evidence from human data that Disperse Red 17 evokes skin sensitisation in humans (shown in > 200 subjects), indicated by dermatological patch tests and performed in a high number of dermatological clinics from several countries as well as published cases. Patch test data reveal mainly low/moderate frequencies of Disperse Red 17 positive patch test reactions among unselected, consecutive dermatitis patients (<1 % positive reactions for consecutive, unselected dermatitis patients, Section 3.4.2.2.3.1, Table 3.2 (ECHA, 2017)). Selected dermatitis patients patch-tested with Disperse Red 17 show mainly high frequencies of positive reactions (≥ 2 % positive reactions for selected dermatitis patients (ECHA, 2017)). The available human data do not give information on the previous level of exposure to Disperse Red 17. Human tests on induction thresholds of Disperse Red 17 are not available. In conclusion, data are insufficient to allow for classification into sub-categories.

9.7.4 Comparison with the CLP criteria

Table 13: Comparison of human and animal data for skin sensitisation of Disperse Red 17 with CLP criteria

Reference(s)	Criteria acc. to CLP regulation, as laid out in (ECHA, 2017)	Results	Resulting Clas- sification
Animal data			
No reliable studies available			
Human data			
Dermatitis patients (unselected, con- secutive) (Lazarov and Cordoba, 2000; Li, 2010; Ryberg et al., 2009; Seidenari et al., 1990; Sousa-Basto and Azenha, 1994; Toholka et al., 2015)	Skin Sens. 1Frequency < 1.0 % and relatively low	Frequency from "relatively low/moderate" to "relatively high" 4/6 studies re- vealed a rela- tively low/mod- erate frequency Exposure un- clear	Skin Sens. 1 (no sub-categori- sation possible)
Selected dermatitis patients (aimed testing) (Balato et al., 1990b; Bauer et al., 2004; Dooms-Goossens, 1992; Foley et al., 2019; Friis et al., 2013; Fuentes Cuesta et al., 2000; Gasperini et al., 1989; Goon et al., 2003; Guin, 2002; Hatch et al., 2003; Heratizadeh et al., 2017; Isaksson et al., 2015; Katugampola and Statham, 2005; Knackstedt and Zug, 2015; Lazarov, 2004; Lazarov and Cordoba, 2000; Lazarov et al., 2000; Lisi et al., 2014; Maurer et	Skin Sens. 1Frequency < 2.0 % and relatively low	Frequency from negative, "rela- tively low/mod- erate" to "rela- tively high" 17/32 studies with a relatively high frequency 13/32 studies with a relatively low/moderate frequency	Skin Sens. 1 (no sub-categori- sation possible)

Reference(s)	Criteria acc. to CLP regulation, as laid out in (ECHA, 2017)	Results	Resulting Clas- sification
al., 1995; Ortiz-Salvador et al., 2017; Pratt and Taraska, 2000; Ryberg et al., 2014; Ryberg et al., 2006; Sei- denari et al., 1997; Seidenari et al., 1991; Serup and Hutton Carlsen, 2014; Slodownik et al., 2011; Smith and Gawkrodger, 2002a; Smith and Gawkrodger, 2002b; Thierbach et al., 1992; Toholka et al., 2015; Uter et al., 2008; Wentworth et al., 2012)		2/32 studies with negative results Exposure un- clear	
Number of published cases (Hausen and Schulz, 1984)	$\frac{\text{Skin Sens. 1}}{\text{Number of published cases} < 100 \text{ and}}$ relatively low exposure or Number of published cases ≥ 100 and relatively high exposure $\frac{\text{Skin Sens. 1A}}{\text{Number of published cases} \geq 100 \text{ and}}$ relatively low exposure $\frac{\text{Skin Sens. 1B}}{\text{Skin Sens. 1B}}$ Number of published cases < 100 and relatively high exposure	Low/moderate frequency Two published cases Exposure un- clear	Skin Sens. 1 (no sub-categori- sation possible)

9.7.4.1 Weight of evidence consideration

According to the CLP regulation "substances shall be classified as skin sensitisers (Category 1) where data are not sufficient for sub-categorisation in accordance with the following criteria:

- a) if there is evidence in humans that the substance can lead to sensitisation by skin contact in a substantial number of persons; or
- b) if there are positive results from an appropriate animal test" (CLP, Annex I, Table 3.4.2).

Classification into sub-categories is required when data are sufficient. Human evidence for sub-categorisation can include data on the induction threshold and/or exposure level a substance (CLP, Section 3.4.2.2.2.). Furthermore, a skin sensitisation potency from animal studies can be used for sub-categorisation (CLP, Annex I, Table 3.4.3 and 3.4.4)."

There is a large number of human studies, including diagnostic patch tests and case reports performed in multiple dermatological clinics from different countries, which show that Disperse Red 17 elicits skin sensitisation in humans. Frequencies of Disperse Red 17 positive patch test reactions are mainly low/moderate for consecutive, unselected dermatitis patients (< 1.0 % occurrence of skin sensitisation). Selected dermatitis patients reveal mostly high frequencies of positive patch test reactions for Disperse Red 17 (\geq 2 % occurrence of skin sensitisation) and low/moderate frequencies (< 2 % occurrence of skin sensitisation), while a low number of studies show negative results for Disperse Red 17. However, available data do not give information on exposure levels or an induction threshold of Disperse Red 17. Therefore, sub-categorisation is not possible based on the available human data.

The available animal data on skin sensitisation are unreliable and cannot be used for classification.

Altogether, based on the positive data from patch testing, obtained in numerous different dermatology clinics it is warranted to classify Disperse Red 17 as a skin sensitiser. However, available data are not sufficient for sub-categorisation. In conclusion, Disperse Red 17 shall be classified as skin sensitiser in Category 1.

9.7.5 Conclusion on classification and labelling for skin sensitisation

Based on the data in Table 13 the DS proposes to classify Disperse Red 17 as skin sensitiser, Skin Sens 1 (H317 - May cause an allergic skin reaction), without sub-categorisation, and with a GCL of 1 % (w/v).

9.8 Germ cell mutagenicity

Not assessed in this dossier

9.9 Carcinogenicity

Not assessed in this dossier

9.10 Reproductive toxicity

Not assessed in this dossier

9.11 Specific target organ toxicity-single exposure

Not assessed in this dossier

9.12 Specific target organ toxicity-repeated exposure

Not assessed in this dossier

9.13 Aspiration hazard

Not assessed in this dossier

10 EVALUATION OF ENVIRONMENTAL HAZARDS

Not assessed in this dossier

11 EVALUATION OF ADDITIONAL HAZARDS

Not assessed in this dossier

12 ADDITIONAL LABELLING

Not relevant

13 REFERENCES

Alberta L., Sweeney S.M., and Wiss K. (2005): Diaper dye dermatitis. Pediatrics 116 (3), e450-e452. DOI: 10.1542/peds.2004-2066

Ameur K., Youssef M., Belhadjali H., Soua Y., Korbi M., Henchi M.A., and Zili J. (2019): Occupational acute generalized exanthematous pustulosis induced by disperse dyes in a textile. Contact Dermatitis 80 (6), 411-412. DOI: 10.1111/cod.13241

Balato N., Lembo G., De Stetano S., Patruno C., De Stefano R., and Ayala F. (1990a): Various Identification of textile dyes by laboratory methods. Contact Dermatitis 23 (4), 266-266. DOI: 10.1111/j.1600-0536.1990.tb05079.x

Balato N., Lembo G., Patruno C., and Ayala F. (1990b): Prevalence of textile dye contact sensitization. Contact Dermatitis 23 (2), 111-112. DOI: 10.1111/j.1600-0536.1990.tb03232.x

Batchelor R.J. and Wilkinson S.M. (2006): Contact allergy to disperse dyes in plastic spectacle frames. Contact Dermatitis 54 (1), 66-67. DOI: 10.1111/j.0105-1873.2006.0729h.x

Bauer A., Geier J., Lessmann H., and Elsner P. (2004): Kontaktallergien gegen Textilfarbstoffe. Ergebnisse des Informationsverbundes Dermatologischer Kliniken (IVDK). Aktuelle Dermatologie 30 (1), 23-27. DOI: 10.1055/s-2004-814276

BVL (2010): Berichte zur Lebensmittelsicherheit 2009 9783034800587 (online) 9783034800570 (print) Bundesamt für Verbraucherschutz und Lebensmittelsicherheit. Springer B. DOI: 10.1007/978-3-0348-0058-7

Chromej I., Frlickova Z., and Chribikova I. (2008): Frequency and relevance of contact sensitization to textile dyes. Contact Dermatitis 58 (s1), 28. DOI: 10.1111/j.1600-0536.2008.01380.x

Clauss B. and Weiss M. (1992): Effect of electron radiation on dyed PET and PA 66 substrates. Die Angewandte Makromolekulare Chemie 197 (1), 185-199. DOI: 10.1002/apmc.1992.051970116

Crichlow S. and Warin A.P. (2004): Allergic contact dermatitis from dyes in wigs following diphencyprone treatment. Contact Dermatitis 51 (3), 148-149. DOI: 10.1111/j.0105-1873.2004.0426b.x

Cunha A.P., Barros M.A., and Resende C. (2003): Contact dermatitis of the feet Journal of the European Academy of Dermatology and Venereology 17 (s3 Posters), 221. DOI: 10.1046/j.1468-3083.17.s3.8.x

de Groot A.C. (2013): Side-effects of henna and semi-permanent 'black henna' tattoos: A full review. Contact Dermatitis 69 (1), 1-25. DOI: 10.1111/cod.12074

Dejobert Y., Martin P., Thomas P., and Bergoend H. (1995): Multiple azo dye sensitization revealed by the wearing of a black "velvet" body. Contact Dermatitis 33 (4), 276-277. DOI: 10.1111/j.1600-0536.1995.tb00490.x

Dooms-Goossens A. (1992): Textile dye dermatitis. Contact Dermatitis 27 (5), 321-323. DOI: 10.1111/j.1600-0536.1992.tb03289.x

ECHA (2014): Guidance on the preparation of CLH dossiers for harmonised classification and labelling. European Chemicals Agency. ISBN: 978-92-9244-748-9

ECHA (2017): Guidance on the Application of the CLP Criteria. In: Guidance to Regulation (EC) No 1272/2008 on classification, labelling and packaging (CLP) of substances and mixtures. European Chemical Agency. ISBN: 978-92-9020-050-5. DOI: 10.2823/124801

ECHA (2019a): Restriction of substances in tattoo inks and permanent make up. European Chemical Agency. <u>https://echa.europa.eu/de/registry-of-restriction-intentions/-/dislist/details/0b0236e180dff62a</u> (last accessed 17 June 2020)

ECHA (2019b): Restriction on the placing on the market of textile, leather, hide and fur articles containing skin sensitising substances. European Chemical Agency. <u>https://echa.europa.eu/restrictions-under-consideration/-/substance-rev/23405/term</u> (last accessed 17 June 2020)

ECHA (2020): Dissemination side. European Chemical Agency (last accessed 16 November 2020)

EU Ecolabel (2015): EU Ecolabel textile products user manual - Commission Decision for the award of the EU Ecolabel for textile products (2014/350/EU), Version 1.0 MArch 2015. <u>https://www.eu-ecolabel.de/eu-ecolabel.html</u> (last accessed 29 Jan. 2020)

Foley C.C., White S., Merry S., Nolan U., Moriarty B., Kirby B., Collins P., and Lally A. (2019): Understanding the role of cutaneous allergy testing in anogenital dermatoses: A retrospective evaluation of contact sensitization in anogenital dermatoses. International Journal of Dermatology 58 (7), 806-810. DOI: 10.1111/ijd.14360

Foussereau J. and Dallara J.M. (1986): Purity of standardized textile dye allergens: A thin layer chromatography study. Contact Dermatitis 14 (5), 303-306. DOI: 10.1111/j.1600-0536.1986.tb05281.x

Friis U.F., Menné T., Flyvholm M.-A., Bonde J.P.E., and Johansen J.D. (2013): Occupational allergic contact dermatitis diagnosed by a systematic stepwise exposure assessment of allergens in the work environment. Contact Dermatitis 69 (3), 153-163. DOI: 10.1111/cod.12102

Fuentes Cuesta M.M., Blanco Carmona J.G., Herrero Gil D., Perez Gimenez R., Garces Sotillos M., Garcia Gonzalez F., Juste Picon S., and Carretero Anibarro P. (2000): Contact allergic dermatitis due to textile fabrics. Alergologia e Inmunologia Clinica 15 (2), 88-92

Garcia-Bravo B., Fernandez-Redondo V., and Sanchez-Pedreño P. (2004): Contact dermatitis from textile colours in three Spanish towns. Contact Dermatitis 50 (3), 177-178. DOI: 10.1111/j.0105-1873.2004.00309ei.x

Gasperini M., Farli M., Lombardi P., and Sertoli A. (1989): Contact dermatitis in the textile and garment industry. In: Current Topics in Contact Dermatitis (Frosch P., Dooms-Goossens A., Lachapelle J.M., Rycroft R.J.G., and Scheper R.J., eds.), pp. 326-329. Springer Berlin Heidelberg, Berlin, Heidelberg. ISBN: 978-3-642-74299-6. DOI: 10.1007/978-3-642-74299-6_68

Gee B.C. and Powell S.M. (2001): A prospective study investigating prevalence of allergic contact dermatitis to textile allergens in Oxford. British Journal of Dermatology 145 (Suppl 59), 94. DOI: 10.1046/j.1365-2133.2001.093ff.x

Goldminz A.M. and Scheinman P.L. (2018): A case series of dupilumab-treated allergic contact dermatitis patients. Dermatologic Therapy 31 (6), e12701. DOI: 10.1111/dth.12701

Goon A.T., Gilmour N.J., Basketter D.A., White I.R., Rycroft R.J., and McFadden J.P. (2003): High frequency of simultaneous sensitivity to Disperse Orange 3 in patients with positive patch tests to paraphenylenediamine. Contact Dermatitis 48 (5), 248-250. DOI: 10.1034/j.1600-0536.2003.00049.x

Guin J.D. (2002): Eyelid dermatitis: Experience in 203 cases. Journal of the American Academy of Dermatology 47 (5), 755-765. DOI: 10.1067/mjd.2002.122736

Hatch K.L., Motschi H., and Maibach H.I. (2003): Disperse dyes in fabrics of patients patch-test-positive to disperse dyes. American Journal of Contact Dermatitis 14 (4), 205-212. <u>https://www.ncbi.nlm.nih.gov/pubmed/14738722</u>

Hausen B. (2006): Die rote Frau. Aktuelle Dermatologie 32 (12), 527-532. DOI: 10.1055/s-2006-945004

Hausen B.M. (1993): Contact allergy to Disperse Blue 106 and Blue 124 in black "velvet" clothes. Contact Dermatitis 28 (3), 169-173. DOI: 10.1111/j.1600-0536.1993.tb03381.x

Hausen B.M. and Schulz K.H. (1984): [Allergy to dyes in stockings]. Deutsche Medizinische Wochenschrift 109 (39), 1469-1475. DOI: 10.1055/s-2008-1069396

Heratizadeh A., Geier J., Molin S., and Werfel T. (2017): Contact sensitization in patients with suspected textile allergy. Data of the Information Network of Departments of Dermatology (IVDK) 2007-2014. Contact Dermatitis 77 (3), 143-150. DOI: 10.1111/cod.12760

Ikarashi Y., Tsuchiya T., and Nakamura A. (1996): Application of sensitive mouse lymph node assay for detection of contact sensitization capacity of dyes. Journal of Applied Toxicology 16 (4), 349-354. DOI: 10.1002/(SICI)1099-1263(199607)16:4<349::AID-JAT351>3.0.CO;2-5

Isaksson M., Ale I., Andersen K.E., Diepgen T., Goh C.L., Goossens R.A., Jerajani H., Maibach H.I., Sasseville D., and Bruze M. (2015): Patch testing to a textile dye mix by the international contact dermatitis research group. Dermatitis : contact, atopic, occupational, drug 26 (4), 170-176. DOI: 10.1097/der.00000000000125

Johansen J.D., Aalto-Korte K., Agner T., Andersen K.E., Bircher A., Bruze M., Cannavó A., Giménez-Arnau A., Gonçalo M., Goossens A., John S.M., Lidén C., Lindberg M., Mahler V., Matura M., Rustemeyer T., Serup J., Spiewak R., Thyssen J.P., Vigan M., White I.R., Wilkinson M., and Uter W. (2015): European Society of Contact Dermatitis guideline for diagnostic patch testing – recommendations on best practice. Contact Dermatitis 73 (4), 195-221. DOI: 10.1111/cod.12432

Karunaratne S.D. (1995): Disperse red 17 skin sensitisation study in the guinea pig. Toxicol. Lab. LTD

Katugampola R.P. and Statham B.N. (2005): A review of allergens found in current hair-care products. Contact Dermatitis 53 (4), 234-235. DOI: 10.1111/j.0105-1873.2005.0670a.x

Kind F., Scherer K., and Bircher A.J. (2012): Contact dermatitis to para-phenylenediamine in hair dye following sensitization to black henna tattoos - An ongoing problem. JDDG: Journal der Deutschen Dermatologischen Gesellschaft 10 (8), 572-577. DOI: 10.1111/j.1610-0387.2011.07882.x

Knackstedt T.J. and Zug K.A. (2015): T cell lymphomatoid contact dermatitis: A challenging case and review of the literature. Contact Dermatitis 72 (2), 65-74. DOI: 10.1111/cod.12294

Koopmans A.K. and Bruynzeel D.P. (2003): Is PPD a useful screening agent? Contact Dermatitis 48 (2), 89-92. DOI: 10.1034/j.1600-0536.2003.480207.x

Kuuliala O., Alanko K., and Jolanki R. (2006): Two types of contact allergy from the same leather/textile gloves. Contact Dermatitis 55 (Poster Presentations s1), 47. DOI: 10.1111/j.1600-0536.2006.00936_2.x

Lazarov A. (2004): Textile dermatitis in patients with contact sensitization in Israel: a 4-year prospective study. Journal of the European Academy of Dermatology and Venereology : JEADV 18 (5), 531-537. DOI: 10.1111/j.1468-3083.2004.00967.x

Lazarov A. and Cordoba M. (2000): Purpuric contact dermatitis in patients with allergic reaction to textile dyes and resins. Journal of the European Academy of Dermatology and Venereology : JEADV 14 (2), 101-105. DOI: 10.1046/j.1468-3083.2000.00025.x

Lazarov A., Trattner A., David M., and Ingber A. (2000): Textile dermatitis in Israel: A retrospective study. American Journal of Contact Dermatitis 11 (1), 26-29. DOI: 10.1053/ajcd.1999.0026

Le Coz C.J. and Tromp G. (2002): Skin paints (pseudo-tattoos) with 'Black Henna': hazards and long-term risks. Exogenous Dermatology 1 (5), 246-252

Li L.F. (2010): Contact sensitization to textile dyes in a self-selected population and a dermatological referral population in Beijing. Contact Dermatitis 63 (5), 291-292. DOI: 10.1111/j.1600-0536.2010.01780.x

Lisboa C., Barros M.A., and Azenha A. (1994): Contact dermatitis from textile dyes. Contact Dermatitis 31 (1), 9-10. DOI: 10.1111/j.1600-0536.1994.tb01896.x

Lisi P., Stingeni L., Cristaudo A., Foti C., Pigatto P., Gola M., Schena D., Corazza M., and Bianchi L. (2014): Clinical and epidemiological features of textile contact dermatitis: An Italian multicentre study. Contact Dermatitis 70 (6), 344-350. DOI: 10.1111/cod.12179

Malinauskiene L., Zimerson E., Bruze M., Ryberg K., and Isaksson M. (2012): Are allergenic disperse dyes used for dyeing textiles? Contact Dermatitis 67 (3), 141-148. DOI: 10.1111/j.1600-0536.2012.02129.x

Maurer S., Seubert A., S. S., and Fuchs T. (1995): Kontaktallergien auf Textilien. Dermatosen Beruf Umwelt 43, 63-68

Mohamoud A.A. and Andersen F. (2017): Allergic contact dermatitis caused by textile dyes mimicking atopic dermatitis. Contact Dermatitis 76 (2), 119-120. DOI: 10.1111/cod.12630

Moro P.A., Morina M., Milani F., Pandolfi M., Guerriero F., and Bernardo L. (2016): Sensitization and Clinically Relevant Allergy to Hair Dyes and Clothes from Black Henna Tattoos: Do People Know the Risk? An Uncommon Serious Case and a Review of the Literature. Cosmetics 3 (3), 23

Mota F., Silva E., Varela P., Azenha A., and Massa A. (2000): An outbreak of occupational textile dye dermatitis from Disperse Blue 106. Contact Dermatitis 43 (4), 235-237. DOI: DOI: 10.1034/j.1600-0536.2000.042004235.x

Nakagawa M., Kawai K., and Kawai K. (1996): Multiple azo disperse dye sensitization mainly due to group sensitizations to azo dyes. Contact Dermatitis 34 (1), 6-11. DOI: 10.1111/j.1600-0536.1996.tb02103.x

Narganes L.M.V., Sambucety P.S., Gonzalez I.R., Rivas M.O., and Prieto M.A.R. (2013): Lymphomatoid dermatitis caused by contact with textile dyes. Contact Dermatitis 68 (1), 62-64. DOI: 10.1111/j.1600-0536.2012.02164.x

OEKO-TEX (2020): Standard 100 by OEKO TEX®, Limit Values of Individual Substances, According to Appendices 5 & 7, Edition 02.2020. OEKO-TEX®. <u>https://www.oekotex.com/en/business/certifications_and_services/ots_100/ots_100_start.xhtml</u> (last accessed 29 Jan. 2020)

Ortiz-Salvador J.M., Esteve-Martínez A., García-Rabasco A., Subiabre-Ferrer D., Martínez-Leboráns L., and Zaragoza-Ninet V. (2017): Dermatitis of the foot: Epidemiologic and clinical features in 389 children. Pediatric Dermatology 34 (5), 535-539. DOI: 10.1111/pde.13203

Patrizi A., Lanzarini M., and Tosti A. (1990): Persistent patch test reactions to textile dyes. Contact Dermatitis 23 (1), 60-61. DOI: 10.1111/j.1600-0536.1990.tb00095.x

Pousa-Martínez M., González-Rodríguez C., Rodriguez-Rodriguez M., Sainz-Gaspar L., Sardina F.J., and Fernández-Redondo V. (2018): Garment allergy caused by Disperse Blue 360: A new sensitizer. Contact Dermatitis 79 (1), 37-38. DOI: 10.1111/cod.12975

Pousa-Martinez M., Rodriguez-Rodriguez M., Vázquez-Veiga H., and Fernández-Redondo V. (2016): Cutaneous reaction to a garment: Unusual clinical presentation. Contact Dermatitis 75, 77. DOI: 10.1111/cod.12637

Pratt M. and Taraska V. (2000): Disperse blue dyes 106 and 124 are common causes of textile dermatitis and should serve as screening allergens for this condition. American Journal of Contact Dermatitis 11 (1), 30-41. DOI: 10.1053/ajcd.2000.0030

Raffi J., Chen R., and Botto N. (2019): Wide dye reactors. JAAD case reports 5 (10), 877-879. DOI: 10.1016/j.jdcr.2019.08.005

Ramírez A., Pérez-Pérez L., Fernández-Redondo V., and Toribio J. (2007): Photoallergic dermatitis induced by diltiazem. Contact Dermatitis 56 (2), 118-119. DOI: 10.1111/j.1600-0536.2007.00967.x

Ryberg K., Agner T., Andersen K.E., Bircher A., Diepgen T., Foti C., Giménez-Arnau A., Gonçalo M., Goossens A., Johansen J.D., Le Coz C., Maibach H.I., and Bruze M. (2014): Patch testing with a textile dye mix - A multicentre study. Contact Dermatitis 71 (4), 215-223. DOI: 10.1111/cod.12244

Ryberg K., Goossens A., Isaksson M., Gruvberger B., Zimerson E., Nilsson F., Bjork J., Hindsen M., and Bruze M. (2009): Is contact allergy to disperse dyes and related substances associated with textile dermatitis? Br J Dermatol 160 (1), 107-115. DOI: 10.1111/j.1365-2133.2008.08953.x

Ryberg K., Gruvberger B., Zimerson E., Isaksson M., Persson L., Sorensen O., Goossens A., and Bruze M. (2008): Chemical investigations of disperse dyes in patch test preparations. Contact Dermatitis 58 (4), 199-209. DOI: 10.1111/j.1600-0536.2007.01298.x

Ryberg K., Isaksson M., Gruvberger B., Hindsen M., Zimerson E., and Bruze M. (2006): Contact allergy to textile dyes in southern Sweden. Contact Dermatitis 54 (6), 313-321. DOI: 10.1111/j.0105-1873.2006.00733.x

Saunders H., O'Brien T., and Nixon R. (2004): Textile dye allergic contact dermatitis following paraphenylenediamine sensitization from a temporary tattoo. Australas J Dermatol 45 (4), 229-231. DOI: 10.1111/j.1440-0960.2004.00110.x

SCCS (2013): Opinion on disperse red 17 - COLIPA no. B5 (Safety S.C.o.C., ed.). European Union. ISBN: 1831-4767. DOI: 10.2772/74722

Seidenari S., Mantovani L., Manzini B.M., and Pignatti M. (1997): Cross-sensitizations between azo dyes and para-amino compound. Contact Dermatitis 36 (2), 91-96. DOI: 10.1111/j.1600-0536.1997.tb00420.x

Seidenari S., Manzini B.M., and Danese P. (1991): Contact sensitization to textile dyes: description of 100 subjects. Contact Dermatitis 24 (4), 253-258. DOI: 10.1111/j.1600-0536.1991.tb01718.x

Seidenari S., Manzini B.M., Danese P., and Motolese A. (1990): Patch and prick test study of 593 healthy subjects. Contact Dermatitis 23 (3), 162-167. DOI: 10.1111/j.1600-0536.1990.tb04777.x

Seidenari S., Manzini B.M., Schoiavi M.E., and Motolese A. (1995): Prevalence of contact allergy to nondisperse azo dyes for natural fibers: A study in 1814 consecutive patients. Contact Dermatitis 33 (2), 118-122. DOI: 10.1111/j.1600-0536.1995.tb00512.x

Serup J. and Hutton Carlsen K. (2014): Patch test study of 90 patients with tattoo reactions: Negative outcome of allergy patch test to baseline batteries and culprit inks suggests allergen(s) are generated in the skin through haptenization. Contact Dermatitis 71 (5), 255-263. DOI: 10.1111/cod.12271

Shehade S.A. and Beck M.H. (1990): Contact dermatitis from disperse dyes in synthetic wigs. Contact Dermatitis 23 (2), 124-125. DOI: 10.1111/j.1600-0536.1990.tb03243.x

Slodownik D., Williams J., Tate B., Tam M., Cahill J., Frowen K., and Nixon R. (2011): Textile allergy--the Melbourne experience. Contact dermatitis 65 (1), 38-42. DOI: 10.1111/j.1600-0536.2010.01861.x

Smith J. and Gawkrodger D.J. (2002a): Contact dermatitis from textile and dye allergens requires a high index of suspicion for diagnosis. A review of 18 cases. Contact Dermatitis 47 (2), 109-125. DOI: 10.1034/j.1600-0536.2002.470210_4.x

Smith J.K. and Gawkrodger D.J. (2002b): Contact dermatitis to textile and dye allergens requires a high degree of suspicion for diagnosis: A review of 18 cases. British Journal of Dermatology 147 (Suppl 62), 72. DOI: 10.1034/j.1600-0536.2002.470210_4.x

Sousa-Basto A. and Azenha A. (1994): Textile dye mixes: Useful screening tests for textile dye allergy. Contact Dermatitis 30 (3), 189-189. DOI: 10.1111/j.1600-0536.1994.tb00716.x

Su J.C. and Horton J.J. (1998): Allergic contact dermatitis from azo dyes. Australasian Journal of Dermatology 39 (1), 48-49. DOI: 10.1111/j.1440-0960.1998.tb01243.x

Su O., Ozkaya D.K., Pirmit S., Ulusal H.A., and Onsun N. (2014): The allergens causing contact sensitization in textile industry workers. Turkderm-Archives of the Turkish Dermatology and Venerology 48 (3), 140-145. DOI: 10.4274/turkderm.35761

Suter H. (1965): Untersuchungen über das Polyamidstrumpfekzem. Dermatologica 130, 411-424. DOI: 10.1159/000254556

Svedman C., Engfeldt M., and Malinauskiene L. (2019): Textile contact dermatitis: How fabrics can induce dermatitis. Current Treatment Options in Allergy 6 (1), 103-111. DOI: 10.1007/s40521-019-0197-5

Thierbach M.A., Geursen-Reitsma A.M., and van Joost T. (1992): Sensitization to azo dyes: Negative patch tests to yellow and red azo dyes in printed paper. Contact Dermatitis 27 (1), 22-26. DOI: 10.1111/j.1600-0536.1992.tb05193.x

Thomas B., White I., White J., McFadden J., and Banerjee P. (2013): Sensitivity to p-phenylenediamine (PPD): Positive relationship of response to PPD on patch testing and cross-reactions with other chemically related allergens. British Journal of Dermatology 169, 136. DOI: 10.1111/bjd.12359

Toholka R., Wang Y.-S., Tate B., Tam M., Cahill J., Palmer A., and Nixon R. (2015): The first Australian Baseline Series: Recommendations for patch testing in suspected contact dermatitis. Australasian Journal of Dermatology 56 (2), 107-115. DOI: 10.1111/ajd.12186

Uter W., Bauer A., Geier J., Lessmann H., and Schnuch A. (2008): Contact allergy to textile and leather dyes. Results of information Network of Departments of Dermatology (IVDK) 2003 to 2006. Allergo Journal 17 (8), 625-630. DOI: 10.1007/BF03361952

Varma D.S., Maheswari A., Gupta V., and Varma I.K. (1980): Poly(ester-ether) fibres. Die Angewandte Makromolekulare Chemie 90 (1), 23-36. DOI: 10.1002/apmc.1980.050900103

Veena S.M., Varma I.K., and Varma D.S. (1979): Polyester fibres with improved dye uptake. Die Angewandte Makromolekulare Chemie 82 (1), 63-78. DOI: 10.1002/apmc.1979.050820106

Walsh G. and Wilkinson S.M. (2006): Materials and allergens within spectacle frames: A review. Contact Dermatitis 55 (3), 130-139. DOI: 10.1111/j.1600-0536.2006.00791.x

Warren L.J. and Marren P. (1997): Textile dermatitis and dyed maggot exposure. Contact Dermatitis 36 (2), 106. DOI: 10.1111/j.1600-0536.1997.tb00424.x

Wentworth A.B., Richardson D.M., and Davis M.D. (2012): Patch testing with textile allergens: The mayo clinic experience. Dermatitis : contact, atopic, occupational, drug 23 (6), 269-274. DOI: 10.1097/DER.0b013e318277ca3d

Wilkinson S.M. and Thomson K.F. (2000): Foot dermatitis due to non-disperse azo dyes - Short communications. Contact Dermatitis 42 (3), 162-185. DOI: 10.1034/j.1600-0536.2000.042003162.x

Wu F.-F., Chen Q.-Y., Ma X.-J., Li T.-T., Wang L.-F., Hong J., Sheng Y.-H., Ye M.-L., and Zhu Y. (2019): N-doped magnetic covalent organic frameworks for preconcentration of allergenic disperse dyes in textiles of fall protection equipment. Analytical Methods 11 (27), 3381-3387. DOI: 10.1039/C9AY00900K

ZENECA (1997): In vitro absorption of various dyes through human and pig epidermis with cover letter dated 07/08/1997. Report no. CTL/L/5926, Microfice No. OTS0001293, NTIS Issue No. 200820, date: 1997-07-08. ZENECA Central Toxicology Laboratory, Technical Report. https://ntrl.ntis.gov/NTRL/dashboard/searchResults/titleDetail/OTS0001293.xhtml

Zhou Y., Du Z., and Zhang Y. (2014): Simultaneous determination of 17 disperse dyes in textile by ultrahigh performance supercritical fluid chromatography combined with tandem mass spectrometry. Talanta 127, 108-115. DOI: 10.1016/j.talanta.2014.03.055