

**Test Report No.:** NGBEC25000784225 **Date:** Feb 20, 2025 Page 1 of 15 **(SVHC)** 

Client Name: HANGZHOU TODAYTEC DOGITAL CO.,LTD

Client Address: 600 KANGXIN RD, HANGZHOU QIANJIANG ECONOMY DEVELOPMENT ZONE

Sample Name: Thermal Transfer Ribbon

Model No.: M33

The above sample(s) and information were provided by the client.

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SGS Job No.: NBP25-000812 Sample Receiving Date: Feb 14, 2025

Testing Period: Feb 14, 2025 ~ Feb 20, 2025

Test Requested: As requested by client, SVHC in Candidate List screening is performed

according to:

(i) Two hundred and forty seven (247) substances in the Candidate List of Substances of Very High Concern (SVHC) for authorization published by European Chemicals Agency (ECHA) on and before Jan 21, 2025 regarding

Regulation (EC) No 1907/2006 concerning the REACH.

As requested by client, Potential SVHC screening is performed according to:

(i) One (1) potential Substances of Very High Concern (SVHC) in the

Identification ongoing.

(ii) Five (5) potential Substances of Very High Concern (SVHC) in the Intention List published by European Chemicals Agency (ECHA) regarding Regulation

(EC) No 1907/2006 concerning the REACH.

Test Method(s): Please refer to next page(s).

Test Result(s): Please refer to next page(s).

#### Summary:

According to the specified scope and evaluation screening, the results of 247 SVHC in the Candidate List are ≤ 0.1% (w/w) in the submitted sample.	Pass
According to the specified scope and evaluation screening, the results of 6 Potential SVHC are ≤ 0.1% (w/w) in the submitted sample.	Pass

Signed for and on behalf of

SGS-CSTC Standards Technical Services Co., Ltd. Ningbo Branch

Hayley Hong

Hayley Hong Approved Signatory





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Remark:

 The chemical analysis of specified SVHC is performed by means of currently available analytical techniques against the following SVHC related documents published by ECHA: http://echa.europa.eu/web/guest/candidate-list-table

These lists are under evaluation by ECHA and may subject to change in the future.

- 2. REACH obligation:
  - 2.1 Concerning article(s):

### Communication:

Article 33 of Regulation (EC) No 1907/2006 requires supplier of an article containing a substance meeting the criteria in Article 57 and identified in accordance with Article 59(1) in a concentration above 0.1% weight by weight (w/w) shall provide the recipient of the article with sufficient information, available to the supplier, to allow safe use of the article including, as a minimum, the name of that substance in the Candidate List.

#### Notification:

In accordance with Regulation (EC) No 1907/2006, any EU producer or importer of articles shall notify ECHA, in accordance with paragraph 4 of Article 7, if a substance meets the criteria in Article 57 and is identified in accordance with Article 59(1) of the Regulation, if (a) the substance in the Candidate List is present in those articles in quantities totaling over one tonne per producer or importer per year; and (b) the substance in the Candidate List is present in those articles above a concentration of 0.1% weight by weight (w/w).

Companies supplying articles containing substances of very high concern (SVHCs) on the Candidate List in a concentration above 0.1% weight by weight (w/w) on the EU market must comply with the Waste Framework Directive 2008/98/EC requirement and submit SCIP notifications on these articles to ECHA, as from 5 January 2021.

### 2.2 Concerning material(s):

Test results in this report are based on the tested sample. This report refers to testing result of tested sample submitted as homogenous material(s). In case such material is being used to compose an article, the results indicated in this report may not represent SVHC concentration in such article. If this report refers to testing result of composite material group by equal weight proportion, the material in each composite test group may come from more than one article.

If the sample is a substance or mixture, and it directly exports to EU, client has the obligation to comply with the supply chain communication obligation under Article 31 of Regulation (EC) No. 1907/2006 and the conditions of Authorization of substance of very high concern included in the Annex XIV of the Regulation (EC) No. 1907/2006.

### 2.3 Concerning substance and preparation:

If a SVHC is found over 0.1% (w/w) and/or the specific concentration limit which is set in Regulation (EC) No 1272/2008 and its amendments, client is suggested to prepare a Safety Data Sheet (SDS) against the SVHC to comply with the supply chain communication obligation under Regulation (EC) No 1907/2006, in which:

- a substance that is classified as hazardous under the CLP Regulation (EC) No 1272/2008.
- a mixture that is classified as hazardous under the CLP Regulation (EC) No 1272/2008, when it contains a substance with concentration equal to, or greater than the classification limit as set in Regulation (EC) No. 1272/2008; or
- a mixture is not classified as hazardous under the CLP Regulation (EC) No 1272/2008, but contains either:



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- (a) a substance posing human health or environmental hazards in an individual concentration of ≥ 1 % by weight for mixtures that are solid or liquids (i.e., non-gaseous mixtures) or ≥ 0.2 % by volume for gaseous mixtures; or
- (b) a substance that is PBT, or vPvB in an individual concentration of ≥ 0.1 % by weight for mixtures that are solid or liquids (i.e., non-gaseous mixtures); or
- (c) a substance on the SVHC candidate list (for reasons other than those listed above), in an individual concentration of ≥ 0.1 % by weight for non-gaseous mixtures; or
- (d) a substance for which there are Europe-wide workplace exposure limits
- 3. If a SVHC is found over the reporting limit, client is suggested to identify the composite component which contains the SVHC and the exact concentration of the SVHC by requesting further quantitative analysis from the laboratory.

### **Test Sample:**

**Testing Group:** 

Test Result ID	Description	Test Part ID	SGS Sample ID
001	Black film	A1	NGB25-0007842-
001	DIACK IIIIII	Ai	0001.C001

### **Test Method:**

With reference to SGS In-House method, analysis was performed by ICP-OES, UV-VIS, GC-MS, HPLC-DAD/MS and Colorimetric Method.



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### Result of SVHC in the Candidate List

Batch	Substance Name	CAS No.	001 Concentration (%)	RL (%)
-	All SVHC in Candidate list	-	ND	-

### **Result of Potential SVHC**

Batch	Substance Name	CAS No.	001 Concentration (%)	RL (%)
/	All Potential SVHC	-	ND	-

#### Notes:

- (1) The table above only shows detected SVHC, and SVHC that below RL are not reported. Please refer to Appendix for the full list of tested SVHC.
- (2) RL = Reporting Limit (Test data will be shown if it ≥ RL. RL is not regulatory limit.) ND = Not detected (lower than RL), ND is denoted on the SVHC substance.
- (3) \* The result is based on the calculation of selected element(s) under the worst-case scenario, and the evaluation of substance usage and material properties.
  - \*\* The result is based on the calculation of selected marker(s) and to the worst-case scenario.

    Calculated concentration of boric compounds are based on water extractive boron detected by ICP-OES.

    Calculated concentration of Barium diboron tetraoxide is based on water extractive boron and barium detected by ICP-OES.
  - RL = 0.005% is evaluated for element (i.e. cobalt, arsenic, lead, chromium, chromium (VI), aluminum, zirconium, boron, strontium, zinc, antimony, titanium, barium and cadmium respectively), except molybdenum RL=0.0005%, boron RL=0.0025% (only for Lead bis(tetrafluoroborate)), fluorine RL=0.050%.
- (4) § The substance is proposed for the identification as SVHC only where it contains Michler's ketone (CAS Number: 90-94-8) or Michler's base (CAS Number: 101-61-1) ≥0.1% (w/w).
- (5) / = Potential SVHC

Unless otherwise stated, the decision rule for conformity reporting is based on Binary Statement for Simple Acceptance Rule (*w*=0) stated in ILAC-G8:09/2019.



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### **Appendix**

### Full list of tested SVHC:

Batch	No.	Substance Name	CAS No.	RL (%)
I	1	4,4'-Diaminodiphenylmethane(MDA)	101-77-9	0.050
I	2	5-tert-butyl-2,4,6-trinitro-m-xylene (musk xylene)	81-15-2	0.050
1	3	Alkanes, C10-13, chloro (Short Chain Chlorinated Paraffins)	85535-84-8	0.050
I	4	Anthracene	120-12-7	0.050
l	5	Benzyl butyl phthalate (BBP)	85-68-7	0.050
I	6	Bis(2-ethylhexyl)phthalate (DEHP)	117-81-7	0.050
	7	Bis(tributyltin)oxide (TBTO)	56-35-9	0.050
	8	Cobalt dichloride*	7646-79-9	0.005
	9	Diarsenic pentaoxide*	1303-28-2	0.005
	10	Diarsenic trioxide*	1327-53-3	0.005
	11	Dibutyl phthalate (DBP)	84-74-2	0.050
I	12	Hexabromocyclododecane (HBCDD) and all major diastereoisomers identified (α-HBCDD, β-HBCDD)	-	0.050
I	13	Lead hydrogen arsenate*	7784-40-9	0.005
I	14	Sodium dichromate*	10588-01-9 /7789-12-0	0.005
I	15	Triethyl arsenate*	15606-95-8	0.005
II	16	2,4-Dinitrotoluene	121-14-2	0.050
II	17	Anthracene oil**	90640-80-5	0.050
<u>II</u>	18	Anthracene oil, anthracene paste**	90640-81-6	0.050
II	19	Anthracene oil, anthracene paste, anthracene fraction**	91995-15-2	0.050
II	20	Anthracene oil, anthracene paste, distn. Lights**	91995-17-4	0.050
II	21	Anthracene oil, anthracene-low**	90640-82-7	0.050
II	22	Diisobutyl phthalate	84-69-5	0.050
II	23	Lead chromate*	7758-97-6	0.005
II	24	Lead chromate molybdate sulphate red (C.I.  Pigment Red 104)*	12656-85-8	0.005
II	25	Lead sulfochromate yellow (C.I. Pigment Yellow 34)*	1344-37-2	0.005
[]	26	Pitch, coal tar, high temp. **	65996-93-2	0.050
П	27	Tris(2-chloroethyl)phosphate	115-96-8	0.050
II	28	Acrylamide	79-06-1	0.050
III	29	Ammonium dichromate*	7789-09-5	0.005
III	30	Boric acid*	-	0.005
Ш	31	Disodium tetraborate, anhydrous*	12179-04-3 /1303-96-4 /1330-43-4	0.005
III	32	Potassium chromate*	7789-00-6	0.005
III	33	Potassium dichromate*	7778-50-9	0.005
III	34	Sodium chromate*	7775-11-3	0.005
III	35	Tetraboron disodium heptaoxide, hydrate*	12267-73-1	0.005



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Batch	No.	Substance Name	CAS No.	RL (%)
III	36	Trichloroethylene	79-01-6	0.050
IV	37	2-Ethoxyethanol	110-80-5	0.050
IV	38	2-Methoxyethanol	109-86-4	0.050
IV	39	Chromic acid, Oligomers of chromic acid and dichromic acid, Dichromic acid*	-	0.005
IV	40	Chromium trioxide*	1333-82-0	0.005
IV	41	Cobalt(II) carbonate*	513-79-1	0.005
IV	42	Cobalt(II) diacetate*	71-48-7	0.005
IV	43	Cobalt(II) dinitrate*	10141-05-6	0.005
IV	44	Cobalt(II) sulphate*	10124-43-3	0.005
V	45	1,2,3-trichloropropane	96-18-4	0.050
V	46	1,2-Benzenedicarboxylic acid, di-C6-8- branched alkyl esters, C7-rich	71888-89-6	0.050
V	47	1,2-Benzenedicarboxylic acid, di-C7-11- branched and linear alkyl esters	68515-42-4	0.050
V	48	1-methyl-2-pyrrolidone	872-50-4	0.050
V	49	2-ethoxyethyl acetate	111-15-9	0.050
V	50	Hydrazine	302-01-2 /7803-57-8	0.050
V	51	strontium chromate*	7789-06-2	0.005
VI	52	1,2-Dichloroethane	107-06-2	0.050
VI	53	2,2'-dichloro-4,4'-methylenedianiline	101-14-4	0.050
VI	54	2-Methoxyaniline; o-Anisidine	90-04-0	0.050
VI	55	4-(1,1,3,3-tetramethylbutyl)phenol	140-66-9	0.050
VI	56	Aluminosilicate Refractory Ceramic Fibres*	-	0.005
VI	57	Arsenic acid*	7778-39-4	0.005
VI	58	Bis(2-methoxyethyl) ether	111-96-6	0.050
VI	59	Bis(2-methoxyethyl) phthalate	117-82-8	0.050
VI	60	Calcium arsenate*	7778-44-1	0.005
VI	61	Dichromium tris(chromate)*	24613-89-6	0.005
VI	62	Formaldehyde, oligomeric reaction products with aniline	25214-70-4	0.050
VI	63	Lead diazide, Lead azide*	13424-46-9	0.005
VI	64	Lead dipicrate*	6477-64-1	0.005
VI	65	Lead styphnate*	15245-44-0	0.005
VI	66	N,N-dimethylacetamide	127-19-5	0.050
VI	67	Pentazinc chromate octahydroxide*	49663-84-5	0.005
VI	68	Phenolphthalein	77-09-8	0.050
VI	69	Potassium hydroxyoctaoxodizincatedichromate*	11103-86-9	0.005
VI	70	Trilead diarsenate*	3687-31-8	0.005
VI	71	Zirconia Aluminosilicate Refractory Ceramic Fibres*	-	0.005
VII	72	[4-[[4-anilino-1-naphthyl][4- (dimethylamino)phenyl]methylene]cyclohexa- 2,5-dien-1-ylidene] dimethylammonium chloride (C.I. Basic Blue 26)§	2580-56-5	0.050



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I-I-I.4.4'-bis(dimethylamino)	Batch	No.	Substance Name	CAS No.	RL (%)
VII   73					· /
VII	\///	70		E40.00.0	0.050
VII         74         1,2-bis(2-methoxyethoxy)ethane (TEGDME; triglyme)         112-49-2         0.050           VII         75         1,2-dimethoxyethane; ethylene glycol dimethyl ether (EGDME)         110-71-4         0.050           VII         76         4,4'-bis(dimethylamino)-4'-(methylamino)trityl alcohols         561-41-1         0.050           VII         77         4,4'-bis(dimethylamino)-4'-(methylamino)trityl alcohols         561-41-1         0.050           VII         78         Diboron trioxide*         1303-86-2         0.005           VII         79         Formamide         75-12-7         0.050           VII         80         Lead(II) bis(methanesulfonate)*         17570-76-2         0.005           VII         81         N,N,N,N'-tetramethyl-4,4'-methylenedianiline (Michler's base)         101-61-1         0.050           VII         82         TGIC (1,3,5-tris(oxiranylmethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)         2451-62-9         0.050           VII         83         (phenylamino)naphthalene-1-methanol (C.I.         6786-83-0         0.050           VIII         84         (phenylamino)naphthalene-1-methanol (C.I.         6786-83-0         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.050	VII	/3		548-62-9	0.050
VII   75					
VII   75   1,2-dimethoxyethane; ethylene glycol dimethylether (EGDME)   110-71-4   0.050     VII   76	1/11	7/	1,2-bis(2-methoxyethoxy)ethane (TEGDME;	112 40 2	0.050
VII   76	VII	74		112-49-2	0.050
VII   76	\/II	75		110-71-4	0.050
VII   77	VII	73		110-71-4	0.030
VII   77	VII	76		90-94-8	0.050
VII   78	V	, ,		00010	0.000
Acconois	VII	77		561-41-1	0.050
VII         79         Formamide         75-12-7         0.050           VII         80         Lead(II) bis(methanesulfonate)*         17570-76-2         0.005           VII         81         N,N,N',N'-tetramethyl-4,4'-methylenedianiline (Michler's base)         101-61-1         0.050           VII         82         TGIC (1,3,5-tris(oxiranylmethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)         2451-62-9         0.050           VII         83         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         6786-83-0         0.050           VII         84         epoxypropyl]-1,3,5-triszine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2)-Jidioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3- oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050					
VII         80         Lead(II) bis(methanesulfonate)*         17570-76-2         0.005           VII         81         N,N,N',N'-tetramethyl-4,4'-methylenedianiline (Michler's base)         101-61-1         0.050           VII         82         TGIC (1,3,5-tris(oxiranylmethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)         2451-62-9         0.050           VII         83         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         6786-83-0         0.050           VII         84         epoxypropyl]-1,3,5-triszine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3- oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Oxydianiline and its salts         101-80-4 <td< td=""><td></td><td></td><td></td><td></td><td></td></td<>					
VII         81         N,N,N,N'-retramethyl-4,4'-methylenedianiline (Michler's base)         101-61-1         0.050           VII         82         TGIC (1,3,5-tris(oxiranylmethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)         2451-62-9         0.050           VII         83         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         6786-83-0         0.050           VII         84         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         59653-74-6         0.050           VIII         84         epoxypropyl]-1,3,5-triazine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.050           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts </td <td></td> <td></td> <td></td> <td></td> <td></td>					
VII   82   TGIC (1,3,5-tris(oxiranylmethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)   2451-62-9   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.	VII	80		17570-76-2	0.005
VIII   82   TGIC (1,3,5-tris(coxiranyImethyl)-1,3,5-triazine-2,4,6(1H,3H,5H)-trione)   2451-62-9   0.050	VII	81		101-61-1	0.050
VII         82         2,4,6(1H,3H,5H)-trione)         2451-62-9         0.050           VII         83         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         6786-83-0         0.050           VII         84         p-TGIC (1,3,5-trisz[(2S and 2R)-2,3-epoxypropyl]-1,3,5-triazine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         96         <		-			
C,q,6(1H,SH,of)-itrione)   C,q,c=Bis[4-(dimethylamino)phenyl]-4   Cphenylamino)naphthalene-1-methanol (C.I.   6786-83-0   0.050   Solvent Blue 4) §   F-TGIC (1,3,5-tris[(2S and 2R)-2,3-epoxypropyl]-1,3,5-tris[arine-2,4,6-(1H,3H,5H)-trione)   Signature   Si	VII	82		2451-62-9	0.050
VII         83         (phenylamino)naphthalene-1-methanol (C.I. Solvent Blue 4) §         6-TGIC (1,3,5-tris[(2S and 2R)-2,3-trione-2,4,6-(1H,3H,5H)-trione)         6-TGIC (1,3,5-tris2ine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         84         epoxypropyl]-1,3,5-triazine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-ph		_			
Solvent Blue 4) §   β-TGIC (1,3,5-tris[(2S and 2R)-2,3-trione)   Post-strione	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	00		0700 00 0	0.050
VIII   84   B-TGIC (1,3,5-tris[(2S and 2R)-2,3-epoxypropyl]-1,3,5-triazine-2,4,6-(1H,3H,5H)-trione)   59653-74-6   0.050   1.2-Benzenedicarboxylic acid, dipentylester, branched and linear   84777-06-0   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050   0.050	VII	83	\(\(\)\(\)	6/86-83-0	0.050
VII         84         epoxypropyl]-1,3,5-triazine-2,4,6-(1H,3H,5H)-trione)         59653-74-6         0.050           VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3- oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120					
VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.	1/11	0.4		E0652 74 6	0.050
VIII         85         [Phthalato(2-)]dioxotrilead*         69011-06-9         0.005           VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4 <td< td=""><td>VII</td><td>04</td><td></td><td>39033-74-0</td><td>0.050</td></td<>	VII	04		39033-74-0	0.050
VIII         86         1,2-Benzenedicarboxylic acid, dipentylester, branched and linear         84777-06-0         0.050           VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050	\/III	85		60011-06-0	0.005
VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, ciscyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anh		00		09011-00-9	0.005
VIII         87         1,2-Diethoxyethane         629-14-1         0.050           VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         100         Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic an	VIII	86		84777-06-0	0.050
VIII         88         1-Bromopropane         106-94-5         0.050           VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         - </td <td>VIII</td> <td>87</td> <td></td> <td>629-14-1</td> <td>0.050</td>	VIII	87		629-14-1	0.050
VIII         89         3-Ethyl-2-methyl-2-(3-methylbutyl)-1,3-oxazolidine         143860-04-2         0.050           VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride					
VIII         99         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050					
VIII         90         4-(1,1,3,3-tetramethylbutyl)phenol, ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         -         0.050	VIII	89		143860-04-2	0.050
VIII         90         ethoxylated         -         0.050           VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, cis-trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         -         0.050					
VIII         91         4,4'-Methylenedi-o-toluidine         838-88-0         0.050           VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, cis-trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxamide (C,C'-         123-77-3         0.050	VIII	90		-	0.050
VIII         92         4,4'-Oxydianiline and its salts         101-80-4         0.050           VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, cis-trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         -         0.050	VIII	91		838-88-0	0.050
VIII         93         4-Aminoazobenzene         60-09-3         0.050           VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           VIII         100         cyclohexane-1,2-dicarboxylic anhydride, cis-trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         123-77-3         0.050	VIII	92			
VIII         94         4-Methyl-m-phenylenediamine         95-80-7         0.050           VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         -         0.050					
VIII         95         4-Nonylphenol, branched and linear         -         0.050           VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxylic anhydride         -         0.050					
VIII         96         6-Methoxy-m-toluidine         120-71-8         0.050           VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxamide (C,C'-approximate)         123-77-3         0.050	VIII	95	4-Nonylphenol, branched and linear	-	0.050
VIII         97         Acetic acid, lead salt, basic*         51404-69-4         0.005           VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxamide (C,C'-approximate)         123-77-3         0.050	VIII	96		120-71-8	0.050
VIII         98         Biphenyl-4-ylamine         92-67-1         0.050           VIII         99         Decabromodiphenyl ether (DecaBDE)         1163-19-5         0.050           Cyclohexane-1,2-dicarboxylic anhydride, cis-cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride         -         0.050           VIII         101         Diazene-1,2-dicarboxamide (C,C'-approximate)         123-77-3         0.050	VIII	97			0.005
VIII 100 Cyclohexane-1,2-dicarboxylic anhydride, cis- cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride Diazene-1,2-dicarboxamide (C,C'- 123-77-3 0.050	VIII	98		92-67-1	0.050
VIII 100 cyclohexane-1,2-dicarboxylic anhydride, trans-cyclohexane-1,2-dicarboxylic anhydride  Diazene-1,2-dicarboxamide (C,C'-	VIII	99	Decabromodiphenyl ether (DecaBDE)	1163-19-5	0.050
trans-cyclohexane-1,2-dicarboxylic anhydride  Diazene-1,2-dicarboxamide (C,C'-			Cyclohexane-1,2-dicarboxylic anhydride, cis-		
VIII 101 Diazene-1,2-dicarboxamide (C,C'-	VIII	100		-	0.050
azodi(formamide))	\/!!!	101		123-77-3	0.050
	VIII	101	azodi(formamide))	120-11-0	0.000



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Batch	No.	Substance Name	CAS No.	RL (%)
VIII	102	Dibutyltin dichloride (DBTC)	683-18-1	0.050
VIII	103	Diethyl sulphate	64-67-5	0.050
VIII	104	Diisopentylphthalate	605-50-5	0.050
VIII	105	Dimethyl sulphate	77-78-1	0.050
VIII	106	Dinoseb	88-85-7	0.050
VIII	107	Dioxobis(stearato)trilead*	12578-12-0	0.005
VIII	108	Fatty acids, C16-18, lead salts*	91031-62-8	0.005
VIII	109	Furan	110-00-9	0.050
VIII	110	Henicosafluoroundecanoic acid	2058-94-8	0.050
VIII	111	Heptacosafluorotetradecanoic acid	376-06-7	0.050
		Hexahydromethylphthalic anhydride,		
VIII	112	Hexahydro-4-methylphthalic anhydride,		0.050
VIII	112	Hexahydro-1-methylphthalic anhydride,	-	0.050
		Hexahydro-3-methylphthalic anhydride		
VIII	113	Lead bis(tetrafluoroborate)*	13814-96-5	0.005
VIII	114	Lead cyanamidate*	20837-86-9	0.005
VIII	115	Lead dinitrate*	10099-74-8	0.005
VIII	116	Lead monoxide*	1317-36-8	0.005
VIII	117	Lead oxide sulfate*	12036-76-9	0.005
VIII	118	Lead tetroxide (orange lead)*	1314-41-6	0.005
VIII	119	Lead titanium trioxide*	12060-00-3	0.005
VIII	120	Lead titanium zirconium oxide*	12626-81-2	0.005
VIII	121	Methoxyacetic acid	625-45-6	0.050
VIII	122	Methyloxirane (Propylene oxide)	75-56-9	0.050
VIII	123	N,N-Dimethylformamide	68-12-2	0.050
VIII	124	N-Methylacetamide	79-16-3	0.050
VIII	125	N-Pentyl-isopentylphthalate	776297-69-9	0.050
VIII	126	o-Aminoazotoluene	97-56-3	0.050
VIII	127	o-Toluidine	95-53-4	0.050
VIII	128	Pentacosafluorotridecanoic acid	72629-94-8	0.050
VIII	129	Pentalead tetraoxide sulphate*	12065-90-6	0.005
VIII	130	Pyrochlore, antimony lead yellow*	8012-00-8	0.005
VIII	131	Silicic acid, barium salt, lead-doped*	68784-75-8	0.005
VIII	132	Silicic acid, lead salt*	11120-22-2	0.005
VIII	133	Sulfurous acid, lead salt, dibasic*	62229-08-7	0.005
VIII	134	Tetraethyllead*	78-00-2	0.005
VIII	135	Tetralead trioxide sulphate*	12202-17-4	0.005
VIII	136	Tricosafluorododecanoic acid	307-55-1	0.050
VIII	137	Trilead bis(carbonate)dihydroxide (basic lead carbonate)*	1319-46-6	0.005
VIII	138	Trilead dioxide phosphonate*	12141-20-7	0.005
IX	139	4-Nonylphenol, branched and linear, ethoxylated	-	0.050
IX	140	Ammonium pentadecafluorooctanoate (APFO)**	3825-26-1	0.050
IX	141	Cadmium oxide*	1306-19-0	0.005
IX	142	Cadmium	7440-43-9	0.005



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Batch	No.	Substance Name	CAS No.	RL (%)
IX	143	Dipentyl phthalate (DPP)	131-18-0	0.050
IX	144	Pentadecafluorooctanoic acid (PFOA)	335-67-1	0.050
X	145	Cadmium sulphide*	1306-23-6	0.005
Х	146	Dihexyl phthalate	84-75-3	0.050
Х	147	Disodium 3,3'-[[1,1'-biphenyl]-4,4'- diylbis(azo)]bis(4-aminonaphthalene-1- sulphonate) (C.I. Direct Red 28)	573-58-0	0.050
х	148	Disodium 4-amino-3-[[4'-[(2,4-diaminophenyl)azo][1,1'-biphenyl]-4-yl]azo] -5-hydroxy-6-(phenylazo)naphthalene-2,7-disulphonate (C.I. Direct Black 38)	1937-37-7	0.050
X	149	Imidazolidine-2-thione; (2-imidazoline-2-thiol)	96-45-7	0.050
X	150	Lead di(acetate)*	301-04-2	0.005
X	151	Trixylyl phosphate	25155-23-1	0.050
ΧI	152	1,2-Benzenedicarboxylic acid, dihexyl ester, branched and linear	68515-50-4	0.050
ΧI	153	Cadmium chloride*	10108-64-2	0.005
XI	154	Sodium perborate; perboric acid, sodium salt*	-	0.005
ΧI	155	Sodium peroxometaborate*	7632-04-4	0.005
XII	156	2-(2H-benzotriazol-2-yl)-4,6-ditertpentylphenol (UV-328)	25973-55-1	0.050
XII	157	2-benzotriazol-2-yl-4,6-di-tert-butylphenol (UV-320)	3846-71-7	0.050
XII	158	2-ethylhexyl 10-ethyl-4,4-dioctyl-7-oxo-8-oxa- 3,5-dithia-4-stannatetradecanoate (DOTE)	15571-58-1	0.050
XII	159	Cadmium fluoride*	7790-79-6	0.005
XII	160	Cadmium sulphate*	10124-36-4 /31119-53-6	0.005
XII	161	Reaction mass of 2-ethylhexyl 10-ethyl-4,4-dioctyl-7-oxo-8-oxa-3,5-dithia-4-stannatetradecanoate & 2-ethylhexyl 10-ethyl-4-[[2-[(2-ethylhexyl)oxy]-2-oxoethyl]thio]-4-octyl-7-oxo-8-oxa-3,5-dithia-4-stannatetradecanoate (reaction mass of DOTE & MOTE)	-	0.050
XIII	162	1,2-benzenedicarboxylic acid, di-C6-10-alkyl esters; 1,2-benzenedicarboxylic acid, mixed decyl and hexyl and octyl diesters with ≥ 0.3% of dihexyl phthalate	-	0.050
XIII	163	5-sec-butyl-2-(2,4-dimethylcyclohex-3-en-1-yl)-5-methyl-1,3-dioxane [1], 5-sec-butyl-2-(4,6-dimethylcyclohex-3-en-1-yl)-5-methyl-1,3-dioxane [2] [covering any of the individual isomers of [1] and [2] or any combination thereof]	-	0.050
XIV	164	1,3-propanesultone	1120-71-4	0.050
XIV	165	2,4-di-tert-butyl-6-(5-chlorobenzotriazol-2-yl) phenol (UV-327)	3864-99-1	0.050



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Batch	No.	Substance Name	CAS No.	RL (%)
XIV	166	2-(2H-benzotriazol-2-yl)-4-(tert-butyl)-6-(sec-	36437-37-3	0.050
XIV	167	butyl) phenol (UV-350) Nitrobenzene	98-95-3	0.050
		Perfluorononan-1-oic-acid and its sodium and	90-90-0	
XIV	168	ammonium salts	-	0.050
XV	169	Benzo[def]chrysene (Benzo[a]pyrene)	50-32-8	0.050
XVI	170	4,4'-isopropylidenediphenol (bisphenol A)	80-05-7	0.050
XVI	171	4-Heptylphenol, branched and linear	-	0.050
XVI	172	Nonadecafluorodecanoic acid (PFDA) and its sodium and ammonium salts	-	0.050
XVI	173	p-(1,1-dimethylpropyl)phenol	80-46-6	0.050
XVII	174	Perfluorohexane-1-sulphonic acid and its salts	-	0.050
7,711	.,,	1,6,7,8,9,14,15,16,17,17,18,18-		0.000
		Dodecachloropentacyclo[12.2.1.16,9.02,13.05		
XVIII	175	,10]octadeca-7,15-diene ("Dechlorane	-	0.050
		Plus"™) [covering any of its individual anti-		
		and syn-isomers or any combination thereof]		
XVIII	176	Benz[a]anthracene	56-55-3	0.050
XVIII	177	Cadmium nitrate*	10325-94-7	0.005
XVIII	178	Cadmium carbonate*	513-78-0	0.005
XVIII	179	Cadmium hydroxide*	21041-95-2	0.005
XVIII	180	Chrysene	218-01-9	0.050
		Reaction products of 1,3,4-thiadiazolidine-2,5-		
XVIII	181	dithione, formaldehyde and 4-heptylphenol,		0.050
AVIII	101	branched and linear (RP-HP) [with ≥0.1% w/w	-	0.050
		4-heptylphenol, branched and linear]		
XIX	182	Benzene-1,2,4-tricarboxylic acid 1,2 anhydride	552-30-7	0.050
		(trimellitic anhydride) (TMA)		
XIX	183	Benzo[ghi]perylene	191-24-2	0.050
XIX	184	Decamethylcyclopentasiloxane (D5)	541-02-6	0.050
XIX	185	Dicyclohexyl phthalate (DCHP)	84-61-7	0.050
XIX	186	Disodium octaborate*	12008-41-2	0.005
XIX	187	Dodecamethylcyclohexasiloxane (D6)	540-97-6	0.050
XIX	188	Ethylenediamine (EDA)	107-15-3	0.050
XIX	189	Lead	7439-92-1	0.005
XIX	190	Octamethylcyclotetrasiloxane (D4)	556-67-2	0.050
XIX	191	Terphenyl, hydrogenated	61788-32-7	0.050
		1,7,7-trimethyl-3-		
XX	192	(phenylmethylene)bicyclo[2.2.1]heptan-2-one	15087-24-8	0.050
		(3-benzylidene camphor)		
XX	193	2,2-bis(4'-hydroxyphenyl)-4-methylpentane	6807-17-6	0.050
XX	194	Benzo[k]fluoranthene	207-08-9	0.050
XX	195	Fluoranthene	206-44-0	0.050
XX	196	Phenanthrene	85-01-8	0.050
XX	197	Pyrene	129-00-0	0.050
XXI	198	2,3,3,3-tetrafluoro-2- (heptafluoropropoxy)propionic acid, its salts	-	0.050
		(Hoptandoropropoxy)propionic acid, its saits		



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Batch	No.	Substance Name	CAS No.	RL (%)
		and its acyl halides (covering any of their		
		individual isomers and combinations thereof)		
XXI	199	2-methoxyethyl acetate	110-49-6	0.050
XXI	200	4-tert-butylphenol (PTBP)	98-54-4	0.050
XXI	201	Tris(4-nonylphenyl, branched and linear) phosphite (TNPP)	-	0.050
		2-benzyl-2-dimethylamino-4'-		
XXII	202	morpholinobutyrophenone	119313-12-1	0.050
		2-methyl-1-(4-methylthiophenyl)-2-		
XXII	203	morpholinopropan-1-one	71868-10-5	0.050
XXII	204	Diisohexyl phthalate	71850-09-4	0.050
XXII	205	Perfluorobutane sulfonic acid (PFBS) and its salts	-	0.050
XXIII	206	1-vinylimidazole	1072-63-5	0.050
XXIII	207	2-methylimidazole	693-98-1	0.050
XXIII	208	Butyl 4-hydroxybenzoate	94-26-8	0.050
XXIII	209	Dibutylbis(pentane-2,4-dionato-O,O')tin**	22673-19-4	0.050
XXIV	210	bis(2-(2-methoxyethoxy)ethyl) ether	143-24-8	0.050
		Dioctyltin dilaurate, stannane, dioctyl-,		
XXIV	211	bis(coco acyloxy) derivs., and any other stannane, dioctyl-, bis(fatty acyloxy) derivs. wherein C12 is the predominant carbon number of the fatty acyloxy moiety**	-	0.050
XXV	212	1,4-Dioxane	123-91-1	0.050
XXV	213	2,2-bis(bromomethyl)propane1,3-diol (BMP); 2,2-dimethylpropan-1-ol, tribromo derivative/3- bromo-2,2-bis(bromomethyl)-1-propanol (TBNPA); 2,3-dibromo-1-propanol (2,3-DBPA)	-	0.050
XXV	214	2-(4-tert-butylbenzyl)propionaldehyde and its individual stereoisomers	-	0.050
XXV	215	4,4'-(1-methylpropylidene)bisphenol; (bisphenol B)	77-40-7	0.050
XXV	216	Glutaral	111-30-8	0.050
XXV	217	Medium-chain chlorinated paraffins (MCCP) [UVCB substances consisting of more than or equal to 80% linear chloroalkanes with carbon chain lengths within the range from C14 to C17]	-	0.050
XXV	218	Orthoboric acid, sodium salt*	13840-56-7	0.005
XXV	219	Phenol, alkylation products (mainly in para position) with C12-rich branched or linear alkyl chains from oligomerisation, covering any individual isomers and/ or combinations thereof (PDDP)	-	0.050
XXVI	220	(±)-1,7,7-trimethyl-3-[(4-methylphenyl)methylene]bicyclo[2.2.1]heptan-2-one covering any of the individual isomers and/or combinations thereof (4-MBC)	-	0.050



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Batch	No.	Substance Name	CAS No.	RL (%)
XXVI	221	6,6'-di-tert-butyl-2,2'-methylenedi-p-cresol	119-47-1	0.050
		(DBMC) S-(tricyclo[5.2.1.0'2,6]deca-3-en-8(or 9)-yl) O-		
		(isopropyl or isobutyl or 2-ethylhexyl) O-		
XXVI	222	(isopropyl or isobutyl or 2-ethylhexyl)	255881-94-8	0.050
		phosphorodithioate		
XXVI	223	Tris(2-methoxyethoxy)vinylsilane	1067-53-4	0.050
XXVII	224	N-(hydroxymethyl)acrylamide	924-42-5	0.050
XXVIII	225	1,1'-[ethane-1,2-diylbisoxy]bis[2,4,6-	37853-59-1	0.050
7(7( ) 111	220	tribromobenzene]	07000 00 1	0.000
XXVIII	226	2,2',6,6'-tetrabromo-4,4'-	79-94-7	0.050
		isopropylidenediphenol		
XXVIII	227	4,4'-sulphonyldiphenol	80-09-1	0.050
XXVIII	228	Barium diboron tetraoxide*	13701-59-2	0.005
XXVIII	229	Bis(2-ethylhexyl) tetrabromophthalate covering any of the individual isomers and/or	_	0.050
AA VIII	229	combinations thereof	-	0.030
XXVIII	230	Isobutyl 4-hydroxybenzoate	4247-02-3	0.050
XXVIII	231	Melamine	108-78-1	0.050
XXVIII	232	Perfluoroheptanoic acid and its salts		0.050
		reaction mass of 2,2,3,3,5,5,6,6-octafluoro-4-		
XXVIII	233	(1,1,1,2,3,3,3-heptafluoropropan-2-		0.050
AAVIII	233	yl)morpholine and 2,2,3,3,5,5,6,6-octafluoro-4-	-	0.050
		(heptafluoropropyl)morpholine*		
XXIX	234	Bis(4-chlorophenyl) sulphone	80-07-9	0.050
XXIX	235	Diphenyl(2,4,6-trimethylbenzoyl)phosphine	75980-60-8	0.050
XXX	236	oxide 2,4,6-tri-tert-butylphenol	732-26-3	0.050
		2-(2H-benzotriazol-2-yl)-4-(1,1,3,3-		
XXX	237	tetramethylbutyl)phenol (UV-329)	3147-75-9	0.050
XXX	238	2-(dimethylamino)-2-[(4-methylphenyl)methyl]-	119344-86-4	0.050
		1-[4-(morpholin-4-yl)phenyl]butan-1-one		0.050
XXX	239	Bumetrizole (UV-326)	3896-11-5	0.050
XXX	240	Oligomerisation and alkylation reaction	-	0.050
		products of 2-phenylpropene and phenol	00.40.0	
XXXI	241	Bis(α,α-dimethylbenzyl) peroxide	80-43-3	0.050
XXXI	242	Triphenyl phosphate 6-[(C10-C13)-alkyl-(branched, unsaturated)-	115-86-6	0.050
XXXII	243	2,5-dioxopyrrolidin-1-yl]hexanoic acid	2156592-54-8	0.050
XXXII	244	O,O,O-triphenyl phosphorothioate	597-82-0	0.050
XXXII	245	Octamethyltrisiloxane	107-51-7	0.050
XXXII	246	Perfluamine	338-83-0	0.050
XXXII	247	Reaction mass of: triphenylthiophosphate and	192268-65-8	0.050
ΛΛΛΙΙ		tertiary butylated phenyl derivatives		
/	248	Resorcinol	108-46-3	0.050
/	249	1,1,1,3,5,5,5-heptamethyl-3- [(trimethylsilyl)oxy]trisiloxane	17928-28-8	0.050
/	250	Decamethyltetrasiloxane	141-62-8	0.050



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Batch	No.	Substance Name	CAS No.	RL (%)
/	251	Dodecamethylpentasiloxane	141-63-9	0.050
/	252	Hexamethyldisiloxane	107-46-0	0.050
/	253	tetra(sodium/potassium) 7-[(E)-{2-acetamido- 4-[(E)-(4-{[4-chloro-6-({2-[(4-fluoro-6-{[4- (vinylsulfonyl)phenyl]amino}-1,3,5-triazine-2- yl)amino]propyl}amino)-1,3,5-triazine-2- yl]amino}-5-sulfonato-1-naphthyl)diazenyl]-5- methoxyphenyl}diazenyl]-1,3,6- naphthalenetrisulfonate; Reactive Brown 51	-	0.050



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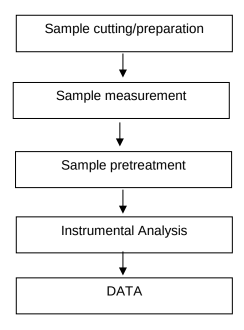
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**Test Report** (SVHC) **ATTACHMENTS** 

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## **Testing Flow Chart**





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Sample photos:



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