26-3-2015 No	Solvay 1.1. Chemical Name		
No	1.1. Chemical Name		
	neodymium trihydroxide	1.2. EC Number 240-514-4	1.3. CAS Number 1.4. Composition Type 16469-17-3 Mono-constituent substance
This Sub	ostance Identification Profile (SIP) is developed	to represent the Identification parameters of the Substance de	escribed in line with the Substance Identification requirements of
Reference	SI Parameter REAC	H Annex VI and relevant Guidances for the purpose to identify Value / Not necessary / Not for SIP	the substance Remark / Justification
	Name or other Identifiers of the substance IUPAC Name	neodymium(3+) trihydroxide	
	Other International chemical name	not relevant	
	Chemical Name	neodymium trihydroxide	
	Abbreviation Other names	not relevant neodymium hydroxide	
1.1.2.0		neodymium(III) hydroxide	
	EC Number	240-514-4	
	EC Name EC Description	neodymium trihydroxide not available	
	CAS Number	16469-17-3	
	CAS Name	neodymium hydroxide	
	CAS Description IUBMB Number	not available not applicable	
	INCI Number	not applicable	
2.1.5.c	Other Catalogue identifiers	not applicable	
		ing under this substance (with justification)	I
	Chemical Name EC Number	not relevant not relevant	-
2.1.6.c	CAS Number	not relevant	
	Information related to molecular and struc		
	Molecular Formula Structural Formula	Nd(OH)3	
		QH	
		но — Na — он	
2.2.1.c	Smiles notation	[Nd+3].[OH-].[OH-].[OH-]	
	Optical activity	none	
	Typical ratio of (stereo) isomers Molecular Weight	not applicable 195.26 g/mol	
	Molecular Weight range	not applicable	
	Chemical Composition of the substance		
	Main Constituent Name -Main Constituent	neodymium trihydroxide	1
	CAS Number -Main Constituent	16469-17-3	
	EC Number -Main Constituent	240-514-4	
2.3.1.d	Concentration range -Main Constituent - Lower value	≥ 80%	
2.3.1.e	Concentration range -Main Constituent	100%	-
	- Upper value		
2.3.1.f	Typical concentration -Main Constituent (=	99,95%	
	Degree of purity) Impurity / Impurities (above 1% or lower if	contributing to the bayard or BBT profile)	
	Agreed strategy for Impurity profile on SIP	The impurity profile is not relevant for the SIP. It can	Each registrant will need to specify the impurities present in their
		however be relevant for Classification and Labelling.	company-specific (confidential) part of the joint registration dossier (section 1-3).
			The registration dossier, and in particular the suggested C&L and the hazard assessment, will assume that the substance as
			placed on the market conforms to:
			 All impurities > 1% do not significantly affect its toxicological and ecotoxicological properties. All hazardous impurities are present at < 0.1%.
			If a registrant's substance does not conform to the above
			specifications then the registrant will have to justify that the
			differences do not modify the IUCLID and CSR conclusions and do not require a different C&L or - if relevant - different exposure scenarios. This information will be reported in the company
			specific (confidential) part of the registration dossier.
	Additive(s) (above 1% or lower if contribut Agreed strategy for Additives profile on SIP	ing to the hazard) No additives above 1% or contributing to the hazard or	
		PBT profile.	
<mark>2,4</mark> 2.4.1	Suggestions for analytical and spectral methor Agreed Spectral data to be used	ods to be used for substance sameness check XRD	
f. I			
	Agreed Analytical Methods to be used	XRF or GDMS + Karl Fischer (for residual water)	
2,5 2.5.1	Substance Sameness Approval Agreed approval method for the sameness	Individual discussions with Consortium members result in a	
	checking procedure using this SIP	generic SIP. This generic SIP, after approval by the	
	(Consortium)	involved Consortium members, is sent to the entire SIEF for approval.	
2.5.2	Agreed approval method for the sameness checking procedure using this SIP (SIEF)	A generic SIP is sent to the entire SIEF. SIEF members that do not agree with the draft generic SIP must notify	
		ARCADIS before the deadline, including any relevant	
		information. SIEF members that agree with the draft	

He agrees that his substance does to the best of his knowledge completely fall under the substance identity being represented by the SIP sufficient for the purpose of meeting the SIEF requirements and opting for the joint submission Registration dossier to be created by the lead registrant in line with the REACH requirements. He agrees that he will inform the Consortium via the Secretariat or the SIEF via the Lead registrant if he has (new) information that might change the content of this SIP or if his Substance is changed in such a way that it might or does no longer fall under the SIP or might potentially have an impact on the content of the Registration dossier. He understands and agrees to be fully responsible for the proper linkage of the substance to the REACH Registration dossier and informing of his supply chain on the safe use of his substance and fulfilling his REACH requirements accordingly.