Section 1: General Form:

Instructions for Section 1: see page 2

Please provide:			
As such, the prompt and accur is treated as critical base data f All information will be treated	ate completion of all section of safety and regulatory we as confidential.	proval and qualification of potential suppliers. In swill facilitate this process and the data on the form. In not applicable, please indicate and explain.	rm
A. Material /Supplier Identifica	ition:		
Trade Name/Product #: AquaS	Style TM SH-100 polymer Harmonized Tariff (Code:	
Color Index Number (if applica	ble): Not Applicable		
Manufacturer Name (if different ASHLAND	t from Supplier): ISP FREET	TOWN FINE CHEMICALS INC. AFFILIATE OF	
Supplier Name: Ashland			
Contact Name: Global Regulat	ory Customer Request Tear	m Title: Ana	lyst
E-mail Address: globalregulate	orycustomerrequest@ashlar	nd.com	
Phone: NA		FAX: NA	
Manufacturer's Plant Address (Address where material is manuf	factured): Refer to CoA	Country of Origin: USA	
B. Required Supporting Docum	entation:		
Material Safety Data Sheet SDS Microbiological Specification Specification Sheet Certificate of Analysis (COA) GMP Certificate BSE Certificate			
C. Applicable Sections: Please	note if any data is restricted	d under confidentiality agreement (Mandatory in re	d).
General Form Regulatory Status Physical Chemical Properties	Restricted Data Restricted Data		

Environmental Impact statement (if available) with this completed questionnaire.

Section 1: General Form

D. Composition & Material Identification:

Total target contents of the components should sum up to 100%. List all intentionally-added constituents in this table. Please be noted the data here is critical base data for safety and regulatory work.

* Definition of Function:

Key Ingredients - defined as chemicals intended to function in finished products.

Carryover Ingredients – defined as chemicals intentionally added to material to maintain its quality or stability, such as preservative, anti-oxidant, UV filter, that does not function in finished product formula.

#	Constituent name (Approved INCI name)	CAS#	EINECS# / ELINCS#	Min (W %)	Target (W %)	Max (W %)	Constituent Function	Feedstock Origin
1	Aqua	7732-18-5	231-791-2	68		72	Solvent	Mineral
2	Acrylates Copolymer	7732-18-5	231-791-2	28		32	Key Ingredient	Synthetic
3	Sorbic Acid	110-44-1	203-768-7		<0.1	15	Additive-Preservative	Synthetic
4							Select	Select
5							Select	Select
6							Select	Select
7							Select	Select
8							Select	Select
9							Select	Select
10							Select	Select

General Comments:

If you selected the Feedstock Origin "Bovine" or "Animal-other", please attach TSE certification.

If you select the Feedstock Origin "Other", please explain:

Is the material a polymer according to the definition (Cefic) Select

Please note if the material is a polymer a detailed impurity profile should be provided in following table G.

Section 1: General Form

Section 1: General Form

F. Global Inventory Status: In the table below, please describe the inventory status of each constituent. Please add in the Inventory Number or choose the Inventory Status in each field below:

#	Constituent name	US	Canada	Australia	China	Jaj	pan	Korea	Philippines
	(Approved INCI name)	TSCA	Inventory	AICS	IECSC	ENCS	JCIA	ECL	PICCS
1	Aqua	Listed	DSL	Listed	Listed	Listed	Select	Listed	Listed
2	Acryaltes Copolymer	Listed	Listed	Listed	Listed	Listed	Select	Listed	Listed
3	Sorbic Acid	Listed	Listed	Listed	Listed	Listed	Select	Listed	Listed
4		Listed	Listed	Listed	Listed	Listed	Select	Listed	Listed
5		Listed	Listed	Listed	Listed	Listed	Select	Listed	Listed
6		Select	Select	Select	Select	Select	Select	Select	Select
7		Select	Select	Select	Select	Select	Select	Select	Select
8		Select	Select	Select	Select	Select	Select	Select	Select
9		Select	Select	Select	Select	Select	Select	Select	Select
10		Select	Select	Select	Select	Select	Select	Select	Select

General Comments/Notes:

G. Impurity Profile: Please refer to General Form Instructions for explanation.

Impurities/Residues/Catalysts/Monomers/By-Products An impurity is any chemical substance that is unintentionally present. List all impurities (regardless of amount).	CAS#	Max Level in RM	% or ppm or ppb	Comments
Methacrylic acide	79-41-4	≤100	ppm	Residual Monomer
Ethyl Acrylate	140-88- 5	≤400	ppm	Resdiual Monomer
Sodium Lauryl Sulphate	68585- 47-7	≤0.5%	ppm	Residue
				Refer to residual solvent statement and Heavy Metal statement

Section 2: Regulatory Status: Instructions for Section 2: see page 4

Instructions: Provide any known restrictions on this material for use in cosmetic products: For example, restrictions relating to UV Filters, colorants or preservatives or other restrictions noted in Annex III of the Cosmetic Directive, CIR. or Canadian Hot List

A. Restrictions:

Country	Restriction
	(% and approved use if any)
US	Sorbic Acid and Acrylates Copolymer -
	CIR reviewed
Canada	
EU	Listed in Annex V of List of
	Preservatives allowed in Cosmetic
	Products - Maximum concentration in
	ready for use preparation: 0.6% (acid)
Australia	
China	Aqua, Acrylates Copolymer and Sorbic
	Acid - Listed in IECIC CFDA list 2015
Japan	
Korea	
Other:	

B. Compliance:

Compliance with Regulations:	Yes or No?	Description
Are <u>any</u> of the ingredients a known Carcinogen, Mutagen or Reproductive Toxicant as defined under European Regulation (EC) No. 1272/2008?	Select	Please refer to the CMR statement.
Does the raw material as supplied contain any ingredients or impurities identified on the Substances of Very High Concern List under the REACH (NUMBER)	Select	Please refer to Positioning Letter on SVHC and REACH letter
Are all of the ingredients compliant with the European Regulation (EC) No 1272/2008 on classification, labelling and packaging of substances and mixtures?	Yes	
Are all of the ingredients in compliance with the European Cosmetic Regulation (EC) No. 1223/2009?	Yes	
Are any of the ingredients or impurities listed by California Proposition 65?	Yes	Ethyl Acrylate listed as Cal prop 65
Are any of the ingredients declarable under the California Safe Cosmetics Act (SB 484)?	No	
Are any of the ingredients or impurities found to be Carcinogenic by NTP?	Select	Refer to US SDS section 11
Are any of the ingredients or impurities found to be Carcinogenic by IARC?	Select	Refer to US SDS section 11

**Please attach the REACH status and summary (including any components that may require authorization) for the material.

Animal Testing:	Yes or	Description
	No?	
Are you in compliance with the European Cosmetic Regulation (EC) No.	Yes	
1223/2009 prohibiting animal testing after a certain date?		
Has your company conducted or commissioned any animal testing on this	Select	Please refer to the Animal Testing
raw material, or are you aware of any such animal testing conducted by a		Statement.
third party? Please specify most recent testing date.		
If testing was conducted post September 11, 2004, please specify date;		fer to the Animal Testing Statement.
describe purpose and endpoints for the test(s).		

Volatile Organic Content	Description

Are any of this material's ingredients classified as a VOC?	No
Total Raw Material VOC content in % (wt/wt) HVOC	
MVOC	
LVOC	
VOC Exempt?	

Section 3: Physical Chemical Properties

Instructions for Section 3: see page 4

A. Physical Form:

The material is supplied as Liquid

The physical parameters reported here are applicable to the material as it is supplied. Select

If no is chosen, explain: Refer to Safety Data Sheet

B. Physical Constants:

		Additional				
Physical Parameter	Value	Information				
Molecular Weight*		Select				
Specific Gravity						
Partition Coefficient (Kow)						
Melting Point	°C	Select				
Boiling Point	°C	Select				
Freezing Point	°C	Select				
Flash Point	°C	Select				
Minimum Ignition Temperature	°C	Select				
Spectral Data	(As applicable)					
UV/Visible	Select					
IR	Select					
Mass	Select					
Fluorescence	Select					
NMR	Select					
Separation/	Separation/ Analysis Data					
GC	Select					
HPLC	Select					
HPTLC	Select					

^{*}Average molecular weight for key component.

Please indicate SMILES notation for the key component:

Note: If physical property data is included in the attached MSDS, check here: \Box

Section 4. Human Safety

Instructions for Section 4: see page 5

Please supply robust summaries of the tests performed as an attachment.

The testing identified here was completed on the material as it is supplied. **Select**

If the answer is "no," please explain. Refer Section 11.0 of SDS

Please specify when *in vitro* models are used.

A. Testing Summary:

Testing Summary.	Protocol	Date	Result
Acute Oral Toxicity			
Sub chronic Toxicity (28 / 90 day test)			
Dermal Irritation			
Dermal Sensitization			
Dermal Absorption			
Mucous Membrane Irritation (Eye)			
Mutagenicity (Ames)			
Genotoxicity			
Carcinogenicity			
Inhalation Toxicity			
Reproductive Toxicity			
Comedogenicity			
Photo-irritation			
Photo-toxicity			
Photosensitization			
Other:			

General Comments/Notes: Refer to SDS. Tox summary will be provided upon separate request

B. Materials of Concern:

Material(s) of Concern	CAS#	ALL SOURCES: Total Inclusion Level %	Added Ingredient Contribution Inclusion Level %	Natural Source Contribution Inclusion Level %
Please refer to the impurity profile section 'G'.	-			

C. Declarable Allergens:

Regulated Allergens	CAS#	Does not contain	ALL SOURCES Total Inclusion Level %	Added Ingredient Contribution Inclusion Level %	Natural Source Contribution Inclusion Level %
Alpha-Isomethyl Ionone	127-51-5				
Amyl cinnamal	122-40-7				
Amyl cinnamyl alcohol	101-85-9				
Anise alcohol	105-13-5				
Benzyl alcohol	100-51-6				
Benzyl benzoate	120-51-4				
Benzyl cinnamate	103-41-3				
Benzyl salicylate	118-58-1				
Butylphenyl methylpropional	80-54-6				
Cinnamal	104-55-2				
Cinnamyl alcohol	104-54-1				
Citral	5392-40-5				
Citronellol	106-22-9				
Coumarin	91-64-5				
Eugenol	97-53-0				
Isoeugenol	97-54-1				
Evernia Furfuracea	90028-67-4				
(Treemoss) Extract					
Evernia Prunastri (Oakmoss)	90028-68-5				
Extract					
Farnesol	4602-84-0				
Geraniol	106-24-1				
Hexyl cinnamal	101-86-0				
Hydroxycitronellal	107-75-5				
Hydroxyisihexyl 3- cyclohexene carboxaldehyde	31906-04-4				
(Lyral)					
Limonene	5989-27-5				
Linalool	78-70-6				
Methyl 2-octynoate	111-12-6				

Section 5. Environmental Impact. *Instructions for Section 5: see page 6*

Please supply robust summaries of the tests performed as an attachment. The testing identified here was completed on the material as it is supplied. Select

If the answer is "no," please explain. Please refer to section 12 of SDS

Endpoint	Protocol	Date	Result
Aquatic Toxicity			
Short-term toxicity testing on Daphnia*			
Long-term toxicity testing on <i>Daphnia</i>			
Growth inhibition study on algae			
Short-term toxicity testing on fish*			
Long-term toxicity testing on fish (OECD 210, OECD 212 or OECD 215)			
Activated sludge respiration inhibition testing			
Nitrification inhibition testing			
Degradation			
Biodegradability Study			
Simulation testing on ultimate degradation in surface water			
Soil simulation testing			
Sediment simulation testing			
Aerobic rate of biodegradation (please specify media)			
Anaerobic rate of biodegradation (please specify media)			
Hydrolysis as a function of pH			
Identification of degradation products			
Fate & Behavior in the Environment			
Life cycle analysis (LCA)			
Adsorption/desorption screening study			
Bioconcentration in an aquatic species			
Effects on Organisms			
Short-term toxicity testing on earthworms*			
Long-term toxicity testing on earthworms			
Long-term toxicity testing on soil invertebrates other than earthworms			
Effects on soil microorganisms			
Short-term toxicity testing to plants*			
Long-term toxicity testing to plants			
Long-term toxicity testing to sediment organisms			
Long-term or reproductive toxicity testing to birds			

Section 6: Plant Derived: *Instructions for Section 6: see page 7.*

Please attach a manufacturing flow chart specifying mode of isolation or extraction. For materials that have identified plant or combination feedstock origin, or contain more than one plant species please fill out one for each species identified.

Please specify plant species Not Applicable							
Please specify plant part(s) used to make this material							
Extract % Solids %							
If a marker compound is included, or available please identify and specify level.							
Please specify Phylogenetic name:							
Genetically modified? Yes ☐ No⊠ Unknown ☐							
If the INCI name includes the phylogenic name of the plant, please answer the following questions:							
1. Is your Raw Material included in the IFRA guidelines? Yes No							
2. If 'Yes', does the Raw Material meet the IFRA guidelines? Yes No							
3. If 'Yes', please specify the amendment:							
4. Revision Date:							
5. Was the plant / botanical /fragrance screened for pesticides? Yes No							
6. If 'Yes', what are the pesticide residue levels?							
7. Do the pesticide residues present meet the European Pharmacoposia Guidelines? Yes No							
8. Does the Raw Material contain any denatured alcohol? Yes No							
9. If "Yes", please specify the denaturant(s):							
Does the hotenical contain, by addition or as a processing aid, any of the following nut/seed products?							

Materials of Concern	Does Not	All Sources Total	Added Ingredient Contribution	Natural Source Contribution
	Contain	Inclusion Level %	Inclusion Level %	Inclusion Level %
Tree Nuts				
Almonds				
Brazil nuts				
Cashews	\square			
Chestnuts	\square			
Cobnuts	\square			
Hazelnuts (Filberts)	\square			
Macadamia nuts	\square			
Pecans	\square			
Pistachio nuts				
Shea nuts	\square			
Seed products, oils or				
derivatives				
Peanuts				
Sesame seed				
Sunflower seed	\square			
Cotton seed	\boxtimes			
Canola (Rapeseed)	\boxtimes			
Other (please specify and describe)				

Section 7: TSE Certification: *Instructions for Section 7: see page 7*

The information captured here is intended to collect details on components that are derived from internal organs or connective tissues of animals.

	Ingredient	Identify Animal Species	Identify Source Organs	Age of Animal	Country of Origin	BSE Country Status GBR#
1	Not applicable "Product is made only from non-animal derived raw materials"	Select				
2		Select				
3		Select				
4		Select				
5		Select				
6		Select				
7		Select				
8		Select				
9		Select				
10		Select				

^{**} Category rating based on the EU Commission scientific steering committee (SCC) Regulation 999/2001-Geographical Risk of Spongiform Encephalopathy (GBR) Adopted on January 11, 2002. Updated on October 31 2007. As noted on the <u>EFSA website</u>.

NOTE: Level I--Highly Unlikely, Level II--Unlikely but Not Excluded, Level III--Likely but Not Confirmed or Confirmed at a Lower Level, Level IV--Confirmed at a Higher Level

Certification of Isolation Method:

The material described in the table above is certified to be classified as a Category 3 under <u>Regulation (EC) No</u> <u>1774/2002</u>. As such the material shall be compliant with Articles 6 & 7 including:

- (b) processed in a processing plant approved in accordance with Article 13 using any of processing methods 1 to 5, in which case the resulting material shall be permanently marked, where technically possible with smell, in accordance with Annex VI, Chapter I, and disposed of as waste either by incineration or by co-incineration in an incineration or co-incineration plant approved in accordance with Article 12 or in a landfill approved under Directive 1999/31/EC;
- (c) processed in a processing plant approved in accordance with Article 17;
- (d) transformed in a technical plant approved in accordance with Article 18;

and suitable for its intended use.

Global Regulatory Customer Request Team_AC Name of Certifying Person

Analyst Title

February 17, 2021 Date

Section 7: TSE Certification