

The "low POP content" threshold of wastes has been set at 15 μ g/kg for dioxin (TEQ PCDD/F) as a provisional value⁴. All residue being bottom ash and flue gas treatment **residue (including fly ash) are expected to be well below the "low POP content"** threshold. Bottom ash is expected at a dioxin content (TEQ) around 0.001-0.030 μ g/kg, and flue gas treatment residue around 1 μ g/kg. This means that the Convention does not require further treatment of the residues prior to disposal when it comes to the dioxin content.

In conclusion all relevant paragraphs of the Stockholm Convention are considered and complied by the EfW facility.

⁴ Basel Convention, Technical Guidelines for the Environmentally Sound Management (ESM) of Wastes Consisting of, Containing or Contaminated with Persistent Organic Pollutants (POPs), cf.



4. Dioxin from EfW

An EfW facility may be illustrated as depicted in the figure below.



Figure 1 Typical Energy from Waste facility consisting of the main functions, 1) Waste bunker, 2) Furnace, 3)Energy recovery in a steam boiler, 4) Flue gas treatment, 5) stack, 6) electricity generation in turbine/generator set, 7) condensation of vapours in an air cooled condenser.

Waste is ignited and burnt on the grate in the furnace at temperatures around 1'100° C and the temperature of the flue gases is thereafter kept above 850 °C for at least 2 seconds in the afterburning chamber.

The flue gas treatment stage consists of a reactor with injection of lime and activated carbon for dioxin adsorption followed by a bag house filter for dust separation, including the activated carbon particles with dioxin adsorbed.

The flue gas treatment system ensures that the stack emissions comply with the emission requirement of 0.1 ng/m³ (at reference conditions) regardless the content in the raw, untreated flue gas within any realistic operational range.

4.1 Destruction and formation of dioxin in the furnace/boiler system

Input waste to the furnace contains dioxin. The content of dioxin is destroyed in the course of the combustion process where the gas temperature reaches above 1'000 °C and also in the afterburning chamber where the temperature is maintained at minimum 850 °C for minimum 2 seconds.



Dioxin is known to oxidise at high temperatures, for instance the Australian Department of Environment indicated thermal decomposition at 700 °C. ⁵ Also Vehlow⁶ illustrated how the temperature affects the dioxin level, Figure 2.



Figure 2 Formation of PCDD and PCDF in fly ashes from waste incineration during annealing in air atmosphere.

Also a report of US EPA⁷ stated that 'a number of studies have provided evidence that most of the CDDs/CDFs present in the MSW are destroyed during combustion'.

The dioxin content in the raw, untreated flue gas originates mostly from formation on the boiler walls in the temperature range 250-400 °C.

The mass balance of dioxin is demonstrated by Vehlow. It indicates that the typical picture is that the dioxin content of the incoming waste grossly exceeds the sum of the outputs, so the EfW-facility is a net destructor of dioxin. The air emission is less than 1% of the content of the incoming waste. As mentioned previously the fly ash and flue gas treatment residue are disposed in a safe manner, outside reach of the environment.



Figure 3 Dioxins (TEQ) in a state of the art EfW-facility. [Footnote 6]

 $^{^{5}\} http://www.npi.gov.au/resource/polychlorinated-dioxins-and-furans$

⁶ Jürgen Vehlow, Dioxins in Waste Combustion – Conclusions from 20 Years of Research, Bioenergy Australia, Melbourne, 2005.

⁷ Mechanisms of formation of dioxin-like compounds during combustion of organic materials, Draft of May 2005.

http://www.epa.gov/ncea/pdfs/dioxin/2k-update/pdfs/Dioxin_Chapter_2.pdf



5. Dioxin emission and inventory

The total dioxin emission in 2014 in Australia[®] is listed as around 300 g per year of TEQ dioxins of which more than half (170 g per year) originates from back yard incinerators.

The total dioxin emission from the TNG EfW facility is estimated as 0.7 g per year TEQ, emission if it operates at full capacity at the emission limit value of 0.1 ng/m³ (ref.) continuously. The typical emissions from EfW facilities are however around a factor 10 lower⁹, causing the expected dioxin emission to be around 0.07 g per year TEQ or around 0.02% of the Australian inventory, and 0.05% of the contribution from back yard incinerators.

6. On domestic burning of waste

Blacktown District Environmental Group argues on domestic incineration, that "Local residents have been prevented from incinerating rubbish in their own backyard - unfair to now impose industrial incinerator on those same residents."

The ban on domestic incineration appears to be in agreement with the provisions of the POP convention to reduce dioxin emissions, as cessation of open burning of waste is specifically mentioned in the convention Annex C (section A of part V), cf. section 3.1 above.

Transfer of waste from domestic incineration to the industrial EfW facility will save the environment for large amounts of dioxin, which is particularly important for the local community where domestic incineration happens.

⁸ Total national facility emissions for the substance: Polychlorinated dioxins and furans (TEQ) http://www.npi.gov.au/npidata/action/load/summary-

result; jsessionid=26410198AE95A174F96749000E3FE4F9/criteria/substance/73/destination/ALL/source-type/ALL/substance/name/Polychlorinated%2Bdioxins%2Band%2Bfurans%2B%2528TEQ%2529/subthreshold-data/Yes/year/2014

⁹ Emissions from decentralised chp plants 2007 - energinet.dk environmental project no. 07/1882, Appendix 3, listing an emission factor of 5 ng/GJ (TEQ) and considering a flue gas flow rate 550 m³/GJ (ref.).



Job	Compilation of COPC Memos
Client	EfW Facility TNG NSW
Date	2016-10-20
То	To Whom it may concern
From	Martin Brunner (Ramboll)
Copy to	Ian Malouf (DADI)
	Phill Andrew (Savills)
	Amanda Lee (AECOM)
	Lesley Randall (AECOM)
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Compilation and update of memos on compounds of potential concern (COPC)

Over the course of the project several memos concerning COPC for the HHRA have been established. Following a summary and overview of the memos with subject, date of issue and revision date is shown.

	Job	Date of Issue	Revision date
Memo 1	Compounds of Potential Concern (COPC) for HHRA	13.09.2015	-
Memo 2	COPC for HHRA	20.09.2015	19.10.2016
Memo 3	COPC for HHRA – Cr(VI)	03.11.2015	19.10.2016
Memo 4	VOC for HHRA	20.10.2016	-
Memo 5	Bromine in Waste	14.10.2016	-

These memos shall serve as an input to the air quality assessment (AQA) and the human health risk assessment (HHRA)

In summary the most important changes compared to the memos edited until end of 2015 are:

- Update of appendix B of Memo 2 (maximum TOC/VOC concentrations)
- Update Cr(VI) emissions
- Evaluation of further VOC compounds
- Assessing bromine emissions

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Attachments

Memos 1 - 5



MEMO

Job	Compounds of Potential Concern (COPC) for HHRA
Client	DADI TNG NSW
Memo no.	1
Date	13/09/2015
То	Lesley Randall (AECOM)
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1. Reference and basis

Reference is made to the following memos:

- a) "TNG Energy from Waste Facility Inputs to Human Health Risk Assessment", dated 11. September 2015 by Damon Roddis (Pacific Environment)
- b) "Advice to address EPA comments", dated 29. January 2015 by Rosalind Flavell (Fichtner)

In line with the above information we have evaluated the in stack concentrations for normal and upset operation based on real data of 4 plants (7 lines and 7 different measuring campaigns) with identical Air Pollution Control system (APC) as planned to be installed at the TNG facility. We have further considered general literature on emission factors of WtE plants. Where no such data was available the concentration was calculated on the expected particulate emission and appropriate concentration of the compound in fly ash. More detailed description of the data used will follow in a separate memo. All values are given based on the following assessment:

Normal operation: Maximum value out of the following:

- Any measured value from the plants with identical APC system
- Literature emission factor for WtE plants

Upset operation: Definition of "Upset Operating Conditions" see memo b) chapter 1. Maximum value out of the following:

- Particulate emission of 150 mg/Nm3, emission based on specific compound concentration in fly ash
- Gas flow of 10% of total gas flow to stack bypassing APC (e.g. bag failure)
- Value of 10 times normal operation

Date 13/09/2015

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When evaluating these data we found that in the memo b) some values were far above operational data. As a result we have re-evaluated the values of memo b) for normal operation. Further some in stack concentrations during upset operation (mainly HF and Dioxins) seem to be highly exaggerated, however (from footnote to table 1) we understand these values were requested by the EPA.

2. Table 1: Missing COPC

Compound	Operation condition			
All values at 11%O2, dry gas		normal	upset	
Beryllium	mg/Nm3	7.00E-06	5.25E-04	
Silver	mg/Nm3	3.40E-04	2.55E-02	
Cobalt	mg/Nm3	4.00E-03	4.00E-02	
PCB (WHO TEQ humans/mammal)	mg/Nm3	1.60E-08	1.60E-07	
PAH (WHO TEQ humans/mammal)	mg/Nm3	5.00E-04	5.00E-03	
Zinc	mg/Nm3	3.70E-02	5.09E+00	
Tin	mg/Nm3	3.33E-03	2.50E-01	
Molybdenum	mg/Nm3	2.20E-05	2.63E-03	
Selenium	mg/Nm3	2.12E-03	2.12E-02	
НСВ	mg/Nm3	8.21E-06	8.21E-05	

3. Table 2: Overestimated COPC

Compound	Operation condition			
All values at 11%O2, dry gas		normal	upset	
Mercury	mg/Nm3	0.004	0.013	
Cadmium	mg/Nm3	0.009	0.090	
Thallium	mg/Nm3	0.001	0.009	
Nickel	mg/Nm3	0.021	0.208	
PCDD/F TEQ (WHO humans/mammal)	ng/Nm3	0.010	0.500	



4. Relevant flue gas volume

For calculation of the ground level concentration the methodology described in memo a) should be used. In case of any doubt the following revised flue gas flow shall be applied.

Parameter		Value		
	Design Point (LPN)			
Number of streams	1	2	4	
Stack Height (m)		100		
Stack Diameter each stream inside (m)		2.2		
Temperature (°C)	120			
Flue Gas Flow (Nm3/s)	57.4	114.8	229.6	
Gas Exit Flow Rate (Am3/s)	82.6	165.2	330.5	
Gas Exit Velocity (m/s)	21.7			
Flue Gas Flow (Nm3/s) @ 11% O2	63.5	127.0	254.0	
Flue gas composition (v/v)		•		
H2O		15.90%		
02		6.60%		
N2		67.80%		
CO2		9.70%		

5. Conclusion

For the further HHRA the data listed in table 1 (above) shall be used.

In case that as a result of the HHRA for one of the compounds listed in table 2 "overestimated COPC" (above) shows to be critical (when using the concentrations in memo b)) we suggest to use the values given in table 2 (above).



MEMO

Job	COPC for HHRA
Client	EfW Facility TNG NSW
Memo no.	2 – Rev 1
Date	19/10/2016
То	To Whom it May Concern
From	Martin Brunner (Ramboll)
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1. Background and goal

In the context of the input to the Human Health Risk Assessment (HHRA) there has been some discussion on the list of Compounds of Potential Concern (COPC). The following memo gives an explanation on why the current COPC's have been chosen.

2. Basis of the current list of COPC's

An initial list of COPC's was established as a basis of the report "energy from waste facility - air quality and greenhouse gas assessment" produced by Pacific Environment in March 2015.

The COPC's chosen where based on the primary emissions from any Energy-from-Waste (EfW) facility, as defined by emission limits for waste incineration set by the European Union (EU) Industrial Emissions Directive (IED; Directive 2010/75/EU).

The emissions defined by the EID and chosen as COPC's are listed in Appendix A.1. In addition to the emissions identified in the IED, the substances listed in A.2 were included. As a result of the submissions from the Public Exhibition the substances listed in A.3 were added.

The current list of COPC's is substantially broader than substances usually taken into account in an Environmental Impact Assessment for an EfW plant in Europe. Nevertheless it is reasonable to question whether this list is complete. The following shall provide the rational for our opinion that the current list is sufficient to perform the HHRA. Date 19/10/2016

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3. The legislation principle of primary emissions and "lead substances"

While emissions in general cover a broad range of toxic, carcinogenic, mutagenic, etc. substances every industry has a specific set of primary emissions which - for reasons of human health and environmental protection - have to be reduced. It is therefore obvious that legislation focuses on the relevant emissions for any industry.

Besides the primary emissions so-called "lead substances" can be defined. Lead substances are representative for an entire group of comparable compounds and either relevant in their toxicity or present in high concentration. These substances are often difficult to capture by an Air Pollution Control (APC) system. Measuring low concentrations of these lead substances therefore is the proof that the separation mechanisms of the APC control are working. Typical lead substances of an EFW plant are: HCl, SO₂, NOx, TOC, CO, dioxins and furans, cadmium, mercury and further heavy metals as nickel, lead or arsenic.

The chosen approach to primarily focus on substances defined in the IED and further lead substances therefore is rational and good industry practice.

4. The implications of the "lead substance approach"

When considering the emissions of an Energy-from-Waste plant the following categories of compounds can be defined:

- particulate matter
- acid gases (HCI, HF, SO₂)
- NO_x
- heavy metals with low boiling point (mercury, cadmium) and volatile compounds, to a high degree present in vaporised form
- heavy metals with high boiling point (nickel, vanadium, etc.), predominantly present in particle form
- Organic substances (expressed as total organic carbon TOC)
- dioxins and furans

Every one of these substances (and therefore the appropriate category) has a specific reduction mechanism in combustion and the APC process. While the lime injection reduces the acid gases, the bag filter eliminates the particulates and any substance in particulate form (mainly heavy metals with high boiling point). Finally the activated carbon injection reduces organic substances and heavy metals with low boiling point by adsorption.

As mentioned earlier a low emission of any lead substance is the proof of an efficient reduction of the category they represent in general.

The COPC's recently added (appendix A.3) all can be classified in the above categories, e.g.:

- Copper, Molybdenum: metals with high boiling point
- PCBs and PAHs: organic substances
- Selenium, Beryllium: metals with low boiling point
- etc.

As a result any further substance can be classified in the above categories and therefore the reduction efficiency (respectively a low emission) can be assured.



5. TOC as guarantee for low organic emissions

While there is a limited number of toxic metals (including their compounds) there is an indefinite number of organic substances. The most important ones have been listed in the relevant regulations and are part of the current list of COPC's. However it is impossible to supply a complete list of individual organic substance and their emission data.

For this reason an additional emission parameter "total organic carbon" (TOC) has been introduced to legislation. The TOC measurement ensures that no relevant amount of organic substances is emitted. The TOC measurement is usually based on "Flame Ionisation Detection" (FID or FIA) and part of the continuous emission monitoring of any EfW plant. The TOC is a summary parameter for organic substances in general, the result is expressed in "carbon equivalent". Average TOC results of energy from waste plants are in the range of $1 - 2 \text{ mg/Nm}^3$ (Nm³ is normal cubic meters, i.e. at standard temperature (0 °C) and pressure (101.3 kPa)).

In Europe extensive research has been done on the composition of the TOC of Energy-from-Waste plants. In total less around 50% of the TOC can be allocated to substances with higher molecular weight (see attachment B). The other 50% (or more) are "light" substances like methane, propane, etc. This is further underlined by theoretical considerations [2] which predict that a part of the TOC will be methane, ethane and propane.

To illustrate the low expected emission level, it can be mentioned that the background ambient air concentration of methane is around 1,800 ppb (volume basis) equivalent to around 1 mg/Nm³ TOC.

6. Operational Data

The above considerations are further supported by operational data (see appendix C). The appendix C.1 shows publicly available emission data from plants exclusively fired by C&I and C&D waste with semi dry APC system (as used for the TNG project) as well as plants with mixed waste (MSW plus C&I, C&D). In summary all values are comparable and far below the emission limits. Further details on operational data are found in appendix C.2 and C.3.

7. Summary and conclusions

The current list of COPC has been established on the following considerations:

- Compounds regulated by recent legislation (in this case the IED; Directive 2010/75/EU) and therefore relevant for the EfW Industry
- Additional COPC's which are not of primary relevance for a EfW plant but might be of public concern
- Lead substances which demonstrate the ability of the APC system to reduce pollutant categories and therefore not only assures a low emission of the substance itself, but also of the entire category
- TOC as an overall guaranty for low organic emissions which as research has shown –contain very low concentrations of potentially harmful substances

As a result of the above and "real data" from comparable plants we are of the opinion that the current list of COPC is exhaustive and a sufficient basis to perform a robust and trustworthy HHRA.



Appendix A

1. The emissions defined by the EID and chosen as COPC's

- Particulate matter (PM), assumed to be emitted as PM10 and PM2.5.
- Hydrogen Chloride (HCl).
- Hydrogen Fluoride (HF).
- Carbon Monoxide (CO).
- Sulfur Dioxide (SO₂)
- Oxides of nitrogen (NOx) (expressed as Nitrogen Dioxide (NO₂)).
- Heavy metals (including Mercury (Hg), Cadmium (Cd), Arsenic (As) and Chromium (Cr).
- Gaseous and vaporous organic substances (expressed as total organic carbon (TOC)).
- Dioxins and furans.

2. In addition the following substances were included:

- Hydrogen sulfide (H₂S).
- Chlorine (Cl₂).
- Ammonia (NH₃).
- Polycyclic aromatic hydrocarbons (PAHs).

3. As a result of the submission the list was amended by the following substances:

- Beryllium (Be)
- Silver (Ag)
- Asbestos
- Copper (Cu)
- Cobalt (Co)
- Manganese (Mn)
- Vanadium (V)
- Polychlorinated Biphenyls (PCBs)
- PAHs (as benzo(a)pyrene equivalent).
- Zinc (Zn)
- Tin (Sn)
- Molybdenum (Mo)
- Selenium (Se)
- Hexachlorobenzene (HCB)



Appendix B

TOC composition in emissions from an EfW plant [1].

Measurement based on adsorption and condensation. Detection limit 5 μ g/Nm³.

Characterisation of Emissions from a Waste Incineration Plant

Total organic carbon (TOC)	1.2 mg/m ³
Identified single components	0.53 mg/m ³
Not identified aliphatic hydrocarbons	56% of TOC

Main Components (µg/m ³)								
Benzoic Acid	100							
Hexadecanoic Acid	37							
Ethyl Benzoic Acid	35							
Toluene	30							
Phthalates	20							
Dichloromethane	20							
Acetone (propanone)	18							
Tetradecanoic Acid	15							
Benzene	15							
Acetonitrile	14							
Xylene	10							
Trichlorophenol	9							
Methylhexane	6							
Trichloroethylene	5							
Heptane	5							

Note: There is little literature on the above subject. Most dates from mid 1990ies, when new emission regulations were issued in Europe. The concentrations of the organic substances were consistently low and therefore no further research or measurements were performed.

For any other TOC compound a maximum in stack concentration of 5 μ g/Nm³ can be assumed. In case of a compound listed as group (e.g. Phthalates) for a conservative approach a maximum concentration for each speciation according to the above value can be chosen.

Literature references

- [1] Ergebnisbericht über Forschung und Entwicklung 1994, Institut für Technische Chemie, Forschungszentrum Karlsruhe, Wissenschaftliche Berichte, FZKA 5531, S. 9
- [2] Stand der Gesamtkohlenstoff-Messung im Abgas von Abfallverbrennungsanlagen, Staub – Reinhaltung der Luft, 49 (1989), S. 221-225
- [3] Emissions from decentralized CHP plants 2007 ENERGINET.DK Environmental project No. 07/1882 – National Environmental Research Institute (NERI) Technical Report no. 786, 2010 (available from http://www2.dmu.dk/Pub/FR786.pdf).



Emission Data from plants with C+I / C+D and/or semi dry APC Publicly available Data

	Plant		h-Knapsack	EEW H	eringen	EEW Premnitz	EEW Gro	ssräschen		Riverside				Vallorca		IED
	Country	[DE	0	DE	DE	0	DE		UK			F	E		
	Waste	C&I,	, C&D	C&I, RDF fro	C&D, m MSW	C&I, C&D	C&I,	C&D	Munici	ipal Solid Was	ite, C&I	Hosp	Municipal soli ital waste, se	id waste, C&I wage sludge	, , tyres	IED limit value
	unit	Line 1	Line 2	Line 1	Line 2		Line 1	Line 2	Line 1	Line 2	Line 3	Line 1	Line 2	Line 3	Line 4	
Total Dust	mg/m³	0.01	0.2	0.4	0	0.2	0.2	0.2	1	1	1	0.3	0	0.4	0.4	10
Total Organic Carbon (TOC)	mg/m³	0.2	0.1	1	0	0.5	0.8	0.5	5	5	3	0.05	0.03	0	0	10
Inorganic chlorine compounds (HCI)	mg/m³	9	9	6	7	6	6.6	3.5	6	3	6	0.5	0.1	0	0.1	10
Inorganic fluorine compounds (HF)	mg/m³	0	0	-	-	0	-	-					ľ		ľ	1
Sulphur dioxide (SO ₂)	mg/m³	2	1	27	11	7.5	18	18	0	0	5	4	2	15	11	50
Oxides of nitrogen (expressed as NO ₂)	mg/m³	188	188	183	185	180	174	176	175	175	175	55	38	60	68	200
Mercury (Hg)	µg/m³	4	23	1	0	3	0.3	0.1								50
Carbon monoxide (CO)	mg/m³	23	23	6	7	12.5	8	8				2.5	5	2.5	2.5	50
Ammonia (NH₃)	mg/m³	2	2				0.4	0.9	0	1	2		ľ		ľ	
													ľ		ľ	
Dioxines and furanes	ng/m3	0.01	0.02	-	-	0.015	-	-	-	-	-	-	- 1	-	- 1	0.1

all values at standard conditions, 11% O2 dry

all values (except Hg and dioxines & furanes) as daily average, Hg and dioxines & furanes as spot sampling all plants except Mallorca with SNCR DeNox, Mallorca with SCR

Sources

EEW Hürth-Knapsackhttp://www.chemiepark-knapsack.de/fileadmin/user_upload/EEW_Emissionswerte_2013.pdfEEW Heringenhttp://www.eew-energyfromwaste.com/de/emissionswerte-heringen.htmlEEW Premnitzhttp://www.eew-energyfromwaste.com/de/standorte/hannover.html#c347bEEW Grossräschenhttp://www.eew-energyfromwaste.com/de/standorte/heringen.html#c399bRiversidehttp://www.coryenvironmental.co.uk/energy-from-waste/riverside-resource-recovery-facility/TIRME Mallorcahttp://www.tirme.com/uk/incineration_02f3s25.html

Hitachi Zosen INOVA

Extended values from plants with semi-dry APC

Detailed emission measurements from HZI plants with semi-dry APC

			Riverside		Newhaven		Cleveland	Evreux	Ingolstadt	Average	EU	
Metal	Symbol	Unit	Line 1	Line 2	Line 3	Line 1	Line 2	Line 3		see note		IED
Mercury	Hg	mg/m ³	0.0015	0.0004	0.0002	0.004	0.003	0.0017			0.002	< 0.05
Cadmium	Cd	mg/m ³	0.00270	0.00085	0.00111	0.009	0.001	0.004	0.004		0.00324	
Thallium	ТІ	mg/m ³	0.00005	0.00003	0.00002	0.000	0.000	0.0009			0.00017	
Sum Cd+Tl	Cd + TI	mg/m ³	0.00275	0.00087	0.00113	0.009	0.001	0.0049			0.003	< 0.05
Arsenic	As	mg/m ³	0.0006	0.0003	0.0004	0.003	0.000	0.0013	0.004		0.0009	
Antimony	Sb	mg/m ³	0.0148	0.0047	0.0047	0.007	0.001	0.0026			0.0058	
Chromium	Cr	mg/m ³	0.0179	0.0115	0.0399	0.014	0.002	0.0467	0.004		0.0220	
Cobalt	Co	mg/m ³	0.0003	0.0002	0.0001	0.003	0.000	0.0006	0.004		0.0007	
Copper	Cu	mg/m ³	0.0085	0.0085	0.0263	0.051	0.001	0.0049			0.0167	
Lead	Pb	mg/m ³	0.0452	0.0137	0.0170	0.172	0.002	0.0094			0.0432	
Manganese	Mn	mg/m ³	0.0084	0.0041	0.0037	0.095	0.005	0.0051			0.0202	
Nickel	Ni	mg/m ³	0.0118	0.0058	0.0041	0.006	0.002	0.0208			0.0084	
Vanadium	v	mg/m ³	0.0003	0.0002	0.0004	0.003	0.000	0.0004			0.0007	
Sum heavy metal	As-V	mg/m ³	0.11	0.049	0.097	0.35	0.015	0.092			0.12	< 0.5
Dioxins and Furanes	PCDD/F TEQ (WHO humans/mammal)	ng/m³	0.004	0.004	0.001	0.0015	0.0004				0.0022	
Dioxin-like PCB's	PCB (WHO TEQ humans/mammal)	ng/m ³	0.016	0.011	0.014	0.00001	0.00001				0.008	
Hexachlorbenzol	НСВ	µg/m3								< 0.0012	0.001	
Benzo(a)pyren	B(a)P	µg/m3							0.002	< 0.0013	0.002	
PAH's	PAH (WHO TEQ humans/mammal)	µg/m³				0.4	0.5				0.45	

all concentrations in gas ref. to STP and 11% O2 dry

note: Ingolstad has APC with wet scrubber and bag house filter

Extract of the revised 2006 (2007 for natural gas fuelled plants) emission factors for Danish decentralised CHP plants < 25MWe. [3]

Note: For calculation to/from GJ to/from mg/m3 the report uses the flue gas amount of 523 Nm3 (dry, at 11% O2) per GJ for MSW.

	Unit	Natural gas fuelled engines	Biogas fuelled engines	Natural gas fuelled gas turbines	Gas oil fuelled engines	Gas oil fuelled gas turbines	Fuel oil, steam turbines	Biomass producer gas, engines	MSW incinera- tion	Straw	Wood
SO ₂	g per GJ	-	-	-	-	-	-	-	< 8.3	49	< 1.9
NO _x	g per GJ	135 ⁸⁾	202	48	942	83	136	173	102	125	81
UHC (C)	g per GJ	435 ⁸⁾	333	2.5 ⁹⁾	(46) ¹⁰⁾	-	(1.6) ¹⁰⁾	12	< 0.68	< 0.94 ⁵⁾	< 6.1 ⁶⁾
NMVOC	g per GJ	92 ^{4) 8)}	10 ⁴⁾	1.6 ⁴⁾	(37) ¹⁰⁾	-	(0.8) ¹⁰⁾	2.3 ⁴⁾	< 0.56 ⁴⁾	< 0.78 ⁴⁾	< 5.1 ⁴⁾
CH ₄	g per GJ	481 ^{4) 8)}	434 ⁴⁾	1.7 ⁴⁾	24	-	< 1.3	13 ⁴⁾	< 0.34 ⁴⁾	< 0.47 ⁴⁾	< 3.1 ⁴⁾
СО	g per GJ	58 ⁸⁾	310	4.8	130	2.6	2.8	586	< 3.9	67	90
N ₂ O	g per GJ	0.58	1.6	1.0	2.1	-	5.0	2.7	1.2	1.1	0.83
NH_3	g per GJ	-	-	-	-	-	-	-	< 0.29	-	-
TSP	g per GJ	-	-	-	-	-	9.5	-	< 0.29	< 2.3	10
As	mg per GJ	< 0.045	< 0.042	-	< 0.055	-	-	0.116	< 0.59	-	-
Cd	mg per GJ	< 0.003	0.002	-	< 0.011	-	-	< 0.009	< 0.44	< 0.32 ³⁾	0.27
Со	mg per GJ	< 0.20	< 0.21	-	< 0.28	-	-	< 0.22	< 0.56	-	-
Cr	mg per GJ	0.048	0.18	-	0.20	-	-	0.029	< 1.6	-	-
Cu	mg per GJ	0.015	0.31	-	0.30	-	-	< 0.045	< 1.3	-	-
Hg	mg per GJ	< 0.098 ³⁾	< 0.12	-	< 0.11	-	-	0.54	< 1.8	< 0.31 ³⁾	< 0.40 ³⁾
Mn	mg per GJ	< 0.046	0.19	-	0.009	-	-	0.008	< 2.1	-	-
Ni	mg per GJ	0.045	0.23	-	0.013	-	-	0.014	< 2.1	-	-
Pb	mg per GJ	0.043	0.005	-	0.15	-	-	0.022	< 5.5	-	-
Sb	mg per GJ	< 0.049 ³⁾	0.12	-	< 0.055	-	-	< 0.045	< 1.1	-	-
Se	mg per GJ	(0.01) ⁷⁾	< 0.21	-	< 0.22	-	-	< 0.18	< 1.1	-	-
TI	mg per GJ	< 0.20 ³⁾	< 0.21	-	< 0.22	-	-	< 0.18	< 0.45 ³⁾	-	-
V	mg per GJ	< 0.048	< 0.042	-	0.007	-	-	< 0.045	< 0.33	-	-
Zn	mg per GJ	2.9	4.0	-	58	-	-	0.058	2.3	0.41	2.3
PCDD/-F	ng per GJ	< 0.57	< 0.96 ¹⁾	-	< 0.99	-	-	< 1.7 ¹⁾	< 5.0	< 19	< 14
PBDD/-F	ng per GJ	-	< 5.0 ¹⁾	-	-	-	-	< 7.2 ¹⁾	< 6.3 ¹⁾	-	-
PAH (BaP)	µg per GJ	< 13	< 4.2	-	< 33	-	-	< 4.9	< 2	< 125	< 13
ΣΡΑΗ	µg per GJ	< 1025	< 606	-	< 8988	-	-	< 181	< 37	< 5946	< 664
Naphthalene	µg per GJ	2452	4577	-	17642	-	-	8492	< 129 ³⁾	12088	2314
HCB	µg per GJ	-	0.19	-	< 0.22	-	-	0.80	< 4.3	< 0.11	-
PCB	ng per GJ	-	< 0.19 ¹⁾	-	< 0.13 ¹⁾	-	-	< 0.24 ¹⁾	< 0.32	-	-
Formalde- hyde	g per GJ	14.1	8.7	-	1.3	-	< 0.002	1.5	-	-	-
HCI	g per GJ	-	-	-	-	-	-	-	< 1.14	56	-
HF	g per GJ	-	-	-	-	-	-	-	< 0.14	-	-

¹⁾ Emission measurements were below detection limits for all congeners.

²⁾ Based on 1 emission measurement. The emission measurement was below the detection limit.

³⁾ All emission measurements were below the detection limit.

⁴⁾ Based on disaggregation of the total unburned hydrocarbon (UHC) emission factor.

⁵⁾ Only 1 out of 7 emission measurement was above the detection limit.

⁶⁾ Two out of three emission measurements were below the detection limit.

⁷⁾ Two emission measurements were performed, both below the detection limit. These results have been ignored and instead the lower emission factor 0.01 mg per GJ based on EEA (2009) have been applied.

⁸⁾ The increased emission level during start up and stop of the gas engines have been included in this emission factor.

⁹⁾ Based on emission measurements performed in 2003-2006.

¹⁰⁾ The emission factor based on emission measurements performed within this project has been ignored. Instead the NMVOC emission factor refers to EEA (2009). The UHC emission factor has been estimated based on the emission factors for NMVOC and CH₄.



MEMO

COPC for HHRA - Cr(VI) Job Client EfW Facility TNG NSW Memo no. 3 – Rev 1 19/10/2016 Date То To Whom it May Concern From Martin Brunner (Ramboll) Copy to Ian Malouf (DADI) Lesley Randall (AECOM) Amanda Lee (AECOM) Damon Roddis (Pacific Environment) Phill Andrew (Savills) Rachael Snape (Urbis) Damon Roddis (Pacific Environment)

1. Background and goal

Chromium (Cr) is widely used metal and appears in different valences, mostly as trivalent Cr(III) but also as hexavalent Chromium Cr(VI). Cr(VI) is toxic and carcinogenic and therefore of major concern.

In case of emissions from Energy from Waste (EfW) facilities Chromium is considered as part of the sum of heavy metals and measured as total Chromium. In the context of the Human Health Risk Assessment (HHRA) for the TNG facility Ramboll has been requested to give a forecast on the emission of Cr(VI). Date 19/10/2016

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2. Basis of the Cr(VI) forecast

As mentioned earlier, Chromium emissions are limited as total Chromium and therefore very little data on Cr(VI) emissions are available. Existing data date back to the 1980 and these emissions are not comparable to today's Air Pollution Control (APC) systems.

The forecast for updated Cr(VI) emissions therefore is based on total Cr emission, the APC removal behaviour and recent data of Cr(VI) values in APC residues.

3. Existing data on Cr(VI) in APC residues

There is a variety of data on total Chromium in EfW fly ash. The values for total Chromium typically range from 500 to 1000 mg/kg of fly ash.

Few measurements exist on Cr(VI) in fly ash. While many are below the detection limit (< 0.05 mg/kg) some values in the range of 1 - 3 mg/kg are found.

Chromium in the flue gas is predominantly present in form of particulates; the vapour pressure is very low and not relevant for the emission level. As a result it can be expected that the in stack concentrations will have a similar distribution as the fly ash.



4. Data considered for total Cr emissions

The following plants with a flue gas cleaning system identical to TNG were considered

Plant	No of Measurements	Max value mg/Nm ³	Mean value mg/Nm ³
Riverside UK	3	0.040	0.023
Newhaven UK	2	0.014	0.008
Cleveland UK	1	0.047	0.047
Mallorca ES	2	0.002	0.002
Phitiviers FR	1	0.002	0.002
Perpignan FR	1	0.024	0.024
Le Mans FR	3	0.009	0.005
Evreux FR	2	0.007	0.006
Ludwigslust DE	6	0.007	0.004
Zorbau DE	6	0.014	0.011
Total/max/mean	27	0.047	0.010

All values refer to 11% O2, dry

5. Evaluation of Cr(IV) in APC residues

The Cr(VI) content in APC residues is in the order of 1-3 mg/kg in relation to a total Chromium content of 500-1000 mg/kg. Expressed as fraction this is 0.1 to 0.3% of the total. To allow for uncertainties due to variations a content of 0.5% as average and 1% as worst case is assumed

6. Conclusion

Based on results of 27 emission measurements of existing plants with identical APC equipment a maximum of total Chromium of 0.047 mg/Nm³ and a mean of 0.010 mg/Nm³ (see table above) is reported. This is well in line with a report of the UK EPA (see attachment) which lists a maximum of 0.052 mg/Nm³ and a mean of 0.011 mg/Nm³ as a result of measurements in 10 plants in the UK.

As a worst case scenario during normal operation therefore a Cr(VI) emission of 0.0005 mg/Nm³ (1% of 0.052 mg/Nm³) and an average of 0.00005 mg/Nm³ (0.5% of 0.010 mg/Nm³) can be assumed.

The above results are well in line with a recent publication by the Environment Agency of the UK (see attached) which predicts maximum Cr(VI) levels of 0.00013 mg/Nm³ and a mean value of 0.000035 mg/Nm³.



Releases from municipal waste incinerators

September 2012 version 3

Guidance to applicants on impact assessment for group 3 metals stack

Scope

This paper provides guidance to Applicants on how we will consider air quality impact assessments from Group 3 metals stack emissions from Municipal Waste Incinerators when we determine permit applications in respect of Schedule 1 activities under the Environmental Permitting Regulations 2010 (EPR). Metals assessments from other plant subject to the Waste Incineration Directive may use the method in this guidance if they can justify the data as representative.

Background

In April 2010, the Environment Agency published revised Environmental Assessment Levels (EALs) for arsenic, nickel and chromium(VI) in our H1 Guidance (<u>H1 Environmental Risks Assessment</u>). The revised EALs are substantially lower than the former EALs:

- Arsenic 3 ng/m³
- Nickel 20 ng/m³
- Chromium (VI) 0.2 ng/m³

The EALs refer to that portion of the metal emissions contained only within the PM10 fraction of particulates in ambient air.

Arsenic, nickel and (total) chromium are three of the nine Group 3 metals whose emissions are subject to a mandatory minimum emission limit by the Waste Incineration Directive (WID). WID sets an aggregate limit of 0.5 mg/m³ for nine "Group 3" metals (Sb, As, Pb, Cr, Co, Cu, Mn, Ni and V and their compounds (total)). Previous air dispersion modelling studies supporting permit applications typically made very conservative assumptions that emissions of each individual metal occurs at the WID aggregate limits. Such an analysis may conclude that there is a risk that the current EALs might be exceeded. Where such a theoretical risk exists, a more detailed assessment is required to determine whether the impact of the release is acceptable.

Detailed Modelling Assessment Methodology

Step 1 - Screening scenario

Predictions made assuming each metal is emitted at 100% of the WID ELV (i.e. 0.5 mg/m³). Where the impact of any metal exceeds the assessment criteria (below), relative to their respective EALs, we consider that there is a potential for significant pollution. Under these circumstances, proceed to Step 2.

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Assessment Criteria:

- Long-term Process Contribution (PC) <1% or Short-term Process Contribution (PC) <10%; or
- Long-term and Short-term Predicted Environmental Concentration (PEC) <100% [taking likely modelling uncertainties into account].

[For screening only, assume Cr(VI) comprises 20% of the total background chromium). Selection of all other background data should be justified.]

Step 2

Worst case scenario based on currently operating plant – make predictions based on assuming each metal comprises 11% of the total group (i.e. 0.5 mg/m³ apportioned across the nine metals). Our emissions monitoring data indicates that it is reasonable to assume that each Group 3 metal comprises no more than 11% of the Group ELV.

Where the impact of any metal is above the assessment criteria given in Step 1 above proceed to Step 3.

Step 3 - Case specific scenario

We will require Applicants to justify their use of percentages lower than 11% of the Group 3 WID ELV or Cr(VI) background levels of <20% for their Step 3 assessment. Assessments should be made using the criteria below Step 1. We will review any use of Applicants' data to identify whether they can achieve the levels proposed and whether that data can be justified as representative.

Appendix A of this guidance contains summary of measured metals stack releases from a range of operating Municipal Waste Incinerators between 2007 and 2009, presented as a range and a mean of actual release and percentage of the WID ELV. The data in Appendix A should be considered as indicative only. Note that although the maximum Nickel concentration is greater than 11%, this represents one single measurement outlier; the mean value is around 4% of the Group ELV.

Appendix B contains data showing the effective Cr(VI) concentration from a range of Municipal Waste Incinerators. Measurement of Cr(VI) at the levels anticipated at the stack emission points is expected to be difficult, with the likely levels being below the level of detection by the most advanced methods. The concentrations presented in the table are based on stack measurements for total chromium and measurements of the proportion of Cr(VI) to total chromium in APC residues collected at the same plant. We have considered the concentration of total chromium and Cr(VI) in the Air Pollution Control (APC) residues collected upstream of the emission point for existing Municipal Waste Incinerators (MWI) and have assumed these to be similar to the particulate matter released from the emission point.

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LIT 7349 2 of 3

	Measured Concentrations mg/m ³			Percentage	ercentage of WID Group 3		
	Mean	Мах	Min	Mean	Мах	Min	
Antimony	0.0033	0.0115	0.0001	0.7%	2.3%	0.02%	
Arsenic	0.0007	0.0030	0.0003	0.14%	0.6%	0.06%	
Chromium	0.0109	0.0521	0.0004	2.2%	10.4%	0.08%	
Cobalt	0.0004	0.0039	0.0002	0.07%	0.8%	0.04%	
Copper	0.0077	0.0163	0.0025	1.5%	3.3%	0.50%	
Lead	0.0158	0.0368	0.0003	3.2%	7.4%	0.06%	
Manganese	0.0172	0.0365	0.0015	3.4%	7.3%	0.30%	
Nickel	0.0220	0.1362	0.0000	4.4%	27.2%	0.00%	
Tin		0.0024	0.0024		0.48%	0.48%	
Vanadium	0.0003	0.0010	0.0002	0.06%	0.20%	0.04%	

Appendix A – Monitoring Data from Municipal Waste Incinerators

Values correspond to the distribution from 19 measurements at 13 plant between 2007 and 2009. The data differs slightly from previous guidance notes.

* Minimum values correspond in some cases to the detection limit

Appendix B – Chromium VI analysis from APC Residues

	Effective Cr(VI) emission concentration ^a (mg/Nm ³)
Mean	3.5*10- ⁵
Minimum	2.3*10- ⁶
Maximum	1.3*10- ⁴

These data are taken from ten MWI plant in England and Wales. We are in the process of gathering more data in order to fully understand the implications of metals emissions.

^a Note the maximum total chromium concentration does not coincide with the plant where the maximum chromium VI fraction in the APC residue was observed.



MEMO

Job Client Memo no. Date To From Copy to COPC for HHRA - VOC EfW Facility TNG NSW 4 2016-10-20 To Whom it may concern Ahmet Erol (Ramboll) Ian Malouf (DADI) Phill Andrew (Savills) Amanda Lee (AECOM) Lesley Randall (AECOM) Rachael Snape (Urbis) Damon Roddis (Pacific Environment)

VOC for HHRA/ Air quality Assessment

Background

In Ramboll Memo 2 COPC for HHRA, dated 20.09.2015 the COPC for AQA and HHRA have been listed. The selected COPC where based on the primary emissions from any Energy-from-Waste (EfW) facility, as defined by emission limits for waste incineration set by the European Union (EU) Industrial Emissions Directive (IED; Directive 2010/75/EU). It has been questioned if this list is covering all necessary compounds.

Further input to the selection of COPC

The study "Site specific risk assessment of an energy-from-waste thermal treatment facility in Durham Region, Ontario, Canada. Part A: Human health risk assessment"¹ (following referred to as "the study") has considered a wide range of COPC. This study is one of the most comprehensive investigations on the relevance of emissions from EfW facilities currently available.

The study categorizes the COPC's in five groups:

- 1. Metals
- 2. Chlorinated Polycyclic Aromatics
- 3. Chlorinated Monocyclic Aromatics
- 4. Poly Aromatic Hydrocarbons (PAH)
- 5. Volatile Organic Chemicals (VOC)

Methodology

COPC listed in the study and not selected so far for the TNG project were extracted. Then they were evaluated if the study found that they contribute to more than 1% of the background concentration.

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¹ Site specific risk assessment of an energy-from-waste thermal treatment facility in Durham Region, Ontario, Canada. Part A: Human health risk assessment, dated 4th of July 2013 (attached)



Evaluation

Below the COPC are listed according to the categories of the study and evaluated if they are already part of the AQA/HHRA. If they have not been selected so far then to what extend they contribute to an increase of the ground level concentration compared to the baseline.

1. Metals

The metals listed in the study are already included in the AQA. Therefore no further assessment is needed.

2. Chlorinated Polycyclic Aromatics

Chlorinated Polycyclic Aromatics listed in the study are already included in the AQA. Therefore no further assessment is needed.

3. Chlorinated Monocyclic Aromatics (CMA)

The table below shows all COPC categorized under CMA in the study and the evaluation if they have already been included in the AQA.

	Listed in the study	TNG Air quality Assessment Memo 2, Appendix A
1,2-Dichlorobenzene	х	not included
1,2,4,5-Tetrachlorobenzene	х	not included
1,2,4 – Trichlorobenzene	х	not included
Pentachlorophenol	х	not included
Hexachlorobenzene	х	х
Pentachlorobenzene	х	not included
2,3,4,6-Tetrachlorophenol	х	not included
2,4,6-Trichlorophenol	X	not included
2,4-Dichlorophenol	Х	not included

Only hexachlorobenzene has been considered for the AQA so far. The assessment concerning impact of the EfW facility on the ground level concentration found in the study is shown below.

The concentration ration values are listed below	(see study	y table 5))
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Chlorinated Monocyclic Aromatics	Baseline	Project Alone	Effect level
1-hour	0.0006	7.5E-05	12,5%
24-hour	0.0001	5.2E-07	0,5%
Annual	0.002	3.0E-06	0,15%

Only the 1-hour value is higher than 1%. All other values are below 1%.

Conclusion

Hexachlorobenzene is the most relevant CMA compound present in the emissions from EfW plants. The concentrations measured during normal operation are in the range of below 1 up to max. 10 ng/Nm³.

The other CMA are mostly expressed as the sum of compounds with identical number of chlorine atoms (dichlorobenze, trichlorobenzene, dichlorphenol, etc.). The measured values for such a group are in the range of below 1 up to max. 10 ng/Nm³. For any assessment a concentration of max. 10 ng/Nm³ per group during normal operation and 100 ng/Nm³ (tenfold value) during upset operation can be assumed.



4. Poly Aromatic Hydrocarbons

Poly Aromatic Hydrocarbons (PAH) listed in the study have been included in the AQA. No further assessment is needed.

5. Volatile Organic Chemicals (VOC)

The table below shows all COPC categorized under Volatile Organic Chemicals (VOC) in the study and the evaluation if they have already been included in the AQA.

	Listed in the study	TNG Air quality Assessment Memo 2, Appendix A
Acetaldehyde	х	not included
Benzene	х	x (Appendix B)
Biphenyl	Х	not included
Bromodichloromethane	Х	not included
Bromomethane	Х	not included
Dichlorodifluoromethane	Х	not included
Dichloroethene	Х	not included
1,1 -, Ethylbenzene	Х	not included
Ethylene Dibromide (1,2-dibromoethane)	Х	not included
Formaldehyde	Х	not included
Tetrachloroethylene	х	not included
Toluene	Х	x (Appendix B)
Trichloroethylene	Х	x (Appendix B)
1,1,2, Vinyl chloride (chloroethene)	Х	not included
Xylenes	Х	x (Appendix B)
m-, p- and o-Bromoform (tribromomethane)	Х	not included
Carbon tetrachloride	Х	not included
Chloroform	Х	not included
Dichloromethane	Х	x (Appendix B)
O-terphenyl	X	not included
Trichloroethane	X	not included
1,1,1 -, Trichlorofluoromethane	x	not included

The concentration ration values are listed below (see assessment EfW facility table 5)

Volatile Organic Chemicals (VOC)	Baseline	Project Alone	Effect level
1-hour	0.55	0.005	0,9%
24-hour	0.41	0.002	0,5%
Annual	0.18	0.0002	0,1%

All effect levels are below 1%. No further assessment for VOC is needed.



Summary and conclusion

Except for chlorinated monocyclic aromatics all compounds evaluated in the study "Site specific risk assessment of an energy-from-waste thermal treatment facility in Durham Region, Ontario, Canada. Part A: Human health risk assessment" are either already included in the AQA or their effect level found in the study is below 1% under all conditions.

For the chlorinated monocyclic aromatics hexachlorobenzene (HCB) - the most relevant compound - is already included in the AQA. Measured values of HCB in EfW plants range from 1-10 ng/Nm³. For any other group of compound (as sum of compounds with identical number of chlorine atoms) a concentration of 10 ng/Nm³ during normal operation and 100 ng/Nm³ (tenfold value) during upset operation can be assumed. Contents lists available at ScienceDirect

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Site specific risk assessment of an energy-from-waste thermal treatment facility in Durham Region, Ontario, Canada. Part A: Human health risk assessment



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HIGHLIGHTS

• Human health risk assessment was performed for an Energy-From-Waste facility

· Results suggest minimal risks to humans expected at approved operating capacity

Future expansion may cause slightly elevated risks under upset conditions

• Further risk assessment required if/when future expansion is pursued

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ABSTRACT

The regions of Durham and York in Ontario, Canada have partnered to construct an energy-from-waste thermal treatment facility as part of a long term strategy for the management of their municipal solid waste. This paper presents the results of a comprehensive human health risk assessment for this facility. This assessment was based on extensive sampling of baseline environmental conditions (e.g., collection and analysis of air, soil, water, and biota samples) as well as detailed site specific modeling to predict facility-related emissions of 87 identified contaminants of potential concern. Emissions were estimated for both the approved initial operating design capacity of the facility (140,000 tonnes per year) and for the maximum design capacity (400,000 tonnes per year). For the 140,000 tonnes per year scenario, this assessment indicated that facility-related emissions are unlikely to cause adverse health risks to local residents, farmers, or other receptors (e.g., recreational users). For the 400,000 tonnes per year scenarios, slightly elevated risks were noted with respect to inhalation (hydrogen chloride) and infant consumption of breast milk (dioxins and furans), but only during predicted 'upset conditions' (i.e. facility start-up, shutdown, and loss of air pollution control) that represent unusual and/or transient occurrences. However, current provincial regulations require that additional environmental screening would be mandatory prior to expansion of the facility beyond the initial approved capacity (140,000 tonnes per year). Therefore, the potential risks due to upset conditions for the 400,000 tonnes per year scenario should be more closely investigated if future expansion is pursued.

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1. Introduction

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The Regions of Durham and York in Ontario, Canada partnered in 2005 to seek a long-term sustainable solution for managing their municipal solid waste. Both Regions have made considerable commitments to decreasing waste production and increasing waste diversion (e.g. through recycling or composting initiatives), but a management strategy is still required for residual waste not diverted through these strategies. Previously, this residual waste was largely exported out of the Regions (primarily to Michigan) for landfill. However, when it was announced that the Michigan border would be closed to municipal waste from Canada as of December 2010, it became imperative to identify a viable waste management alternative.

Abbreviations: CAC, Criteria air contaminant; COPC, Contaminant of potential concern; CR, Concentration ratio; CSF, Cancer slope factor; EA, Environmental assessment; EFW, Energy-from-waste; ERA, Environmental risk assessment; HHRA, Human health risk assessment; HQ, Hazard quotient; ILCR, Incremental lifetime cancer risk; LADD, Lifetime average daily dose; LCR, Lifetime cancer risk; LRASA, Local risk assessment study area; MDL, Method detection limit; RfC, Reference concentration; RfD, Reference dose; TEF, Toxic equivalency factor; TRV, Toxicity reference value; UR, Unit risk.

Due to public opposition, establishment of a new local landfill was considered unacceptable. In addition, it was recognized that continuing to ship the waste to an external landfill could not provide a stable and secure alternative due to the vulnerability of this option to public policy decisions made by external governments. Therefore, processing and treatment options such as mechanical, biological, and thermal treatment were considered. Through an extensive public consultation process as well as a detailed evaluation of environmental, social and economic considerations, the preferred option was determined to be the construction of an Energy-From-Waste (EFW) thermal treatment plant. Such facilities have the capacity to reduce the volume of waste by >90% while also recovering metals and producing energy that can be sold to offset annual operating costs (Rushton, 2003).

EFW facilities are widespread in Europe and other jurisdictions (Bogner et al., 2008). Research and monitoring programs around these facilities suggest that in light of strict emissions guidelines and modern engineering controls, these facilities are unlikely to be hazardous to human health or the environment (Bordonaba et al., 2011; Cangialosi et al., 2008; Lee et al., 2007; Morselli et al., 2011; Rovira et al., 2010; Schuhmacher and Domingo, 2006). However, a new EFW facility had not been built in Ontario for over 20 years. As part of the approval process for construction of this new facility in Ontario, extensive human health and ecological risk assessments were performed to determine the potential effects of this project on surrounding communities and ecosystems. This paper describes the methods and results of human health risk assessment; the methods and results of the ecological risk assessment are provided in a separate publication (Ollson et al., 2014). These risk assessments formed an important component of the final Environmental Assessment for this project, which was submitted to the Ontario Ministry of the Environment (MOE) in 2009 and received final approval in 2010. On the basis of this approval, the project was permitted to proceed to the construction phase, which was initiated in 2011. Facility start-up is currently projected to occur by the end of 2014.

2. Material and methods

2.1. Scope of the assessment

This risk assessment examined the potential for emissions from the proposed project (i.e., construction, operation, and eventual decommissioning of a modern EFW thermal treatment facility) to pose an unacceptable risk to human health over both short-term and long-term (i.e., after 30 years of operation). Existing conditions at the proposed location for the facility were also assessed in order to provide a baseline for the assessment (Table 1). The entire assessment was carried out following the US EPA human health risk assessment protocol for hazardous waste combustion facilities (US EPA, 2005).

The initial operating design capacity of the proposed facility was 140,000 tonnes per year, with a capacity for expansion to 400,000 tonnes per year within the 30-year planning period. As the expansion

of the facility beyond the initial approved capacity of 140,000 tonnes per year would require additional environmental screening under provincial regulations, the present risk assessment focused primarily on the potential risks from the facility with respect to operation at the 140,000 tonnes per year level. However, for comparison purposes, consideration was also given to the potential risks associated with the maximum design capacity of 400,000 tonnes per year.

2.2. Facility description

Facility design information for this assessment was provided by Covanta Energy Corporation, which was selected by the Regions as the preferred vendor for this project. Covanta, the largest provider of thermal treatment services in North America (with 40 facilities in the United States and one in Canada), was contracted by the Regions to direct the design, engineering, construction and operation of the facility. Therefore, they were able to provide detailed information, specific to the planned facility, which also reflects the features and functionality of existing modern EFW facilities elsewhere in North America.

This facility will be accepting municipal solid waste from typical Ontario curbside waste collection (i.e. household waste excluding separated recyclable materials and organics). No additional feed stock separation will occur at the facility. The facility will use a thermal mass burn technology, wherein municipal solid waste is fed into a furnace and burned at very high temperatures. For the initial operating design capacity of 140,000 tonnes per year, there will be two independent waste processing trains consisting of a feed chute, stoker, integrated furnace/boiler, dry recirculation acid gas scrubber, a fabric filter bag house and associated ash and residue collection systems. Expansion to the maximum design capacity (400,000 tonnes per year) would include the addition of two more waste processing trains. Steam produced in each boiler will drive a turbine-generator to produce electricity for delivery to the grid, for in-plant use and/or district heating. After the removal of residual metals for recycling, ash produced by the process will be shipped to landfill for use as daily cover or will be reused, possibly as road construction material or other civil projects. Air pollution control equipment throughout the facility will ensure that emissions do not exceed the provincial guidelines outlined by the Ontario Ministry of the Environment (MOE, 2004a) and specific conditions of Certificate of Approval 7306-8FDKNX issued June 28, 2011 for the Facility.

2.3. Identification of chemicals of potential concern (COPC)

Chemicals that could potentially be released by the facility to the atmosphere were identified by reviewing sources such as existing provincial guidelines for municipal incinerators (MOE, 2004a), the Canadian National Pollutant Release Inventory for waste incinerators (Environment Canada, 2007), and the results of stack testing of an existing waste incinerator in nearby Brampton, Ontario. From this review, a COPC list consisting of 87 chemicals was developed (Table 2) that consisted of both Criteria Air Contaminants (CACs,

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Project scenarios considered in the human health risk assessment.

Project Scenarios	Case	Conditions assessed
Existing Conditions	Baseline	Existing conditions in the assessment area. No Facility-related emissions or exposures
		were included as this was completed prior to construction and operation of the Facility.
	Baseline Traffic	Offsite vehicle traffic emissions prior to the start-up of the Facility.
Construction	Construction	Construction and commissioning of the Facility.
Operation	Project Alone	Emissions from the Facility alone.
	Project (Baseline + Project)	Emissions from the Facility combined with existing/baseline conditions.
	Process Upset	Emissions from the Facility operating at upset conditions (i.e., Facility start-up, shutdown,
		and loss of air pollution control).
	Process Upset Project (Baseline + Upset)	Emissions from the Facility operating at upset conditions combined with existing/baseline conditions.
	Traffic	Emissions from offsite and onsite traffic associated with the Facility combined with baseline
		traffic conditions and onsite stationary source emissions for the Facility.
Decommissioning	Decommissioning (Closure Period)	Emissions related to the removal of infrastructure and rehabilitation of the Site.

for which regulatory limits already exist) and non-CACs (substances that are capable of causing environmental or health effects for which no regulatory limits were identified).

All COPC were evaluated for their potential to pose a risk to human health via inhalation as this was expected to be the primary route of human exposure to facility-related air emissions (Table 2). In addition, COPC that were considered to be persistent and/or bioaccumulative (i.e., half-life in soil \geq 6 months and/or Log K_{ow} \geq 5) were also included in a multi-pathway risk assessment that addressed the possibility that these compounds may persist in and/or be transferred to various environmental media (e.g., soil, water, and food) following their release to air (Table 2).

2.4. Study area

The selected location for the facility is located within the Municipality of Clarington, Ontario, Canada (approximately 80 km east of Toronto, Ontario). This location is bordered by Lake Ontario to the south, commercial properties to the north and agricultural lands to the east and west. The Darlington Nuclear Generating Station is located approximated 2 km to the east.

In order to define the study area, the CALPUFF dispersion model (Scire et al., 1995) was applied to predict ground level concentrations of COPC as well as wet and dry deposition fluxes over a 40×40 km grid around the proposed facility location. The inputs to this model included geophysical (terrain and land use) and meteorological data specific to the region (Environment Canada, 2008; USGS, 2007; UCAR, 2008) as well as COPC physical-chemical properties. Stack parameters (i.e., location, base elevation, stack height, stack diameter, gas exit velocity, gas exit temperature, and emission rates) were provided by the vendor with respect to the planned facility. Potential stack emissions of COPC were estimated based on manufacturer's guarantees of maximum emissions, emission levels measured by the preferred vendor at one or more of their existing facilities that utilise similar technologies (measured at maximum load), and literature sources for other facilities.

Results of the CALPUFF model showed that the highest concentrations of emissions and depositions would be located in the area immediately surrounding the facility with a radius of approximately 10 km. Therefore, this area was defined as the Local Risk Assessment Study Area (LRASA) for consideration in this risk assessment. This LRASA includes the urban centers of Oshawa, Courtice, Bowmanville, and Port Darlington, Ontario.

2.5. Receptor identification and exposure pathways

Residential land use in the LRASA is mainly suburban residential and rural residential. The rural residential areas include large, dispersed lots that may be used for agricultural purposes (e.g., cash crops or livestock). Within the larger urban centers there are numerous commercial and institutional developments. Recreational opportunities in the area include hiking, camping, equestrian activities, hunting, fishing and swimming.

In light of these identified land uses, the human receptors considered in this risk assessment included local residents, local farmers, daycare/school attendees, and recreational users (sport and/or camping) (Table 3). Potential exposure pathways determined for each receptor included inhalation of vapours and particulate emissions, ingestion and dermal exposure to soil and/or dust, and food chain exposures (Table 3). It was also assumed that some receptors may incur additional exposures to COPC via hunting, fishing, or swimming within the LRASA. Therefore, additional exposures related to these activities that can be added to any of the identified receptors were also assessed (Table 3). Consumption of local drinking water was not considered since it was found that residents in the LRASA obtain their drinking water from municipal water supply services, which would not be affected by facility-related emissions. Similarly, consumption of grocery store bought foods was not considered.

The life stages considered for each receptor and for the hunting/ angling and swimming additional exposures were selected to represent those with the greatest sensitivity and/or exposure to each COPC. For non-carcinogenic COPC, which act via a threshold mechanism, the

Table 2

Contaminants of potential concern (COPC) considered in this assessment.

COPC	Inhalation	Multi-Pathway
Criteria Air Contaminants:		
Sulfur Dioxide (SO ₂), Hydrogen Chloride (HCl), Hydrogen Fluoride (HF), Nitrogen Dioxide (NO ₂),		
Particulate Matter (PM ₁₀), Particulate Matter (PM _{2.5}), Total Particulate Matter (TSP),		
Ammonia (Slip at Stack)		
Chlorinated Polycyclic Aromatics:		
Dioxins and Furans as Toxic Equivalents (TEQ), Total PCBs (as Aroclor 1254)		
Metals:		
Antimony, Arsenic [®] , Barium, Beryllium [®] , Boron, Cadmium [®] , Chromium (hexavalent) [®] ,	Land Contract of C	
Total Chromium (and compounds) ^o , Cobalt, Lead, Mercury ^a , Nickel, Phosphorus, Silver,		
Selenium, Thallium, Tin, Vanadium, Zinc		
Chlorinated Monocyclic Aromatics:		
1,2-Dichlorobenzene, 1,2,4,5-Tetrachlorobenzene, 1,2,4 – Trichlorobenzene, Pentachlorophenol ¹⁰ ,	Land Contract of C	
Hexachlorobenzene", Pentachlorobenzene		
2,3,4,6-Tetrachlorophenol, 2,4,6-Trichlorophenol ⁹ , 2,4-Dichlorophenol	Land Contract of C	
Poly Aromatic Hydrocarbons:		
Acenaphthylene ⁰ , Acenaphthene ⁰ , Anthracene, Benzo(a)anthracene ⁰ , Benzo(b)fluoranthene ⁰ ,		
Benzo(k)fluoranthene ^b , Benzo(a)fluorene, Benzo(b)fluorene, Benzo(ghi)perylene ^b ,		
Benzo(a)pyrene 1EQ ⁰ , Benzo(e)pyrene ⁰ , Chrysene ⁰ , Dibenzo(a,C)anthracene ⁰ ,		
Dibenzo(a,h)anthracene ", Fluoranthene ", Fluorene, Indeno(1,2,3 – cd)pyrene",		
Perylene ", Phenanthrene ", Pyrene "		
1 – methylnaphthalene, 2 – methylnaphthalene, Naphthalene		
Volatile Organic Chemicals (VOC):		
Acetaldehyde ⁹ , Benzene ⁹ , Biphenyl, Bromodichioromethane, Bromomethane, Dichlorodifluoromethane,		
Dichloroethene, I, I -, Ethylbenzene, Ethylene Dibromide (1,2-dibromoethane) ¹⁰ ,		
Formaldenyde ⁵ , letrachioroethylene ⁵ , loluene, lrichloroethylene, 1,1,2 ⁻⁵ ,		
Vinyl chloride (chloroethene) ³ , Xylenes, m-, p- and o-		
Bromotorm (tripromometnane), Carbon tetrachioride b, Chlorotorm b, Dichloromethane b,		
O-terpnenyi, Iricnioroetnane, 1,1,1 -, Iricniorofiuoromethane		

^a Inorganic and methylmercury.

^b This chemical was evaluated as a non-carcinogen and a carcinogen.

Table 3

Exposure pathways and life stages evaluated for identified receptor types.

	Receptor Type			Additional Exposures ^a			
	Resident	Farmer	Recreation User – Sport	Recreation User - Camping	Daycare	Swimming	Hunting/Angling
Exposure Pathway							
Direct Inhalation							
Soil Ingestion							
Dermal Contact – Soil							
Dermal Contact – Water							
Incidental Surface Water Ingestion							
Garden Produce							
Fish							1
Breast Milk							
Wild Game							1
Agriculture							
Life stage considered for threshold (not	1-carcinogenic) COPC					
Infant (0 to 6 mo)		/					
Toddler (7 mo to 4 vr)	-	-		100		1	
Life stage considered for non-threshold	(carcinogenic) COPC					
Adult (20 to 75 yr)							
Composite							

^a Exposures through these pathways can be added to identified receptors.

toddler life stage (i.e., 6 months to 4 years) was considered to represent the most sensitive life stage based on receptor characteristics (e.g., lower body weights) combined with behavioural patterns (e.g., higher soil ingestion rates). Therefore, all health risks associated with exposures to non-carcinogenic COPC were estimated for the toddler receptor (Table 3). In addition, the infant life stage (i.e., 0 to 6 months) was evaluated for farmer and resident receptors in the multi-pathway risk assessment for non-carcinogenic COPC in order to address the potential health risks associated with consumption of breast milk (Table 3). For carcinogenic COPC (non-threshold), a composite life stage for most receptors was considered that combines the characteristics of infant (i.e., 0 to 6 months), toddler (i.e., 7 months to 4 years), child (i.e., 5 years to 11 years), adolescent (i.e., 12 to 19 years), and adult (i.e., 20 years to 75 years) life stages (Health Canada, 2007) (Table 3). However, for the daycare/school receptor, exposure to carcinogenic COPC was assessed only for the adult stage (Table 3) since this class of receptor has the potential to have the longest duration of exposure to the daycare/school conditions (assuming employment from youth to retirement at that location).

2.6. Collection of baseline data

In order to characterize pre-project baseline conditions, ambient air monitoring and soil, water, and biota sampling was performed in the vicinity of the proposed facility location. All laboratory analyses of the collected samples were conducted by ALS Laboratory Group using standard methods (See Supporting Information Section S1).

2.6.1. Baseline ambient air monitoring

An air monitoring station was set up approximately 2 km southwest of the proposed facility location. Data was collected and analyzed over a 15 month period (September 2007 to December 2008). The station continuously monitored Sulfur Dioxide (SO₂), Nitrogen Oxides (NOx), Carbon Monoxide (CO), Ozone (O₃), and Particulate Matter smaller than 2.5 microns (PM_{2.5}). Hi-volume air samplers were also installed to collect 24-hour average samples of Total Suspended Particulate (TSP) and metals, Polycyclic Aromatic Hydrocarbons (PAHs), and Dioxins and Furans (PCDD/F).

In addition, baseline offsite vehicle emissions prior to the start up of the facility were estimated using traffic volume estimates provided by URS Canada Inc. These traffic estimates were combined with the existing baseline ambient air conditions in the airshed to produce the baseline traffic case.

2.6.2. Baseline soil and biota sampling

Additional baseline soil and biota samples were collected and analyzed for the COPC identified for consideration in the multi-pathway risk assessment. The sampling program included collection of soil, terrestrial vegetation (forage, browse, and crops), small mammals, surface water, sediment and fish sampled within a 1 km radius of the proposed facility location. Where possible, samples were collected in areas where air modeling predicted maximum rates of deposition for various COPC, and locations were also selected to be representative of different land uses. In addition, agricultural products (beef, chicken, pork, dairy and eggs) and produce were collected from farms and markets located outside a 1 km radius due to limited availability. However, efforts were made to ensure that farms were located as close as possible to the proposed facility location, and therefore the collected samples are considered sufficient to represent baseline conditions for this assessment.

2.7. Fate and transport modeling of COPC from project-related emissions

The potential impacts of facility-related emissions on the concentrations of COPC in the surrounding environment were predicted using best available data (i.e., results of the CALPUFF modeling described in Section 2.4, physical-chemical properties of the COPC, and detailed geophysical and meteorological data specific to the LRASA) and accepted modeling techniques as described in the US EPA human health risk assessment protocol for hazardous waste combustion facilities (US EPA, 2005). Specifically, the contributions of facility-related emissions to ambient air concentrations were predicted for all COPC at 309 distinct receptor locations selected to represent a variety of land uses as well as areas where initial modeling suggested the highest acute (1-hr or 24-hr) or chronic (annual) ground level concentrations were likely to occur. Additionally, for the persistent and/or bioaccumulative COPC considered in the multi-pathway risk assessment (Table 2), facility-related changes in COPC concentrations in soil, surface water, garden and farm produce and fruit, agricultural products (i.e., beef, chicken, pork, dairy and eggs), wild game, fish, and breast milk were predicted at 133 of the 309 locations.

In addition to predictions made for emissions from the normal operating scenarios at both 140,000 and 400,000 tonnes per year, the potential emissions under 'process upset' conditions (i.e., facility start-up, shutdown, and loss of air pollution control) were modeled following protocol suggested by the US EPA (2005). Specifically, for determining short-term (1-hour to 24-hour average) ground level COPC concentrations under upset conditions, the emission rates for

the facility under normal operation were conservatively increased by a factor of ten. This factor was applied to all COPC except for SO₂ and NO_x for which emissions were increased by factors of 16 and 1.63 respectively, based on data received from the vendor. As per US EPA (2005) guidance, for metals and CACs it was assumed that the facility would operate under upset conditions for 5% of the year. Therefore, emission rates for these COPC were increased by a factor of 1.45 [(0.95 x 1) + (0.05 x 10) = 1.45], with the exception of SO₂ and NO_x, for which emission rates were increased by factors of 1.75 and 1.03, respectively using the same assumptions. For the remaining COPC (organics), annual average concentrations for the process upset case were increased by a factor of 2.8 based on an assumption that the facility would operate under upset conditions for 20% of the year $[(0.80 \times 1) + (0.20 \times 10) = 2.8]$ (also as suggested by US EPA, 2005). This upset case is considered an absolute extreme scenario, given that the Ministry of the Environment would not allow the facility to operate in upset conditions for 20% of the year.

2.8. Exposure assessment

The sources of chemical concentrations used in the exposure assessment are described in Sections 2.5 and 2.6. In order to ensure a conservative estimate of risk, all exposure assessments were conducted deterministically using exposure point concentrations representative of reasonable maximum exposure. For the baseline values (described in Section 2.6), a single baseline exposure point concentration (i.e., the maximum detected concentration, 95% upper confidence limit of the mean, or method detection limit as described in Supporting Information, Section S2) was used to model exposure for each environmental medium collected for all receptor types. Although individual baseline concentrations were not obtained at the location of each receptor group evaluated, the baseline exposure point concentrations used are considered representative of reasonable maximum exposure, to all receptors, from background concentrations. A different approach was applied for the modeled facilityrelated contributions of COPC to the environment. In this case, the receptor locations were grouped by similar land use and the maximum or 95% upper confidence limit of the mean (selected as described in Supporting Information, Section S2) of the air and/or deposition concentration of each COPC within each receptor grouping was used to calculate the level of exposure for the entire grouping.

Physiological and behavioural characteristics of the receptors (e.g., respiration rate, soils/dusts intake, time spent at various activities and in different areas) were selected, if available, from existing guidance documents (Health Canada, 1994, 2007; MOE, 2005; Richardson, 1997; US EPA, 1997, 2005). In addition, oral and dermal bioavailability factors were compiled from Health Canada (2007) or the US Department of Energy's Oak Ridge National Laboratory Risk Assessment Information System (RAIS) database (ORNL, 2008). Whenever possible, preference was given to Canadian guidance documents and literature (e.g. Health Canada, 2007; Richardson, 1997). More details regarding the specific assumptions, input parameters and calculations used for each exposure pathway and receptor are provided in the Supporting Information (Section S3).

Exposure estimation was facilitated through the use of an integrated multi-pathway environmental risk assessment model developed by the Study Team. The model is spreadsheet based (Microsoft Excel™) and incorporates the techniques and procedures for exposure modeling developed by the MOE and Health Canada, and the US EPA (Health Canada, 1994; 2007; MOE, 2005; Richardson, 1997; US EPA, 1997, 2005).

2.9. Hazard assessment

2.9.1. Identification of toxicity reference values (TRVs)

For chemicals that follow a threshold dose-response (i.e., noncarcinogens), a threshold level must be exceeded in order for toxicity to occur, and it is possible to derive a reference concentration (RfC, for inhalation receptors) or reference dose (RfD, for multi-pathway receptors) that is expected to be safe to sensitive subjects following exposure for a prescribed period of time (US EPA, 1989). For chemicals that follow non-threshold dose-responses (i.e., carcinogens), a specific dose where toxic effects manifest themselves cannot be identified as any level of long-term exposure to carcinogenic chemicals is associated with some hypothetical cancer risk. As a result, risk assessment of these types of chemicals typically considers evaluation of the incremental lifetime cancer risk (ILCR) associated with exposure to the chemical (US EPA, 1989). This may be estimated based on the unit risk (UR) or cancer slope factor (CSF) of the chemical, where UR represents the upper-bound excess lifetime cancer risk estimated to result from continuous exposure to an agent at a concentration of 1 μ g/L in water, or 1 μ g/m³ in air and CSF provides an upper bound estimate of the increased cancer risk from lifetime exposure to an agent (US EPA, 1989).

Literature and public guidance documents were reviewed to identify RfCs, RfDs, URs or CSFs for inclusion as toxicity reference values (TRVs) for each COPC. Regulatory benchmarks, which are also health-based but often also policy derived, were also considered as TRVs for some COPC. A summary of the non-carcinogenic and carcinogenic TRVs used in both the inhalation and multi-pathway exposure assessment are presented in Supporting Information (Section S4).

2.9.2. Chemical mixtures and additivity of risks

In order to properly assess health risks to the human receptors, certain groups of chemicals were assessed as mixtures. Specifically, dioxin and furan congeners and carcinogenic PAHs were assessed using the toxic equivalency factor (TEF) approach (Supporting Information, Section S5). TEFs for dioxin and furan congeners represent their potency relative to 2,3,7,8 TCDD (Van den Berg et al., 2006), while TEFs for carcinogenic PAHs represent their toxicity relative to benzo(a)pyrene (IPCS, 1998).

Additional groups of chemicals were identified that may have additive, synergistic, or antagonistic effects due to their similar toxic modes of action (see Table S7 in Supporting Information, Section S5). However, there is currently very little available toxicological data or regulatory guidance to support the prediction of the effects of simultaneous exposure to these chemicals. In the original risk assessment an approach assuming additivity of the effects was used (see details in Supporting Information, Section S5). However, as this approach is not based on actual toxicological study results and cannot consider more complex interactions (i.e. synergism or antagonism), it is considered highly speculative and was presented for information purposes only. In light of these uncertainties, the effects of simultaneous exposure to multiple pollutants are not discussed further in the present manuscript. It is acknowledged that the interpretation of the potential effects of simultaneous exposure to chemical mixtures remains a considerable source of uncertainty in human health risk assessments conducted in Ontario.

2.10. Risk characterization

2.10.1. Threshold chemicals (non-carcinogens)

The risk associated with threshold chemicals was assessed using a Concentration Ratio (CR) for the inhalation pathway. CR values were calculated by dividing the predicted ground level air concentration (1-hour, 24-hour or annual average) by the appropriate toxicity reference value (reference concentration [RfC] or health based inhalation benchmark), according to Eq. (1):

$$CR_{duration} = \frac{[Air]_{duration}}{RfC_{duration} \text{ or health benchmark}}$$
(1)

Where CR _{duration} represents a duration specific Concentration Ratio (unitless), calculated for 1-hr, 24-hr and chronic durations as appropriate; [Air] $_{duration}$ represents the predicted ground-level air concentration (µg/m³) for that duration and RfC $_{duration}$ represents the selected (duration specific) reference concentration (µg/m³). A CR less than or equal to one signifies that the estimated exposure is less than or equal to the exposure limit; therefore, no adverse health risk is expected. Conversely, a CR greater than one signifies the potential for adverse health effects.

For the multi-pathway risk assessment, a Hazard Quotient (HQ) approach was applied. HQ values were calculated by dividing the predicted exposure dose (via multiple pathways) by the appropriate toxicity reference value (reference dose [RfD]), according to Eq. (2):

$$HQ = \frac{\sum Exp}{RfD}$$
(2)

Where $\sum Exp$ represents the chronic exposure estimate resulting from the sum of multiple exposure pathways (µg/kg/day) and *RfD* represents the selected chronic reference dose (µg/kg/day). For the purposes of this assessment, it was considered that the intake of the COPC by all routes of exposure was unlikely to exceed the tolerable intake level when the HQ was less than 0.2. This conservative approach allows 80% of the tolerable daily intake of a COPC to be received from other sources not considered in this risk assessment.

2.10.2. Non-threshold chemicals (carcinogens)

Incremental lifetime cancer risk (ILCR) and lifetime cancer risk (LCR) estimates resulting from direct air inhalation were calculated described in Eqs. (3) and (4):

$$ILCR = [Air]_{project \ alone} \times UR \tag{3}$$

$$LCR = [Air]_{all \ sources} \times UR \tag{4}$$

Where $[Air]_{project\ alone}$ represents the predicted annual average ground-level air concentration from the Project Alone (μ g/m³), $[Air]_{all\ sources}$ represents predicted annual average ground-level air concentrations from all sources, and *UR* represents COPC-specific unit risk (μ g/m³)⁻¹.

For the multi-pathway risk assessment, ILCR/LCR estimates resulting from a lifetime of exposure through multiple pathways were calculated using Eqs. (5) and (6):

$$ILCR = \sum LADD_{project alone} x CSF$$
 (5)

$$LCR = \sum LADD_{all \ sources} \ x \ CSF$$
(6)

Where $\sum LADD_{project\ alone}$ represents the sum of average daily dose via multiple pathways from the project alone (µg/kg/day), $\sum LADD_{all\ sources}$ represents the sum of average daily dose via multiple pathways from the all sources (µg/kg/day), and CSF represents the cancer slope factor (µg/kg/day)⁻¹.

In this risk assessment, an ILCR of 1-in-1,000,000 was considered acceptable, as outlined in relevant provincial guidelines (MOE, 2005). As no regulatory guidance exists for LCRs, this value was compared with the typical observed cancer incidence in the Canadian population, which is 38% for women and 44% for men (Canadian Cancer Society, 2007).

3. Results and discussion

3.1. Risk characterization: Existing conditions

Human health risks resulting from baseline exposures to individual COPC in the baseline scenario (prior to construction of the facility) were estimated using the results of the baseline ambient air monitoring and the baseline soil and biota sampling (Supporting Information, Section S6).

3.1.1. Inhalation risk assessment: Non-carcinogens

For criteria air contaminants (CACs, for which regulatory limits already exist), no baseline case acute (1-hr or 24-hr) or chronic (annual) CR risk estimates exceeded the regulatory benchmark (CR = 1), therefore no adverse health risks were expected from exposure to baseline air concentrations of these compounds (Table 4). Additionally, baseline case CACs (including NO₂, SO₂, PM_{2.5}, and PM₁₀) were also compared to WHO benchmarks for informational purposes and no exceedances were observed (Table 4). Similar results were noted for the baseline traffic case, in which estimated offsite vehicle emissions were added to the measured baseline ambient air conditions, except for a slight exceedance (CR = 1.1) for annual nitrogen dioxide compared to the WHO benchmark (Supporting information Section S8). However, the concentration of nitrogen dioxide measured in the baseline ambient air monitoring program in the LRASA was similar to that observed in other urbanized areas such as Toronto, Hamilton, and Windsor (Supporting information, Section S7), therefore this observation does not represent a unique property-specific risk. For non-criteria air contaminants (for which no relevant criteria were identified) baseline case concentrations were also shown not to exceed the acute (1-hr or 24-hr) or chronic (annual) CR regulatory benchmark (Table 5).

3.1.2. Inhalation risk assessment: Carcinogens

For non-criteria air contaminants assessed as possible carcinogens, the estimated lifetime cancer risk (LCR) values associated with their baseline ambient air concentrations were calculated (Supporting information Section S8). Because there are no acceptable benchmarks for comparison of LCR values, the implications of baseline results for each receptor group and scenario are not discussed in detail. However, to put these values in context, the maximum LCR associated with an individual baseline ambient air concentration for a COPC addressed in this study was 3.1×10^{-3} % (Supporting information Section S8), while the typical observed cancer incidence in the Canadian population is 38% for women and 44% for men (Canadian Cancer Society, 2007).

3.1.3. Multi-pathway risk assessment: Non-carcinogens

For all non-carcinogens, baseline chronic risk estimates (via multiple exposure pathways) were expressed as HQ values (Tables 6, 7, and Supporting Information Section S8). For most receptors and COPC, the predicted hazard quotients did not exceed the regulatory benchmark of 0.2 for the Baseline Case. However, some exceedances were noted for resident and farmer infants and toddlers. Also, addition of the swimming or hunting/angling exposures to the toddler receptor also led to some exceedances. Therefore, these cases were examined further.

3.1.3.1. Resident infant. For the resident infant receptor, the multipathway assessment indicated that potential risks may exist from exposure to baseline concentrations of PCBs and dioxins/furans (Table 6, HQ values of 11 and 3.8, respectively). The identified risk from these compounds was entirely related to the ingestion of breast milk, for which the COPC concentrations had been predicted based on exposure of the infant's mother to measured or estimated background COPC concentrations in relevant exposure media (i.e., soil) and food items (e.g., produce, poultry, etc.). However, in the results of the baseline sampling program, concentrations of PCBs, dioxins and furans were frequently below detection limit for these exposure media and food items (Supporting Information, Section S6). In these cases, the method detection limit (MDL) was substituted for the contaminant concentration in order to provide a 'worst-case scenario' estimate of exposure. However, it is possible that actual contaminant concentrations were significantly lower than the MDL (or not present at all). Therefore, the HQ values for PCBs and dioxins/furans that were calculated in this assessment for the resident infant receptor may represent a significant overestimation of the actual risk.

Table 4

Concentration Ratio (CR) Values for Baseline and 140,000 tpy for Criteria Air Contaminants at the Maximum Ground Level Concentration. A bolded cell indicates exposure for that particular scenario and COPC exceeded the selected benchmark.

COPC	Concentration Ratio (CR) Values					Concentration Ratio (CR) Values –WHO Benchmarks ^f						
	Baseline	Project Alone	Project	Process Upset	Process Upset Project	Baseline	Project Alone	Project	Process Upset	Process Upset Project		
1-Hour												
Ammonia ^a	-	0.0006	0.0006	0.006	0.006	-	-	-	-	-		
Carbon Monoxide (CO)	0.07	0.001	0.07	0.01	0.08	-	-	-	-	-		
Hydrogen Chloride (HCl) ^a	-	0.04	0.04	0.44	0.44	-	-	-	-	-		
Hydrogen Fluoride (HF) ^a	-	0.01	0.01	0.13	0.13	-	-	-	-	-		
Nitrogen Dioxide (NO ₂)	0.16	0.11	0.27	0.18	0.34	0.32	0.22	0.54	0.36	0.68		
Particulate Matter - PM ₁₀ ^{a, b, e}	-	-	-	-	-	-	-	-	-	-		
Particulate Matter - PM _{2.5} ^{b, e}	-	-	-	-	-	-	-	-	-	-		
Particulate Matter - Total ^{b, e}	-	-	-	-	-	-	-	-	-	-		
Sulfur Dioxide (SO ₂)	0.03	0.02	0.05	0.29	0.32	-	-	-	-	-		
24 Hour												
24-noui		0.002	0.002	0.02	0.02							
Carbon Monovide (CO) ^c	-	0.005	0.005	0.05	0.05	-		-	_			
Hydrogen Chloride (HCl) ^a	-	0.02	-	- 0.23	0.23	-		-	_			
Hydrogen Eluoride (HE) ^{a, c}	_	-	-	-	-		_		_	_		
Nitrogen Dioxide (NO ₂)	0.29	0.03	0.32	0.05	0.34		_		_	_		
Particulate Matter - PM ₁₀ ^{a, e}	-	0.01	0.02	0.05	0.11		0.01	0.01	0.11	0.11		
Particulate Matter - PM_{25}^{e}	0.68	0.02	0.70	0.18	0.86	0.82	0.02	0.84	0.21	10		
Particulate Matter - Total ^e	0.29	0.004	0.30	0.04	0.34	-	-	-	-	-		
Sulfur Dioxide (SO_2)	0.07	0.006	0.08	0.10	0.17	0.15	0.01	0.17	0.22	0.38		
(2)												
Annual												
Ammonia ^a	-	7.8E-05	7.8E-05	0.0001	0.0001	-	-	-	-	-		
Carbon Monoxide (CO) ^d	-	-	-	-	-	-	-	-	-	-		
Hydrogen Chloride (HCl) ⁴	-	0.0007	0.0007	0.0010	0.0010	-	-	-	-	-		
Hydrogen Fluoride (HF) ^{ad}	-	-	-	-	-	-	-	-	-	-		
Nitrogen Dioxide (NO ₂)	0.62	0.003	0.62	0.003	0.62	0.93	0.005	0.93	0.005	0.93		
Particulate Matter - PM ₁₀ ^{a, u, e}	-	-	-	-	-	-	0.0008	0.0008	0.001	0.001		
Particulate Matter - PM _{2.5} ^{d, e}	-	-	-	-	-	0.98	0.002	0.98	0.002	0.98		
Particulate Matter - Total ^e	0.35	0.0003	0.35	0.0004	0.35	-	-	-	-	-		
Sultur Dioxide (SO ₂)	0.20	0.002	0.21	0.003	0.21	-	-	-	-	-		

^a Baseline Data Not Available.

^b 1-hr TRV Not Available.

^c 24-hr TRV Not Available.

^d Annual TRV Not Available.

^e Particulate Matter results include contribution of Secondary Particulate.

^f "-" indicates WHO benchmark not available.

3.1.3.2. Resident toddler. The multi-pathway assessment for exposure of the toddler resident receptor to COPC indicates that potential risks may exist from exposure to baseline concentrations of PCBs (HQ = 0.49), arsenic (HQ = 0.32) and thallium (HQ = 0.25) (Table 6). For PCBs, it was determined that the majority of risk was associated with ingestion of homegrown produce and fruit. However, as was previously noted in the discussion of the risk of PCBs to resident infants, the PCB concentrations in these media in the baseline sampling program were below detection limits and were replaced with the value of the MDL in the risk assessment. Therefore, the HQ value for PCB exposure for the toddler resident likely overestimates the actual risk.

For arsenic, risk to the toddler resident receptor was attributed to incidental ingestion of soil. In contrast to PCBs, arsenic was widely detected in soil in the baseline sampling program. However the maximum detected soil arsenic concentration (8 mg/kg) used in the risk characterization was within the range of concentrations previously reported in natural, uncontaminated soils in Canada (Wang and Mulligan, 2006) and was less than the current Ontario Ministry of the Environment regulatory soil chemical standard of 11 mg/kg for arsenic at sensitive sites (MOE, 2004b). Therefore, this soil is not likely to cause any undue risk to human receptors within the LRASA. The elevated HQ values observed for the resident toddler receptors for arsenic can likely be attributed to conservative model assumptions applied throughout the risk assessment process.

For thallium, the relevant exposure pathways that contributed to the potential risk to resident toddlers were incidental soil ingestion and produce and fruit ingestion. However, none of the soil, produce, or fruit samples collected during the baseline sampling program had detectable levels of thallium. Therefore, the risk assessment for thallium was based entirely on the substitution of the method detection limit (1 mg/kg) for the undetected values and likely provides a significant overestimation of risk. In addition, the detection limit (1 mg/kg) was less than the Ontario Ministry of the Environment regulatory soil chemical standard for sensitive sites of 2.5 mg/kg (MOE, 2004b). This also suggests that the elevated HQ values observed in this assessment for thallium for the resident toddler are likely due to conservative model assumptions applied throughout the risk assessment process.

3.1.3.3. Farmer Infant. The multi-pathway assessment for exposure of the farmer infant receptor to COPC also suggested potential risks may exist from exposure to baseline concentrations of PCBs, dioxins/furans, and 1,2,4-trichlorobenzene (Table 6, HQ values of 118, 20, and 0.21, respectively). However, as was noted for the resident infant receptor, PCBs and the majority of dioxins/furans were not detected in any media relevant to exposure of farmers (i.e., soil, home-grown produce, or farm-raised livestock) (Supporting Information, Section S6). Furthermore, 1,2,4-trichlorobenzene was also not detected in any samples collected in the baseline sampling program (Supporting Information, Section S6). Therefore, these HQ values may also represent a significant

$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	Ratio (CR) Values – 140,000 tpy									
Baseline Project <		24-hour				Annual				
Metals 0.04 0.03 0.04 0.25 0.27 0.03 0.01 0.04 Chlorinated Polycyclic Aromatics 0.001 0.003 0.001 0.003 0.003 0.006 0.001 5.2E-07 0.001 <th>iject Project Process Process Upset ine Upset Project</th> <th>Baseline Project Alone</th> <th>Project</th> <th>Process F Upset F</th> <th>rocess Upset roject</th> <th>Baseline F</th> <th>^oroject Alone</th> <th>Project</th> <th>Process Upset</th> <th>Process Upset Project</th>	iject Project Process Process Upset ine Upset Project	Baseline Project Alone	Project	Process F Upset F	rocess Upset roject	Baseline F	^o roject Alone	Project	Process Upset	Process Upset Project
Chlorinated Polycyclic Aromatics 0.001 0.003 0.001 0.003 0.001 0.006 0.006 0.006 0.006 0.000 0.001	3 0.04 0.25 0.27	0.03 0.01	0.04	0.14 0	0.16	0.12 0	0.003	0.12	0.004	0.12
Chlorinated Monocyclic Aromatics 0.0006 7.5E-05 0.0007 0.0008 0.001 5.2E-07 0.00	003 0.001 0.003 0.004	0.005 0.0006	0.005	0.006 0	.01	0.002 5	9.0E-06	0.002	2.0E-05	0.002
	E-05 0.0007 0.0008 0.001	0.0001 5.2E-07	0.0001	5.0E-06 (0001	0.002 3	3.0E-06	0.002	1.0E-05	0.002
Polycyclic Aromatic Hydrocarbons (PAH) 0.01 6.9E-05 0.01 0.0007 0.01 0.07 0.07 0.002 0.07	E-05 0.01 0.0007 0.01	0.07 0.0002	0.07	0.002 0	.07	0.002 1	1.0E-06	0.002	2.0E-06	0.002
Volatile Organic Chemicals (VOC) 0.55 0.005 0.55 0.05 0.41 0.002 0.41	05 0.55 0.05 0.56	0.41 0.002	0.41	0.02 (.41	0.18 C	0.0002	0.18	0.0005	0.18

overestimation of the actual risk due to the substitution of the MDL for non-detect values.

3.1.3.4. Farmer Toddler. HQ values greater than 0.2 were observed for the farmer toddler receptor for total PCBs, bromoform, carbon tetrachloride, chloroform, dichloromethane, 1,2,4,5-tetrachlorobenzene, 1,2,4-trichlorobenzene, antimony, arsenic, beryllium, thallium, and dioxins/furans (Table 6). When the risks to the farmer toddler from each COPC were apportioned into their respective exposure pathways, it was observed that ingestion of dairy was the primary exposure pathway associated with risks to the farmer toddler (>65% of total exposure for all chemicals except for arsenic for which only 47% of exposure was related to ingestion of dairy). However, none of these chemicals were actually detected in dairy products in the baseline sampling program and risk assessment was performed using the method detection limit. Therefore, as has been observed for other receptors and COPC in this assessment, the hazard quotients resulting from this substitution likely represent overestimations of the true risk. Furthermore, as toddlerspecific ingestion rates for food items produced on farms were not available, child-specific ingestion rates were adopted from US EPA (2005) as a conservative measure that may also have resulted in an overestimate of exposure since ingestion rates are typically proportional to body weight (Health Canada, 2007).

The farmer toddler also received a significant proportion of its exposure to arsenic via soil and dust ingestion (26%). As was previously discussed with respect to the resident toddler, the maximum soil arsenic concentration used for risk characterization in this assessment (8 mg/kg) is within the expected range for uncontaminated soils in Canada and is also less than the Ontario Ministry of the Environment regulatory soil chemical standard for sensitive sites (MOE, 2004b). Therefore, it is not considered likely that soil and dust ingestion will pose significant undue risk with respect to arsenic exposure for any of the human receptors in the LRASA.

3.1.3.5. Additional Risks Related to Swimming and Hunting/Angling. Additional risks from exposure to surface water while swimming, wading or playing in surface water bodies, as well as from engaging in hunting and angling activities within the LRASA were assessed (Table 7). Results of the swimming exposure assessment indicate that the incremental risks associated with exposure to surface water are between one to six orders of magnitude less than the acceptable multi-pathway HQ benchmark of 0.2 (Table 7). When this additional exposure pathway was added to an existing receptor (e.g., the resident Toddler), the only HQ exceedances noted were for COPC that exceeded the regulatory guideline prior to addition of the swimming pathway (Table 7). In contrast, results of the hunter/angler assessment suggested that this pathway alone may be sufficient to increase COPC exposure above the regulatory guideline for arsenic, cadmium, total PCBs and dioxins/furans (Table 7, HQ values of 0.43, 0.46, 0.67, and 0.38, respectively). Some of these contaminants were not detected in small mammals or fish collected in the baseline sampling program (Supporting Information, Section S6), therefore some of the perceived risk may relate to the replacement of non-detect values with the method detection limit. Furthermore, the concentrations of COPC that were detected in fish (PCBs, arsenic, cadmium, and certain dioxins/furans) and small mammals (arsenic and cadmium), were similar to what would be expected at other areas across Ontario and are therefore not unique to this project (Supporting Information, Section S7).

3.1.4. Multi-pathway risk assessment: Carcinogens

The baseline case multi-pathway assessment also provided oral/ dermal lifetime cancer risk (LCR) estimates for all carcinogenic COPC for the defined multi-pathway receptors and for the incremental exposures resulting from recreational swimming and/or hunting/angling (Supporting Information, Section S8). As discussed in Section 3.1.2, there is no acceptable benchmark for comparison of LCR values, as

Concentration Ratio (CR) Values for Baseline and 140,000 tonnes per year operating scenarios at the Maximum Ground Level Concentration. Each value represents the maximum observed CR value for an individual COPC within each chem-

they represent an individual's lifetime cancer risks associated with all potential exposures to a given carcinogenic COPC within the environment. However, the maximum LCR observed under baseline conditions for these COPC was 0.03%, which is much lower than the typical observed rates of cancer in Canada (38% for women and 44% for men) (Canadian Cancer Society, 2007).

3.2. Risk characterization: Construction case

For consideration of the construction case, it was assumed that construction activities would occur intermittently, during daylight hours, over a period of approximately 30 months. The primary concerns related to these activities with respect to human health were considered to be dust emissions from construction activities and exhaust emissions from fuel combustion by vehicles on the site. In addition, construction activities such as welding, use of solvents, sand blasting and painting may also affect air quality in the construction area. However, relative to the anticipated operational emissions, construction emissions will be minor, short-term and transitory. Therefore, it was expected that the assessment of operational scenarios (Sections 3.3-3.4) will be protective of any potential health risks that could arise during periods of construction and this case was not assessed in detail.

3.3. Risk characterization: Operational scenarios (140,000 tonnes per year)

3.3.1. Inhalation risk assessment: Non-carcinogens

For CACs, predicted maximum 1-hour, 24-hour and annual air concentrations for predicted operational scenarios at 140,000 tonnes per year (i.e. Project Alone Case, Project Case, Process Upset Case or Process Upset Project Case) did not exceed their relevant exposure limits (Table 4); therefore, no adverse health risk is expected from potential exposure to CACs. Additionally, when predicted CAC concentrations were compared to WHO benchmarks for informational purposes, no exceedances were noted for any of the considered assessment scenarios, except for PM2.5 in the Process Upset Project Case (CR = 1.01, Table 4). The exceedance of fine particulate matter is driven by baseline concentrations as the CR for baseline conditions alone is 0.82, while the CR for process upset conditions is only 0.21 (Table 4). However, the baseline concentration of PM_{2.5} in this area is similar to other urban areas in Ontario (Supporting Information, Section S7). In addition, frequency analysis of the baseline monitoring performed as part of this assessment showed that 24-hour PM₂₅ concentrations exceeding the WHO benchmark of 25 μ g/m³ are very rare (Supporting Information, Section S9). No exceedance was noted in comparison to the selected 24-hour PM2.5 Canada-Wide Standard (Table 4).

In addition, for the CACs, the Traffic Case (which combined emissions from offsite and onsite traffic with the anticipated onsite stationary source emissions for the facility) was contrasted with the baseline traffic case. In this case, the predicted 1-hour, 24-hour and annual air concentrations for the CAC at 140,000 tonnes per year did not exceed their relevant exposure limit for either the Baseline Traffic Case, or the Traffic Case (Supporting Information, Section S8). Therefore, no adverse health risk is expected from potential exposure to CACs due to the combined effect of facility emissions at 140,000 tonnes per year and local vehicular traffic. When compared to WHO benchmarks for informational purposes, an exceedance was noted for annual nitrogen dioxide (CR = 1.2) for both the baseline traffic case and the traffic case (Supporting Information, Section S8). However, as discussed in Section 3.1.1, this exceedance was driven by baseline concentrations, which were within a normal range for an urban area in Ontario (Supporting Information, Section S7). Therefore, this does not represent an unusual level of risk associated with this location.

For remaining COPC, none of the predicted maximum 1-hour, 24-hour or annual air concentrations exceeded their relevant exposure limit for any of the operational scenarios (Table 5).

3.3.2. Inhalation risk assessment: Carcinogens

For all carcinogenic COPC, chronic incremental lifetime cancer risks (ILCR) values were calculated for the 140,000 tonnes per year Project Alone Case and Process Upset Case at the maximum predicted ground level concentration (Supporting Information, Section S8). As outlined in Section 2.10.2, an ILCR less than or equal to 1-in-1,000,000 (i.e., 1×10^{-6}) signifies that the incremental lifetime cancer risk is less than the regulatory benchmark (i.e., the assumed safe level of exposure); therefore, no adverse risk is expected. Conversely, an ILCR greater than 1 x 10⁻⁶ indicates that the potential for an elevated level of risk may be present and suggests further investigation should be pursued to confirm the identified risk. In this assessment, none of the predicted ILCR exceeded the regulatory benchmark for the carcinogenic COPC in either the Project Alone Case or Process Upset Case (Supporting Information, Section S8). Therefore, it is not expected that concentrations of carcinogenic COPC from the facility at 140,000 tonnes per year will pose any individual adverse carcinogenic risk to the health of human receptors via inhalation.

3.3.3. Multi-pathway risk assessment: Non-carcinogens

For most receptors, COPC, and operational scenarios, the HQ values did not exceed the regulatory benchmark of 0.2 (Tables 6, 7). The only exceedances noted were for operational scenarios that also incorporated the baseline conditions (i.e., the Project Case and Process Upset Project Case). In these cases, the source of the exceedance was always the baseline case. For instance, for the local resident infant and toddler receptors neither the Project Alone Case nor the Process Upset Case ever represented more than approximately 0.5% of the Project Case or Process Upset Project Case risk, respectively. Similarly, for the farmer infant and toddler receptors, the Project Alone Case or Process Upset Case never represented more than approximately 2% of the Project Case or Process Upset Project Case risk, respectively.

As discussed in Section 3.1.3, the exceedances observed in the baseline conditions were related to a number of issues such as the use of laboratory method detection limits as environmental media concentrations and the conservative nature of risk assessment exposure calculations. In addition, some COPC concentrations actually exceeded relevant guidelines in specific media. However, the baseline COPC concentrations were found to be no different in the LRASA than in other similar areas of Ontario and are therefore not unique to this project.

3.3.4. Multi-pathway risk assessment: Carcinogens

Incremental lifetime cancer risks (ILCR) were estimated for all receptors under the Project Alone Case and Process Upset Case assessment scenarios (Supporting Information, Section S8). In addition, activity specific ILCR values were calculated with respect to hunting/ angling and swimming and were added to that of the worst case resident receptor. None of the predicted ILCR values exceeded the accepted regulatory benchmark for the Project Alone Case or Process Upset Case; therefore, it is not expected that the facility will pose any additional adverse cancer risk to the health of local receptors at 140,000 tonnes per year.

3.4. Risk characterization: Operational scenarios (400,000 tonnes per year)

For comparison purposes, a human health risk assessment was also performed that considered the possible expansion of the facility to its maximum design operating capacity of 400,000 tonnes per year. This assessment was performed using identical methods and assumptions as those described for the 140,000 tonnes per year assessment, except that the facility related emissions were increased. Most of the conclusions of this assessment were similar to those identified for operational scenarios at 140,000 tonnes per year (i.e., most observed risks were related to existing baseline conditions rather than facility-related emissions). However, in the Process Upset Case,

Table 6

b.

Summary of Multi-Pathway Risk Assessment Hazard Quotient (HQ) Results for Baseline and 140,000 tonnes per year operating scenarios for a. the worst-case resident infant and toddler and b. farmer infant and toddler receptors. Each value represents the maximum observed HQ value for an individual COPC within each chemical class. A bolded cell indicates exposure for that particular scenario and COPC exceeded the selected benchmark.

d.										
	Worst-case	e resident inf	ant			Worst-case	e resident too	ldler		
	Baseline	Project Alone	Project	Process Upset	Process Upset Project	Baseline	Project Alone	Project	Process Upset	Process Upset Project
PAHs Maximum observed	6.3E-06	3.4E-11	6.3E-06	9.6E-11	6.3E-06	2.0E-05	5.7E-10	2.0E-05	1.6E-09	2.0E-05
PCBs Aroclor 1254 (Total PCBs)	10.8	0.0003	10.8	0.0008	10.8	0.49	3.4E-05	0.49	9.6E-05	0.49
VOCs Max	0.0002	1.0E-12	0.0002	2.8E-12	0.0002	0.03	2.7E-09	0.03	7.6E-09	0.03
Chlorinated Monocyclic Aromatics Maximum observed	0.003	1.2E-08	0.003	3.4E-08	0.003	0.06	1.2E-07	0.06	3.5E-07	0.06
<i>Inorganics</i> All except Arsenic and Thallium Arsenic Thallium	0.02 0.10 0.05	4.0E-05 5.0E-07 0.0004	0.02 0.10 0.05	5.9E-05 7.3E-07 0.0006	0.02 0.10 0.05	0.07 0.32 0.25	0.0002 3.2E-06 0.002	0.07 0.32 0.25	0.0004 4.6E-06 0.003	0.07 0.32 0.26
Dioxins/Furans and Lead 2,3,7,8-TCDD Equivalent Lead	3.8 0.04	0.002 0.0002	3.8 0.04	0.004 0.0002	3.8 0.04	0.17 0.12	0.0002 0.0005	0.17 0.12	0.0006 0.0007	0.17 0.12

	Farmer inf	ant				Farmer too	ddler			
	Baseline	Project Alone	Project	Process Upset	Process Upset Project	Baseline	Project Alone	Project	Process Upset	Process Upset Project
PAHs Maximum observed	6.8E-06	4.7E-11	6.8E-06	1.3E-10	6.8E-06	5.8E-05	1.5E-09	5.8E-05	4.1E-09	5.8E-05
PCBs Aroclor 1254 (Total PCBs)	117.5	0.004	117.5	0.01	117.5	4.2	0.0001	4.2	0.0004	4.2
VOCs 1,1,1-Trichloroethane Bromoform Carbon Tetrachloride Chloroform Dichloromethane Trichlorofluoromethane	1.8E-07 6.6E-05 0.003 3.1E-05 2.8E-05 5.9E-06	1.6E-14 4.4E-11 4.0E-11 2.3E-13 2.1E-12 1.2E-11	1.8E-07 6.6E-05 0.003 3.1E-05 2.8E-05 5.9E-06	4.6E-14 1.2E-10 1.1E-10 6.4E-13 6.0E-12 3.4E-11	1.8E-07 6.6E-05 0.003 3.1E-05 2.8E-05 5.9E-06	0.0006 0.32 4.6 0.32 0.65 0.02	5.1E-11 1.9E-07 6.3E-08 2.0E-09 4.9E-08 3.8E-08	0.0006 0.32 4.6 0.32 0.65 0.02	1.4E-10 5.3E-07 1.8E-07 5.6E-09 1.4E-07 1.1E-07	0.0006 0.32 4.6 0.32 0.65 0.02
Chlorinated Monocyclic Aromatics Maximum observed (excepting 1,2,4,5-Tetrachlorobenzene and 1,2,4-Trichlorobenzene) 1,2,4,5-Tetrachlorobenzene 1,2,4-Trichlorobenzene	0.03 0.02 0.21	4.0E-08 1.6E-08 1.7E-10	0.03 0.02 0.21	1.1E-07 4.4E-08 4.8E-10	0.03 0.02 0.21	0.17 0.40 20.1	3.2E-07 2.4E-07 1.3E-08	0.17 0.40 20.1	9.0E-07 6.8E-07 3.7E-08	0.17 0.40 20.1
Inorganics Maximum observed (excepting antimony, arsenic, beryllium, and thallium) Antimony	0.02	4.2E-05	0.02	6.1E-05	0.02	0.18	0.0006 8 3E-05	0.18	0.0009	0.18
Arsenic Beryllium Thallium	0.001 0.001 0.005	7.0E-07 6.6E-07 0.0006	0.10 0.001 0.05	1.0E-06 9.6E-07 0.0008	0.01 0.001 0.005	0.57 0.42 1.2	7.6E-06 2.8E-06 0.01	0.24 0.57 0.42 1.2	1.1E-05 4.1E-06 0.02	0.24 0.57 0.42 1.2
Dioxins/Furans and Lead 2,3,7,8-TCDD Equivalent Lead	20.3 0.04	0.05 0.0002	20.3 0.04	0.13 0.0003	20.4 0.04	0.72 0.20	0.002 0.0010	0.72 0.20	0.004 0.001	0.73 0.20

slightly elevated potential risks above the government benchmarks for human health were noted that were not explained by baseline conditions. Maximum exposure to the 1 hour hydrogen chloride concentration at the commercial/industrial receptor location resulted in a CR of 1.0 (benchmark CR = 1.0) and exposure of farmer infant to breast milk of a mother living in close proximity to the facility under the Process Upset Case resulted in an infant dioxin and furan HQ of 0.22, which was slightly in excess of the government benchmark of 0.2. However, these slight exceedances of benchmark risk levels were seen only under upset conditions, it is possible that they may be prevented through the application of adequate engineering controls. Regardless, in the event that a 400,000 tonnes per year expansion of the facility is eventually contemplated, special consideration should be given at that time to ensure that Process Upset Conditions do not result in an undue risk to people living and working in the area surrounding the facility. Overall, the results suggest that a 400,000 tonnes per year facility could be safely sited in Clarington, Ontario using the pollution control technology suggested by Covanta.

Table 7

2

b.

Summary of multi-pathway risk assessment hazard quotient (HQ) results for baseline and 140,000 tonnes per year operating scenarios for additional exposure via a. swimming and b. hunting/angling. The results of adding these exposure pathways to the worst case resident toddler are also shown. Each value represents the maximum observed HQ value for an individual COPC within each chemical class. A bolded cell indicates exposure for that particular scenario and COPC exceeded the regulatory benchmark.

	Hazard qu	otients for	swimming	exposure al	one (toddler)	Swimmin	g exposure	added to w	orst case re	sident toddler
	Baseline	Project Alone	Project	Process Upset	Process Upset Project	Baseline	Project Alone	Project	Process Upset	Process Upset Project
PAHs Maximum observed	1.2E-06	2.8E-11	1.2E-06	7.8E-11	1.2E-06	2.1E-05	5.7E-10	2.1E-05	1.6E-09	2.1E-05
PCBs Aroclor 1254 (Total PCBs)	0.03	6.8E-07	0.03	1.9E-06	0.03	0.52	3.5E-05	0.52	9.8E-05	0.52
<i>VOCs</i> Maximum observed	0.001	2.1E-08	0.001	5.8E-08	0.001	0.03	2.6E-08	0.03	7.3E-08	0.03
Chlorinated Monocyclic Aromatics Maximum observed	0.0007	1.1E-07	0.0007	3.0E-07	0.0007	0.06	2.3E-07	0.06	6.5E-07	0.06
Inorganics Maximum observed excepting arsenic, cadmium. and thallium	0.02	1.3E-05	0.02	1.9E-05	0.02	0.07	0.0002	0.07	0.0003	0.07
Arsenic Cadmium Thallium	0.01 0.0003 0.005	2.7E-06 2.6E-05 0.001	0.01 0.0003 0.006	3.9E-06 3.8E-05 0.001	0.01 0.0003 0.008	0.33 0.03 0.26	5.8E-06 0.0003 0.003	0.33 0.03 0.26	8.5E-06 0.0004 0.004	0.33 0.03 0.26
Dioxins/Furans and Lead 2,3,7,8-TCDD Equivalent Lead	0.003 0.0008	2.8E-07 2.3E-05	0.003 0.0008	8.0E-07 3.4E-05	0.003 0.0008	0.17 0.12	0.0002 0.0005	0.17 0.12	0.0006 0.0007	0.17 0.12

	Hazard quotients for hunter/angler exposure alone (toddler)						Hunter/angler exposure added to worst case resident toddler					
	Baseline	Project Alone	Project	Process Upset	Process Upset Project	Baseline	Project Alone	Project	Process Upset	Process Upset Project		
PAHs Maximum observed	2.1E-05	3.4E-12	2.1E-05	9.6E-12	2.1E-05	4.1E-05	5.7E-10	4.1E-05	1.6E-09	4.1E-05		
PCBs Aroclor 1254 (Total PCBs)	0.67	0.002	0.67	0.006	0.68	1.20	0.002	1.20	0.006	1.20		
<i>VOCs</i> Maximum observed	-	6.2E-09	-	1.7E-08	_	0.03	6.2E-09	0.03	1.7E-08	0.03		
Chlorinated Monocyclic Aromatics Maximum observed	0.06	8.3E-06	0.06	2.3E-05	0.06	0.11	8.4E-06	0.11	2.4E-05	0.11		
Inorganics Maximum observed excepting arsenic, cadmium and thallium	0.16	0.001	0.16	0.002	0.16	0.17	0.001	0.17	0.002	0.17		
Arsenic Cadmium Thallium	0.43 0.47 0.17	3.3E-05 0.008 0.002	0.43 0.47 0.17	4.7E-05 0.01 0.003	0.43 0.48 0.17	0.75 0.49 0.42	3.6E-05 0.008 0.004	0.75 0.50 0.42	5.2E-05 0.01 0.006	0.75 0.50 0.43		
Dioxins/Furans and Lead 2,3,7,8-TCDD Equivalent Lead	0.38 0.04	0.002 0.0006	0.38 0.04	0.005 0.0009	0.38 0.04	0.54 0.15	0.002 0.001	0.54 0.15	0.005 0.002	0.55 0.15		

3.5. Risk characterization: Decommissioning and abandonment

Decommissioning and abandonment of the facility is not expected to occur for several decades. Similar to the construction case, it is expected that this process would entail short-term, localized emissions of air contaminants. While it is unlikely that these activities would significantly increase any potential risk to human health, it is expected that a more current assessment of these potential risks would be conducted prior to the commencement of decommissioning activities. Consequently, the prediction of risks to human health from decommissioning and abandonment were not undertaken in this assessment.

4. Uncertainty Analysis

As part of this risk assessment, it was necessary to make certain assumptions in order to be able to quantitatively evaluate the risks to human health from exposure to the Project. These assumptions inherently add an element of uncertainty to the risk assessment. Where variability and uncertainty are known to exist, it is standard risk assessment practice to make assumptions and select data that are likely to overestimate, rather than underestimate, potential exposure and effects. As a result, risk assessments tend to overstate the actual level of risk. Some of the conservative assumptions applied in this risk assessment include the use of method detection limits to represent chemical concentrations and use of child-specific ingestion rates to represent toddler rate of ingestion. A full accounting of the assumptions and uncertainties relied upon in this HHRA is provided in the Supporting Information (Section S10).

5. Conclusions

Overall, the results of the human health risk assessment indicate that it is not expected that the proposed project (i.e., construction, operation, and eventual decommissioning of a modern EFW thermal
treatment facility) will result in any adverse health risk to local residents, farmers or other receptors in the Local Risk Assessment Study Area at 140,000 tonnes per year. Although some risk has been identified through the assessment of Baseline Case concentrations, this risk can be attributed to conservative modeling assumptions that overestimate the actual risk present (e.g., use of method detection limits to represent chemical concentrations and use of child-specific ingestion rates to represent toddler rate of ingestion) and/or pre-existing natural or anthropogenic conditions that correlate to baseline risk. These pre-existing natural or anthropogenic conditions were generally shown not to differ from those of similar urbanized areas in Ontario.

Based on the success of this human health risk assessment and an accompanying ecological risk assessment (see Ollson et al., 2014), the regions of Durham and York were able to move forward with this project, and the described facility is currently under construction, with operational start-up anticipated in Fall 2014. This facility will be capable of processing 140,000 tonnes of post-diversion residual waste annually while recovering metals and energy.

Conflict of interest

The authors have no actual or potential conflicts of interest to declare.

Acknowledgements

The authors wish to acknowledge the contribution of Greg Crooks and his Air Quality team at Stantec who provided the required air input data. We would also like to thank the Chair Anderson, Council, and Cliff Curtis and his staff at Durham Region; without them this project would not have been undertaken. The direction of the overall environmental assessment was provided by James McKay, now with HDR Canada.

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.scitotenv.2013.07.019.

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MEMO

Job Client Memo no. Date To From Copy to Bromine in Waste EfW Facility TNG NSW 5 2016-10-14 To Whom it may concern Ahmet Erol (Ramboll) I an Malouf (DADI) Phill Andrew (Savills) Amanda Lee (AECOM) Lesley Randall (AECOM) Rachael Snape (Urbis) Damon Roddis (Pacific Environment)

Bromine Emissions from WtE

Background

The most common use of brominated flame retardants (BFRs) is in building materials, textiles and electronic supplies, e.g. TVs, PCs and photocopiers. In incineration plants with good combustion BFRs will decompose and form other brominated compounds (Söderström, G. et al, 2000), mainly hydrogen bromide (HBr) (Vehlow, J. et al, 1998).

In addition, other brominated compounds will also be formed, in particular brominated organic compounds, such as dioxins where chlorine is fully or partly substituted by bromine (brominated and brominated/chlorinated dioxins).¹

Characterization of the brominated waste

Data from literature regarding bromine content in municipal waste from households and small businesses indicates typical bromine content of 0,003-0,006 % by weight of bromine.

Floc waste TNG

Analysis of 17 floc samples from TNG facility done by HRL Technology shows that the average bromine content is 0,01 % on dry basis (db). Maximal bromine content was 0,04 % (db). TNG has 14,4% floc waste in the design waste.

Increase of bromine content by floc waste

Assuming an average content of 0.0045% bromine (average of reported minimum and maximum content of 0.003% and 0.006%) in all waste streams except floc waste and 0,01% bromine in floc this results in an increase of bromine in the total waste from 0.0045% to 0.00529%. The final concentration of 0.00529% is still within the reported range of MSW.

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¹ Emission Measurements During Incineration of Waste Containing Bromine, TemaNord 2005: 529



Effect of increased bromine content in waste

The most extensive measurements were performed at the largest municipal waste incineration plant in Oslo (Klemetsrud Plant). The plant has two incinerator lines, each with the capacity of incinerating 10 tons of waste per hour. Each line is equipped with a flue gas cleaning system, consisting of a bag house filter with active coal injection, and a wet scrubber.

At this plant sampling and analysis were carried out in three different situations:

- No addition of brominated waste
- Mix with 5 % by weight brominated waste; i.e. approximately 0,05 % by weight bromine in total waste.
- Mix with 10 % by -weight brominated waste; i.e. approximately 0,1 % by weight bromine in total waste.

Results

The in stack (after flue gas treatment) concentration of gaseous bromine (HBr and Br_2) was reported to be < 2,2 mg/Nm³ even in the case of addition of 10% brominated waste.

The in stack concentration of BFR in case of the Klemetsrud Plant was 14-22 ng/Nm³, in case of the Energos Plant (Ranheim) <5 ng/Nm³. The BFR detected where DekaBDE and TBBPA, in the flue gas DekaBEDE has the highest concentration level.

Brominated waste has no adverse effect on dioxin formation nor on additional formation of brominated and chlorinated/brominated dioxins.

Conclusions

Using floc waste as a fuel has no adverse effect on the emissions of a WtE facility.

In case of TNG (with a comparable flue gas cleaning technology as the Energos plant) the in stack concentrations for gaseous bromine can be assumed to be below 2 mg/Nm³, BFR below 5 ng/Nm³ and total brominated dioxins far below 0,05 ng/Nm³.

Reference (attached):

Emission Measurements During Incineration of Waste Containing Bromine, TemaNord 2005:529, © Nordic Council of Ministers, Copenhagen 2005

Emission Measurements During Incineration of Waste Containing Bromine

TemaNord 2005:529

Emission Measurements During Incineration of Waste Containing Bromine

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Nordic Environmental Co-operation

Environmental co-operation is aimed at contributing to the improvement of the environment and forestall problems in the Nordic countries as well as on the international scene. The co-operation is conducted by the Nordic Committee of Senior Officials for Environmental Affairs. The co-operation endeavours to advance joint aims for Action Plans and joint projects, exchange of information and assistance, e.g. to Eastern Europe, through the Nordic Environmental Finance Corporation (NEF-CO).

Nordic Co-operation

Nordic co-operation, one of the oldest and most wide-ranging regional partnerships in the world, involves Denmark, Finland, Iceland, Norway, Sweden, the Faroe Islands, Greenland and Åland. Co-operation reinforces the sense of Nordic community while respecting national differences and similarities, makes it possible to uphold Nordic interests in the world at large and promotes positive relations between neighbouring peoples.

Co-operation was formalised in 1952 when *the Nordic Council* was set up as a forum for parliamentarians and governments. The Helsinki Treaty of 1962 has formed the framework for Nordic partnership ever since. The *Nordic Council of Ministers* was set up in 1971 as the formal forum for co-operation between the governments of the Nordic countries and the political leadership of the autonomous areas, i.e. the Faroe Islands, Greenland and Åland.

Content

Preface	7
Abstract	9
1. Background and Objective	15
2. Terms and Abbreviations	17
3. Incineration of Plastics Containing Brominated Flame-Retardants	19
3.1 Plastics from EE-waste	19
3.2 Emissions of brominated flame-retardants	20
3.3 Formation and emissions of chlorinated, brominated and brominated/chlorinated dioxins	20
4. Former Incineration Tests	23
4.1 Brominated and brominated/chlorinated dioxins	23
4.1.1 Incineration tests in pilot plants	23
4.1.2 Measurements at full scale plants	25
4.1.3 Comparison of test programs from earlier tests	25
4.2 Brominated flame-retardants	27
4.3 Studies of other bromine compounds at Klemetsrud Plant, Norway	29
5. Incineration Tests at Three Norwegian Waste Incineration Plants	31
5.1 Measurement program	31
5.2 Characterization of the brominated waste	32
5.3 Description of plants and sampling points	34
5.3.1 Klemetsrud Plant	34
5.3.2 Energos Plant	37
5.3.3 FREVAR Plant	38

6.Results	39
6.1 Operating conditions	39
6.2 Measurements of gaseous bromine in flue gas before cleaning	39
6.3 Mass balance for bromine	40
6.4 Brominated, chlorinated and brominated/chlorinated dioxins	41
6.5 Brominated flame-retardants (BFRs)	43
Sammendrag	47

Preface

Background

The most common use for BFRs is in building materials, textiles and electronic supplies, e.g. TVs, PCs and photocopiers. In incineration plants with good combustion BFRs will decompose and form other brominated compounds, mainly hydrogen bromide (HBr).

In addition, other brominated compounds will also be formed, in particular brominated organic compounds, such as dioxins where chlorine is fully or partly substituted by bromine (brominated and brominated/chlorinated dioxins).

There have been few Studies regarding incineration of plastics containing BFRs at full-scale incineration plants with a modern flue gas cleaning system.

The Norwegian Ministry of the Environment presented in the autumn of 2002 a working plan for reducing the emissions and discharges of BFRs. One action is to investigate the emissions from incineration of waste containing BFRs.

The project has been organised as follows:

Management group:

- Håkon Jentoft, Norwegian Solid Waste Assosiation (NRF)
- Bernt Ringvold, Norwegian Pollution Control Authority (SFT)
- Ole Viggo Svendsen, Elektronikkretur AS
- Tor Christian Svendsen, Hvitevareretur AS
- Hallgeir Betele, Renas
- Fredrik Eide Aas, Stena Miljø AS
- Gerhard Dürbeck, Oslo kommune renovasjonsetaten
- Nordic working group for Thermal Treatment

Reference group:

- NRF's working group for Thermal Treatment
- NRF has been the secretary for the project.

The work has been done by Kjelforeningen Norsk Energi. Authors are Dag Borgnes and Bente Rikheim.

Abstract

Project objective

The objective of the project is to investigate the emissions of dioxincompounds that may occur from incineration of plastic waste containing brominated flame retardants (BFRs) together with waste from households and the commercial sector. The decomposition of BFRs will also be investigated.

The project results will serve as a basis for both the authorities and the owners of incineration plants, to make decisions about whether, and under what conditions, this type of waste may be incinerated.

Literature search and initial studies

The objective of the literature search and initial studies is to establish a detailed program for measurements. It should also be the basis for comparison and evaluation of the results from the measurements.

Studies in small-scale pilot plants

Incineration tests with waste containing BFRs have been carried out in small-scale pilot plants in Sweden (University of Umeå) and in Germany (TAMARA Plant).

The results from Sweden, where the content of bromine was increased up to 1-2 % by weight, showed that the concentration of halogenated dioxins in untreated flue gas was significantly higher with BFRs than without.

At the TAMARA-Plant, the content of bromine varied from 0 to approximately 0,2 % by weight. Increasing the content of bromine showed no increase in the concentration of chlorinated dioxins, or in brominated or brominated/chlorinated dioxins in untreated flue gas.

Measurements on full-scale plants

Studies of emissions of brominated dioxins to air were earlier carried out on incineration plants in Denmark, Sweden and Norway. Measurements performed in Denmark also included brominated/chlorinated dioxins. All plants were equipped with advanced flue gas treatment systems. Measurements were performed during incineration of waste from households and the commercial sector (waste with low BFR content), and results showed very low levels for all analysed dioxins.

There is little relevant data of emissions of BFRs from waste incineration plants. We have found results from emission measurements carried out at a Japanese incineration plant burning plastic waste containing BFRs, mixed with waste from households and the commercial sector. Total input of BFRs was less than 500 g/hr, and the emission to air of PBDE (polybrominated diphenyl ethers) and TBBPA (tetrabrombisphenol) was respectively 3,5 and 8 ng/Nm³.

Incineration tests at three Norwegian plants

The main goal of the incineration tests was to establish the flue gas concentration of brominated, chlorinated and brominated/chlorinated dioxins before and after flue gas cleaning, and with different proportions of plastic waste containing BFRs. To verify the input, the contents of bromine and chlorine in all output flows (bottom ash, fly ash, scrubber water and flue gas) were analysed. The decomposition of BFRs was investigated by analysing BFRs in output flows.

Execution of tests

The incineration test included sampling and analysis at two larger plants for mixed municipal waste, and one smaller plant for ground/shredded industrial waste. The brominated waste added was waste from a plant for demolition of electric and electronic devices. It was estimated to contain approximately 1 % by weightbromine. Approximately 80% of this contained PBDE.

The most extensive measurements were performed at the largest municipal waste incineration plant in Oslo (Klemetsrud Plant). The plant has two incinerator lines, each with the capacity of incinerating 10 tons of waste per hour. Each line is equipped with a flue gas cleaning system, consisting of a bag house filter with active coal injection, and a wet scrubber.

At this plant sampling and analysis were carried out in three different situations:

- No addition of brominated waste
- Mix with 5 % by weight brominated waste; i.e. approximately 0,05 % by weight bromine in total waste.
- Mix with 10 % by -weight brominated waste; i.e. approximately 0,1
 % by weight bromine in total waste.

At the second plant (FREVAR Plant, Fredrikstad) measurements were carried out with no addition of BFRs.

At the third and smaller plant (Energos Plant, Ranheim) measurements were performed incinerating a mix with 0 and 20 % by weight bromine containing waste (i.e. 0,2 % by weight bromine in the total mix).

Results and conclusions

The incinerating conditions during sampling and measurements at Klemetsrud Plant (Oslo) were normal for the plant, with average CO-levels at approximately 20-30 mg/Nm³. During sampling at the FREVAR Plant average CO-levels were approximately 50 mg/Nm³. At FREVAR Plant they also experienced some problems with the fabric filters during the measurements.

At the Energos Plant (Ranheim) CO was not detectable, which indicates that incineration was good.

Bromine in output flows

Measured results for gaseous bromine in untreated flue gas during incineration of normal waste mix indicates a bromine content equal to or lower than what is common for waste from households and the commercial sector.

Measured results of bromine in output flows at Klemetsrud Plant (Oslo) and at the Energos Plant (Ranheim) indicate that the content of bromine in the plastic mixture was correctly estimated.

Brominated flame retardants (BFRs)

The amount of BFRs in the waste mixture used in the tests at Klemetsrud Plant (Oslo) was not analysed, but calculated/estimated to be approximately 30 kg/hr. The measured results confirm that BFRs decompose in the incineration process. The amount of BFRs in output flows is less than 0,001 % by weight of the total amount of BFRs in the waste mix (see figure below).



Observed input and output flows of brominated flame retardants at Klemetsrud Plant (Oslo) with 10 % by weight addition of brominated waste.

The concentration of BFRs in flue gas from Klemetsrud Plant (Oslo) was 14-22 ng/Nm³. This equals 0,9-1,4 mg/hour and approximately 0,01 kg/year, assuming 8000 running hours/year at the same emission level. A Danish study (Miljøstyrelsen, 1999) estimates the total national Danish emissions of BFRs from incineration to be < 0,04 tons. A report from the Norwegian National State Pollution Control Authority (SFT), estimates the national emissions from combustion in Norway to be < 0,01 tons (1998), i.e. < 10 kg/year.

At the Energos Plant (Ranheim) the reported concentration of BFRs in the flue gas was $<5 \text{ ng/Nm}^3$.

The concentration of BFRs in bottom ash from the tests at Klemetsrud Plant (Oslo) shows levels far below the threshold value stated in the Hazardous Waste Directive.

DekaBEDE and TBBPA (Tetrabrombisphenol A) are the dominating compounds of BFRs in the bottom ash at Klemetsrud Plant (Oslo). In the flue gas dekaBEDE has the highest concentration level.

Concentration of dioxins in emissions to air (after cleaning)

The figure below shows emissions of chlorinated, brominated and chlorinated/brominated dioxins without any addition of brominated waste, and with the addition of 5 % by weight, 10 % by weight and 20 % by weight bromine containing waste.



Emissions of chlorinated, brominated and chlorinated/brominated dioxins. The results are reported as actual emission, not toxic equivalents.

Emissions of chlorinated dioxins (PCDDs/Fs), in terms of Nordic toxic equivalents, resulting from the addition of brominated waste, are presented in the figure below.

Emissions of chlorinated dioxins (PCDDs/Fs), in terms of Nordic toxic equivalents, resulting from addition of brominated waste.



EU threshold value

Uncertainty in sampling and analysis, variations in operating conditions and waste mixture, differences between laboratories with respect to methods of analysis (especially dioxins), makes comparison of results difficult. We may although draw the following main conclusions:

- Increasing the content of BFRs in the waste gave no significant increase in the emissions of chlorinated dioxins, or either brominated and chlorinated/brominated dioxins
- The emission level is highest for chlorinated dioxins, lower for chlorinated/brominated dioxins and lowest for brominated dioxins

- The emission levels for chlorinated dioxins, reported as Nordic toxic equivalents, are low compared to emission threshold value in the EU-directive for incineration of waste. The reported emission levels were 0,03 ng/Nm³ and 0,006 ng/Nm³ respectively for the Klemetsrud Plant (Oslo) and Energos Plant (Ranheim), and the EU threshold value is 0,1 ng/Nm³.
- The emission measurement results indicate that the incineration efficiency and the operating conditions of the flue gas treatment systems are of greater importance to the resulting emission levels for dioxins, rather than the bromine content level.

Concentration of dioxins in emissions before and after cleaning

Measurements of dioxins in the flue gas before and after flue gas cleaning were carried out with addition of 10 % by weight bromine containing waste at the Klemetsrud Plant (Oslo)

The concentration of chlorinated/brominated dioxins before cleaning was approximately 28 ng/Nm³, which was three times the concentration of chlorinated dioxins. After cleaning the concentration was approximately 0,1 ng/Nm³. This gives a removal efficiency for chlorinated/brominated dioxins of >99% and for chlorinated dioxins approximately 93%. The removed dioxins end up in the fly ash from the fabric filter, which is treated as hazardous waste.

Concentration of chlorinated, brominated and chlorinated/brominated dioxins in flue gas before and after flue gas cleaning, Klemetsrud Plant (Oslo). The levels are given as actual measured levels, not corrected for toxicity.



1. Background and Objective

The most common use of BFRs is in building materials, textiles and electronic supplies, e.g. TVs, PCs and photocopiers. In incineration plants with good combustion BFRs will decompose and form other brominated compounds (Söderström, G. et al, 2000), mainly hydrogen bromide (HBr) (Vehlow, J. et al, 1998). Additionally, other brominated compounds will also be formed, in particular brominated organic compounds, such as dioxins where chlorine is fully or partly substituted by bromine (brominated and brominated/chlorinated dioxins).

The formation of brominated/chlorinated dioxins during incineration of waste with BFRs has been proven earlier, in the project "Coincineration of brominated flame-retardants and MSW in small-scale reactor" in 2000 (financed by the Nordic PA-group and documented in TEMA-Nord Report No 2001:512). The tests were carried out at Umeå University, in at laboratory pilot plant with no flue gas treatment.

There have been few studies regarding incineration of plastics containing BFRs at full-scale incineration plants with a modern flue gas cleaning system.

The Norwegian Ministry of the Environment presented in the autumn of 2002 a working plan for reducing the emissions and discharges of BFRs. One action is to investigate the emissions from incineration of waste containing BFRs.

The objective of the project is to investigate the emissions of dioxincompounds that may occur from incineration of plastic waste containing brominated flame-retardants (BFRs) together with waste from households and the commercial sector. The decomposition of BFRs will also be investigated.

The project results will serve as a basis for both the authorities and the owners of incineration plants, to make decisions about whether, and under what conditions, this type of waste may be incinerated.

This report is based on separate reports from incineration tests at Klemetsrudanlegget (Oslo) (Kjelforeningen-Norsk Energi, 2004), and at FREVAR (Kjelforeningen-Norsk Energi, 2004b), which also include detailed description of measurement methods and analysis results.

The incineration tests at Energos Ranheim are reported in a report from TÜV Nord Umweltschutz (2003).

2. Terms and Abbreviations

Brominated flame-retardants (BFRs)

Name specific compound	IUPAC-no [*] .	Abbreviation	Abbreviation groupname	Groupname		
ТВА		ТВА	ТВА	Tribromanisol		
4,4'-DiBB	15	DiBB				
2,2',4,5'-TetBB	49	T (DD DDD		Polybrominated		
2,2',5,5'-TetBB	52	TELDD	FBB	Biphenyls		
2,2',4,4',5,5'-HexBB	153	HeksaBB				
2,4,4'-TriBDE	28	TriBDE				
2,2',4,4'-TetBDE	47					
2,3',4',6-TetBDE	71	TetBDE				
3,3',4,4'-TetBDE	77					
2,2',4,4',5-PenBDE	99					
2,2',4,4',6-PenBDE	100	PeBDE		Polybrominated		
2,3',4,4',6-PenBDE	119		FBDL	diphenyl ethers		
2,2',3,4,4',5'-HexBDE	138					
2,2',4,4',5,5'-HexBDE	153	HexBDE				
2,2',4,4',5,6'-HexBDE	154					
2,2',3,4,4',5',6-HepBDE	183	HepBDE				
DecaBDE	209	DecaBDE				
ТВВРА		ТВВРА	ТВВРА	Tetrabrombisphenol A		
alpha-HBCD						
beta-HBCD		HBCD	HBCD	Hexabrom-cyklododecane		
gamma-HBCD]				

* Indexes according to International Union of Pure and Applied Chemistry (IUPAC).

Chlorinated dioxins (PCDD+PCDF)

- PCDD = polychlorinated dibenzo-p-dioxins
- PCDF = polychlorinated dibenzofurans

Brominated dioxines (PBDD+PBDF)

- PBDD = polybrominated dibenzo-p-dioxins
- PBDF = polybrominated dibenzofurans

Brominated/chlorinated dioxins (ClxBryDD+ClxBryDF)

ClxBryDD, PXDD = polychlorinated/brominated dibenzo-p-dioxins ClxBryDF, PXDF = polychlorinated/brominated dibenzofurans

Incineration of Plastics Containing Brominated Flame-Retardants

3.1 Plastics from EE-waste

Brominated flame-retardants are being found in i.e. electric and electronic (EE) products. In Norway and Sweden there is established extensive collection systems for discarded EE products. As a consequence of new EU regulations, similar systems will have to be established in all EU/EEA countries within the end of 2005.

Table 1 shows the amounts of plastics from EE-waste, based on the information from collection companies in Norway (Svendsen, T. C., 2003).

	Separated plastics	Plastics in shredderfluff
Collection company	(tons/år)	(tons/år)
Elektronikkretur AS	1800	180
Hvitevareretur AS	150	4000
RENAS AS Total	45 1995	239 4419

Table 1 Amounts of plastics from EE-waste, based on the information from collection companies (tons/år)

As will be seen from Table 1, approximately 2000 tons separated plastics is generated yearly from EE-waste. This is bigger plastic items with and without BFRs, which relatively easy may be sorted out manually, for example the cover of a data monitor, back-cover of a TV and soap container in a dishwasher. These plastic components will mainly be incinerated in advanced waste incinerators with adequate flue gas cleaning.

The largest amount of plastics will however be found in the so-called shredderfluff, with an amount of approximately 4400 tons/year. Shred-derfluff is the waste fraction from wrecked car/scrap metal fragmenting plants, and it contains (among other things) a mix of plastics, rubber, wood, concrete, and small amounts of metals. In Norway, shredderfluff is normally deposited in landfills.

3.2 Emissions of brominated flame-retardants

The most applied brominated flame-retardants are the group polybrominated diphenylethers (PBDE) and the compunds tetrabromobisphenol A (TBBPA) og hexabromocyclododecane (HBCD).

Figure 1 shows and example of polybrominated diphenylethers (DeBDE), TBBPA and HBCD.

Figure 1 Polybrominated diphenylether (DeBDE), TBBPA og HBCD.



Decabromodiphenyl ether (DeBDE) Hexabromocyclododecane (HBCD)

Tetrabromobisphenol A (TBBPA)

In incinerators with good combustion, the BFRs will decompose and form other brominated compounds (Söderström, G. et al, 2000), mainly hydrogenbromide (HBr) (Vehlow, J. et al, 1998).

3.3 Formation and emissions of chlorinated, brominated and brominated/chlorinated dioxins

Chlorinated dioxins is a collective term for organic compounds consisting of dibenzo-p-dioxins and dibenzofurans with 1-8 chlorine substituents in different positions. This gives a total 210 different plychlorinated dibenzo-p-dioxins (PCDD) and polychlorinated dibenzofurans (PCDF). Chlorinated dioxins and furans are often referred to as "dioxins".

Brominated dioxines is a collective term for the corresponding 210 organic compounds substituted with bromine instead of chlorine. These are normally also reffered to as PBDD og PBDF (polybrominated dibenzo-pdioxins og polybrominated dibenzofurans). *Brominated/chlorinated dioxins* includes dibenzo-p-dioxins and dibenzofurans with both bromine and clorine substituents, in total 4600 different compounds. See figure 2.

Air emissions of chlorinated dioxins are regulated through the standard regulations for incineration, and the emissions are checked every 6 or 12 months. The emission limit value is given in terms of toxic equivalents, which is generated by weighted calculation, giving each compound a relative weight value between 0 and 1, depending on the toxicity. 2,3,7,8-tetrachlorine dibenzo-p-dioxin is known as the most toxic compound, and therefore has the relative weight value 1.

Toxic equivalents terms is however not established for compounds of brominated and brominated/chlorinated dioxins.

Figure 2 Brominated/chlorinated dibenzo-p-dioxins and dibenzofurans



Formation of dioxins

The formation of dioxins during incineration has been studied extensively for nearly 30 years.

Different formation mechanisms has been found:

- De Novo synthesis
- Precursor reactions
- Secondary halogenations in flue gas

In the de Novo synthesis, the formation takes place by chlorination of compounds in flue gas containing carbon. The chlorination step is assumed to occur as HCl combine to form Cl_2 (the Deacon-reaction), with subsequently chlorination of aromatics. The Deacon-reaction is catalysed by (among others) compounds of copper, and is favoured by oxygen excess.

Chlorinated dioxins may also be formed from precursors, for example chlor-phenols, which can condensate on particle surfaces, and also by halogenation of non-chlorinated dioxins and furans in flue gas. Studies have shown that the highest formation of dioxins takes place at temperatures between 200-600° C.

Emissions of dioxins may also occur if the incinerated waste or the added combustion air contains dioxins. To minimize the formation of chlorinated dioxins, one has found the following to be important:

- Short residence time at temperatures favouring dioxin formation
- Efficient combustion
- Minimize chlorine content in waste
- Minimize particle content in flue gas
- Increased sulphur/chlorine content ratio
- Minimize oxygen excess
- Minimize content of metals which can act as catalysers (especially copper)

Several studies have shown that there is no clear relation between the rate of dioxin formation and the chlorine content (SFT, 1994) (Wikström, E., 1999). However, some waste incineration tests indicate that dioxin formation increases with increasing chlorine content, when the chlorine content exceeds a certain value.

Several studies of waste incineration have shown that combustion efficiency is of greater importance for the dioxin formation, rather than the chlorine content.

Formation mechanisms for brominated and brominated/chlorinated dioxins are less investigated. It is however reasonable to assume that mechanism has similarities with the formation mechanism of chlorinated dioxins.

Incineration plants which comply with the regulations of the EUdirective for waste incineration, are all equipped with flue gas cleaning systems which reduce the concentration of chlorinated dioxins in the flue gas substantially. The similarities between chlorinated, brominated and brominated/chlorinated dioxins indicates strongly that the cleaning efficiency is also high for brominated and brominated/chlorinated dioxins.

4. Former Incineration Tests

Swedish, Danish and Finnish environmental authorities has been contacted to get data from incineration tests including measurements of brominated and brominated/chlorinated dioxins.

Information has also been gathered from universities and research communities in Sweden, Denmark and Germany, and also through search on the Internet. Articles from the last three Dioxin conferences (2001, 2002, 2003) are also examined.

The objective of this work has been to establish a detailed measurement program. The possibility to estimate input amounts of bromine from measurements/calculations of output bromine containing flows (bromine in bottom ash, fly ash, flue gas and scrubber water) is also investigated. The work should also form basis for comparison and evaluation of measurement results from the incineration test.

4.1 Brominated and brominated/chlorinated dioxins

4.1.1 Incineration tests in pilot plants

Emissions and formation of brominated dioxins is investigated in a pilot plant in Germany. The TAMARA Plant has a capacity of 250 kgs of waste per hour, and is equipped with textile filter, quenching and a wet scrubber. At this plant incineration tests of polystyrene- and polyurethane foam containing BFRs has been carried out, together with waste from households and smaller industries/businesses (Vehlow, J. et al, 1996).

The incineration temperature was ranging from 850 to 950 °C. The additional inputs of bromine during the tests were ranging up to 6 times the original bromine content in the household waste.

Measurement results showed low concentrations of brominated dioxins, and the study concluded that incineration of limited amounts of specific foams in efficient plants with "state of the art" flue gas cleaning, is environmentally acceptable.

Tests with incineration of plastics from EE-waste, together with waste from households and smaller industries/businesses, have also been carried out in the TAMARA Plant (Vehlow, J. et al, 1997). The tests included four different types of plastics, with different contents of bromine.

Measurements of brominated, chlorinated, and brominated/chlorinated dioxins in flue gas, both prior to and after cleaning, were carried out. The tests also included analysis of bromine content in the plastics, bromine-, chlorine- and antimony-content in bottom ash, fly ash and flue gas before cleaning. Antimony is often added to enhance the effect of BFRs. It was concluded that EE-waste containing bromine and bromine and chlorine did not increase the total formation of dioxins. Figure 3 shows concentration levels of PCDDs/Fs and Cl_xBr_yDDs/Fs in flue gas before cleaning as a function of the bromine content in the waste.

Figuer 3 Concentration levels of PCDDs/Fs and Cl_xBr_yDDs/Fs in flue gas before cleaning as a function of the bromine content in the waste, recorded from incineration tests with EE-waste at TAMARA Plant, Germany (Vehlow, J. et al, 1997).



The formation of brominated dioxins from co-incineration of household waste and brominated flame-retardants is investigated in a 5kW incineration reactor (fluidised bed) in pilot scale at the university of Umeå (Söderström, G. et al, 2000). Different types of flame-retardants were added in amounts corresponding to a "worst case scenario" for batch wise incineration of flame retarded products med BFRs. The incineration temperature was slightly above 800 °C.

The results from the studies in Umeå showed that the formation of halogenated dioxins were much higher when adding BFRs, than with only chlorine present. Additionally, the study showed that bromine caused significant higher formation of halogenated dioxins than the equal amount of chlorine, which is assumed to relate to the ratio between Br₂ and HBr, which again is substantially different from the ratio between Cl₂ and HCl. The conclusion from the study is that batch incineration of wastes containing BFRs should be avoided.

4.1.2 Measurements at full scale plants

In 2002 measurements of emissions of brominated dioxins were carried out at Energos Hurum Plant, Norway (Energos Hurum Energigjenvinning) with normal waste composition (wastes from households and small industries/businesses).

It was found 0,003 ng/Nm³ tetrabrominated dibenzo-p-dioxins. Concentrations of other single compounds were lower than the detection limit, i.e. than 0,0001-0,02 ng/Nm³. Note that the concentrations is given as actual measured values, not as toxic equivalents (Energos ASA, 2002).

In 1999 measurements of brominated dioxins were carried out at Uppsala Energi, Sweden and at Renova, Gothenburg, during incineration of ordinary municipal waste. Both plants are equipped with advanced flue gas cleaning systems, with low emissions of chlorinated dioxins (substantially lower than 0,1 ng/Nm³ in 1999). The emissions of brominated dioxins were lower than the detection limit for the measurements, i.e. < 0,05 ng/Nm³ for all measured dioxins (Westas, H., 2000).

Autumn 2002, Danmarks Miljøundersøgelser carried out a study regarding the content of brominated, chlorinated and brominated/chlorinated dioxins in flue gas and in remains from flue gas cleaning at Vestforbrænding (VF) (wastes from households and small industries/businesses) and Kommunekemi (KK) (hazardous waste) (Vikelsøe, J., 2000). Both plants were equipped with advanced flue gas cleaning systems.

Figure 4 shows comparison of PBDDs/Fs and PCDDs/Fs in flue gas from the plants after cleaning

Figur 4 Comparison of PBDDs/Fs and PCDDs/Fs in flue gas from Vestforbrænding and Kommunekemi (after cleaning). PBDDs/Fs total "B-TEQ" og PCDD/F I-TEQ, ng/Nm³. (Vikelsøe, J., 2000).



4.1.3 Comparison of test programs from earlier tests

Table 2 shows the added amount of bromine/brominecontaining plastics in earlier testprograms in small-scale pilot plants.

Table 2 Former test programs/measurements

	Share BFR-plastics	Bromine content in BFR-plastics	Bromine content in waste
Plant/test description	[% by weight]	[% by weight]	[% by weight]
TAMARA			
Polystyrene- and polyurethanefoam + waste from housholds and small industry/ businesses TAMARA Plastics from EE-waste + waste from housholds and small industry/ businesses Umeå University	1-3 3-12	2,2-4,2 0,4-1,5	0,02-0,08 0,01-0,18
Waste from housholds and small in- dustry/businesses added different BFRs	-	-	0,9-1,7

Table 3 shows the measurements included in incineration tests with brominated dioxins and/or chlorinated/brominated dioxins.

Table 3 Measurements included in incineration tests with brominated dioxins and/or chlorinated/brominated dioxins

	Analyzed parame- ters	Analyzed parame- ters	Analyzed parame- ters	Additional analy- zed parameters
	in waste	Raw flue gas	Clean flue gas	
TAMARA	Bromine, chlorine	PXDDs/Fs		Bromine in fly ash
PS- og PU- foam + waste from housholds and small industry/ businesses				
TAMARA	Bromine, chlori-	Bromine	PCDDs/Fs	Bromine, chlori-
Plastics from EE-waste +	ne, antimony	(HBr,Br ₂), chlori-	Non-brominated	ne, antimony (Sb)
industry/ businesses	and flame- retardants ¹⁾	(Sb), PXDDs/Fs	CIONINS	flyash
Umeå Universitet Waste from housholds and small husinesses added different BERs	Bromine, chlorine and flame- retardants ²⁾	Cl ₂ , HCl, Br ₂ , HBr, PXDDs/Fs ³⁾		
Uppsala Energi,Renova Waste from housholds and small	Tetardants		PXDDs/Fs	
Vestforbrænding Waste from housholds and small			PXDDs/Fs	PXDDs/Fs in residue from flue
industry/businesses KommunekemiHazardous waste			PXDDs/Fs	gas cleaning PXDDs/Fs in residue from flue gas cleaning

0) PXDDs/Fs: Dioxins containing bromine and/or chlorine PBB, PBDE, TBBA DeBDE, TBBP-A, HBCD PCDD, PCDF, TeBCDD, TeBCDF, TeBDD, TeBDD Relatively extensive incineration tests, with different input of bromine, have been carried out at pilot plants both in Sweden and Germany. In both tests analysis were carried out with respect to brominated and brominated/chlorinated dioxins on raw flue gas only (before cleaning).

Analysis of brominated dioxins in flue gas after cleaning is carried out on some Swedish and Danish waste incineration plants, with normal waste composition.

Our literature search and preliminary studies indicated clearly a need for more incineration tests and studies of brominated and brominated/chlorinated dioxins, especially in full scale plants. The scope of former studies indicates that the main goal with such tests should be to decide the concentrations in flue gas of brominated, chlorinated and brominated/chlorinated dioxins before and after cleaning, and at different levels of bromine content in the waste. To verify the input of bromine and chlorine, all output flows (bottom ash, fly ash, water from scrubbers and flue gas) should also be analyzed for bromine and chlorine.

4.2 Brominated flame-retardants

An article presented by Chen, Y. et al (Dioxin 2003) reports from sampling and analysis of emissions of BFRs and brominated dioxins carried out at an incineration plant burning wastes from households and small industry/businesses. The plant is not described in the article, nor the waste or the operating conditions of the plant. The method used to determine BFRs and dioxins is not the same as the methods used in the tests at Klemetsrud Plant, Oslo and Energos Plant, Ranheim. It is carried out five series of measurements of air emissions from a waste incinerator and 3 series from an electric smelter. Average results are quoted in Table 4. Seven congeners of PBDE (BDE-28, -47, -100, -99, -154, -153, -183) was detected in all the samples. The three most dominating congeners is BDE -47, -99 og -28 and both the tests shows equal distribution between the congeners.

Table 4 Emissions of BFRs and brominated dioxins from waste incinerator and electric smelter. (Chen, Y. et al, 2003).

	BFRs	Brominated dioxins
	(ng/m³)	(ng/m³)
Waste incinerator Electric smelter	99±31 68±25	0,275-4,01 0,079-0,485

An article presented by Tamade, Y. et al, Japan (Dioxin 2003) reports from measurements during incineration of plastic waste with BFRs. The measurements include analysis of brominated dioxins and furans, PBDE and TBBPA on the input waste, such as back covers from TVs, dust from TVs, and also in mass flows from a recovery plant for plastics, and finally in mass flow from an incineration plant.

The incineration plant was equipped with an electric precipitator and a fabric filter. The incinerated waste was a mixture of residues from the plastic waste recovery plant (with BFRs) and waste from households and small industry/businesses. The waste was analysed with respect to content of PBDE, TBBPA and brominated dioxins. Total input amount of PBDE and TBBPA were 18-360 g/hr and 6,2-96 g/hr respectively. Analysis of brominated dioxins and furans in air emissions showed a total concentration of 0,014 ng/Nm³. Air emission of PBDE and TBBPA was 3,5 and 8 ng/Nm³ respectively. Bottom ash and filter dust showed a content of PBDE of 300 and 470 ng/g respectively. The content of TBBPA in bottom ash and filter dust was 20 and 1,3 ng/g respectively.

Due to few studies and lack of emission limit values for BFRs from waste incineration, we have also looked at reported concentration levels in other types of samples.

A study of indoor dust in common households in Germany includes analysis of 40 samples (taken from vacuum cleaners) with respect to 10 different PBDE congeners (BDE-28, -47, - 49, -85, -99, -100, -153, -154, -183, -209) (Knoth, W. et al, 2003). The results show huge variations in concentrations between the different congeners, and also between the samples. BDE-209 was the dominating congenere in 35 of the 40 samples, as BDE-99 dominated in 4 of the samples. The source for PBDE in the samples was reported unknown, with exception for some samples of dust from mattresses which showed high levels of dekaBDE. Average total concentration of the 10 PBDEs in the 40 samples was determined to 1404 ng/g.

The Norwegian Institute for Air Research (NILU) has taken samples of sediments for analysis of BFRs in the Drammen river, Norway (Fjeld et al, 2004).

Samples of sediment were taken at seven different spots in the river, four samples from the inner Drammensfjord and one sample in the marine environment of the fjord. The sum of PBDEs analysed showed a concentration level of 4-80 ng/g. The BDE-209 congenere dominated in all samples.

NILU have also made studies of BFRs in leachate from landfills (Schlabach. M. et al, 2002).

Samples were taken from sediments in leachate from 6 larger landfills. PBBs were not detected in any of the samples. PBDE-209 was detected in all samples, with a concentration level in the range of 0,49-91 ng/g wet weight. The three HBCD-isomeres was detected in almost all samples, and the concentrations was in the range <0,1-84 ng/g wet weight for HBCD. TBBPA was detected in all sediment samples from the landfills, with a concentration level in the range of 01,9-44 ng/g wet weight.

PBDE-209 and HBCD are also detected in samples of moss, which implies that the compounds may be transported by air. NILU has estimated that maximum discharge from a larger landfill might rate up to 1-10 g/year per single compound of PBDE, HBCD and TBBPA. The concentrations found in the inestigations are at the same levels as concentrations found in sewage sludge in Sweden.

Table 5	Results	from	different	studies	of	BFRs
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Type of study	BFR-compound	µg/g	ng/m³
Air emissions from waste incineration	PBDE		99
Air emissions from electric smelter	PBDE		68
Dust from households	PBDE	1,4	
River sediments	PBDE	0,004-0,08	
	PBDE	0,0005-0,09	
Sediments from landfill leachate	HBCD	<0,0001-0,08	
	TBBPA	0,001-0,044	
Bottom ash from incinorator Japan	PBDE	0,3	
Bottom ash non incinerator, Japan	TBBPA	0,02	
Elv ash from incinorator Japan	PBDE	0,47	
Fly ash non incinerator, Japan	TBBPA	0,0013	
	PBDE		3,5
Air emissions from incinerator, Japan	TBBPA		8

Our preliminary studies and literature search indicated clearly a need for more incineration tests and studies of emissions and decomposition of BFRs in connection with waste incineration.

4.3 Studies of other bromine compounds at Klemetsrud Plant, Norway

In 2002 measurements of brominated and brominated/chlorinated organic compounds were carried out at the Klemetsrud Plant in Norway. The measurements were performed by Kjelforeningen-Norsk Energi (Kjelforeningen-Norsk Energi, 2002).

In 1998 incineration tests with EE-waste were carried out at the same plant, and online measurements of a variety of brominated components in flue gas were performed (not dioxins). A portable GC was used for the measurements (Det Norske Veritas, 1998). In parallel measurements of emissions of chlorinated dioxins after flue gas cleaning was performed. The results indicated an increased level of dioxins during incineration of EE-waste. It should be noted that this was before active coal injection and fabric filtration were introduced at the plant.

5. Incineration Tests at Three Norwegian Waste Incineration Plants

Measurements of emissions of brominated and brominated/chlorinated dioxins, and brominated flame-retardants (BFRs) are carried out during incineration of waste with both normal and increased content of BFRs in the waste. For verification of input, analysis of bromine and chlorine in output flows were made. Decomposition of BFRs during incineration was also investigated by analysis of BFRs in output flows.

Tests and measurements were carried out at the following Norwegian plants:

- Klemetsrud Plant, Oslo : municipal waste incinerator, capacity 2 x 10 tons/hr
- Energos Plant, Ranheim : municipal waste incinerator*, capacity 1,5 tons/hr
- FREVAR Plant, Fredrikstad : municipal waste incinerator, capacity 2 x 5 tons/hr

* source separated and shredded waste

5.1 Measurement program

The program for measurements during tests at three Norwegian waste incineration plants are shown in Table 6 below.

	Type of waste	Analysed parameters	Analysed parameters	Other analysed parame-
		in flue gas before cleaning	in flue gas after cleaning	
	Waste from households and small industry/businesses	HCI, Cl ₂ , HBr, Br ₂	Chlorinated, brominated and chlorinated / bromina- ted dioxins	-
Klemetsrud- Plant	Waste from households and small industry/businesses + 5 % by weight brominated waste	HCl, Cl ₂ , HBr, Br ₂	Chlorinated, brominated and chlorinated / bromina- ted dioxins	-
	Waste from households and small industry/businesses + 10 % by weight brominated waste	Chlorinated, brominated and chlorinated / bromina- ted dioxins HCI, Cl ₂ , HBr, Br ₂	Chlorinated, brominated and chlorinated / bromina- ted dioxins BFRs	Bromine, Chlorine and BFRs in bottom ash, fly ash, flue gas, scrub- berwater
FREVAR Plant	Waste from households and small industry/businesses Hospital waste	HCI, Cl ₂ , HBr, Br ₂	Chlorinated, brominated and chlorinated / bromina- ted dioxins	-
Energos Plant	Industrial waste	HCI, Cl ₂ , HBr, Br ₂	Chlorinated and bromina- ted dioxins	-
	Industrial waste +		Chlorinated, brominated	Bromine, Chlorine and
	20 % by weight brominated waste	HCI, Cl ₂ , HBr, Br ₂	and chlorinated / bromina- ted dioxins	BFRs in bottom ash
			HCI, Cl ₂ , HBr, Br ₂	
			BFRs	

Table 6 Program for measurements during tests at three Norwegian waste incineration plants

Sampling and analysis of dioxins during tests with no addition of brominated waste at Klemetsrud Plant, and all sampling and analysis at Energos Plant, were performed by the German consultancy TÜV. Kjelforeningen-Norsk Energi did all other sampling at Klemetsrud Plant, and at FREVAR Plant. NILU laboratory made the analysis of dioxins and BFRs, Eurofins laboratory (Oslo) analysed the flue gas samples, and Analytica laboratory made the bottom ash, filter dust and scrubber water analysis.

5.2 Characterization of the brominated waste

Data from literature regarding bromine content in municipal waste from households and small businesses, indicates typical bromine content of 0,003-0,006 % by weight of bromine. One source (Söderström, G. et al, 2000) reports typical content of 0,004 % by weight from a study in 1992, but that the level has increased the last decade. In comparison is normal chlorine level in municipal waste approximately 0,75 % by weight (Söderström, G. et al, 2000).

The brominated waste added to the municipal waste was generated in a demolition plant for electric and electronic waste (Stena Miljø AS, Oslo). In total approximately 70 tons brominated waste were generated for the incineration tests.

Stena Miljø AS has calculated the level of bromine in the actual mixture (brominecontaining plastics) which were used in the tests at Klemetsrud and Energos Plants. The calculations gave the following levels : 27 % by weight brominated plastics, 16 % by weight wooden material, 57 % by weight plastics without bromine (Aass, F.E., 2003a). The bromine level in plastics is reported to be 3-4,5 % by weight, and the bromine level in the total mixture approximately 1 % by weight (Sjølin, S., 2003).

Approximately 80 % by weight of the brominated plastics is reported to contain PBDE (polybrominated diphenylethers).

Exact level of BFRs in the plastics is not known, but is earlier reported to be approximately 12 % by weight (SFT, 2003) (Aass, F.E., 2003b). This is determined mainly from PCs and monitors, and the level relatively uncertain.

A Danish report from 1999 reports the content of TBBPA and other BFRs separately in different electronic products (Miljøstyrelsen i Danmark, 1999). Reported levels are:

Colo	our TVs :	
a)	TBBPA	12 % by weight
b)	BFRs	12 % by weight
PCs		
a)	TBBPA	12-14 % by weight
b)	BFRs	12-14 % by weight

•

Levels in printers, photocopiers and fax-machines are reported to be lower.

As the brominated plastics used in the tests mainly origins from TVand monitor-cabinets, it is assumed a BFR-level in the plastics of 12 % by weight.

Stena Miljø AS has also reported that the waste mix may contain approx. 1 % by weight PVC (Aass, F.E., 2004), from which on may derive that the chlorine level in the waste is significantly lower than in municipal waste from households and businesses.

The Norwegian State Pollution Control Authority (SFT) has done a preliminary analysis of BFRs in 60-100 kgs of plastic waste from the Stena Miljø AS demolition plant.

The total concentration level of BFRs was determined to be approximately 20 000 mg/kg, i.e. 2 % by weight.

The level of BFRs in the these plastics is therefore lower than the assumed levels for the plastics used in the tests. Still, the samples analysed by SFT may not necessarily be representative for the brominated waste used in the tests. Further, SFT showed that octaBDE and decaBDE was the dominating BFR-compounds, with a level of 8 000 - 9 000 mg/kg for each of the compounds.

Figure 5 shows brominated waste used in the incineration tests (before shredding).



Figur 5 Brominated waste used in the incineration tests (before shredding).

5.3 Description of plants and sampling points

5.3.1 Klemetsrud Plant

Measurements are carried out at Oslo Municipality's incineration plant at Klemetsrud in Oslo. The plant incinerates untreated municipal waste from households and businesses in 2 lines, each with a capacity of approx. 10 tons/hr. Each line is equipped with a flue gas cleaning system, consisting of active coal injection, a fabric filter and a wet scrubber.

A sketch of the plant, with marking of the sampling points, is shown in figure 6.



Figure 6 Sketch of Klemetsrud Plant, with marking of the sampling points.

1 Bottom ash Total bromine, total chlorine. Brominated flame-retardants.

2 Raw flue gas Gaseous bromine and chlorine. Brominated, chlorinated, brominated/chlorinated dioxins

3 Filter dust Total bromine, total chlorine. Brominated flame-retardants.

4 Scrubber water Total bromine, total chlorine. Brominated flame-retardants.

5 Cleaned flue gas Brominated, chlorinated, brominated/chlorinated dioxins. Brominated flame-retardants.

Measurements without addition of bromine containing waste

Measurements of brominated, chlorinated, brominated/chlorinated dioxins were done on line 1 by TÜV in parallel to the annual emission control measurements October 16th -17th 2003. Measurements of total bromine and chlorine in raw flue gas were done on line 2 December 18th 2003 by Kjelforeningen-Norsk Energi.

Measurements with addition of bromine containing waste

The measurements with addition of bromine containing waste were done October 28th and 30th 2003 by Kjelforeningen Norsk Energi. Measurements were done with two different mixtures:

- Low addition: 5 % by weight addition of bromine containing waste. This mixture gives a feed rate for bromine containing waste of approx. 0,5 tons/hr, i.e. slightly above 5 % by weight. The resulting bromine feed rate was approx. 5 kg bromine/hr.
- High addition: 10 % by weight addition of bromine containing waste. It is possible to feed up to 2 tons/hr of bromine containing waste at line 2 at the Klemetsrud Plant. This is however an unrealistic high share, because it may significantly affect the incineration conditions. A realistic maximum addition is approx. 1 ton/hr (10 % by weight

bromine containing waste). This mixture gives a feed rate of approx. 10 kg bromine/hr.

In order to maintain stable concentrations in output flows from the process, feeding of bromine containing waste to the incinerator has to start in due time before sampling.

Figure 7 shows the calculated theoretical change in concentration of bromine in filter dust, and in circulating fluids in scrubber (HCl-step). One can see from the figure that the concentration level in filter dust during sampling period is approx. 80-90 % by weight of maximum concentration level, and that the actual time of feed start, 24 hours before sampling, was sufficient to maintain a stable concentration level in circulating fluids in scrubber.

Figure 7 Calculated change in concentrations of bromine in filter dust and circulating fluids in scrubber (HCI-step).



Concentration change in filter dust



Concentration change in circulating fluids in scrubber

5.3.2 Energos Plant

The Energos Plant at Ranheim incinerates annually approx. 10 000 tons of waste. The plant has approx. 4 MW thermal output, and a steamproduction of approx. 25 GWh/year, supplied to a neighbouring industrial plant, Peterson Linerboard Ranheim (PLR). The waste is a mixture of waste from PLR and other industries.

The flue gas is treated in a fabric filter after injection of coal and lime. Output flows are bottom ash/slag, fly ash and emissions to air from stack.

A flow sheet for the plant is shown in Figure 8. The waste is fed from the storage silo (1) with the conveyor (2) into the 2nd storage (3). From this storage the waste is fed in portions onto the fire grate in the primary chamber of the furnace (4). On the fire grate the waste is dried, gasified and burned-out at sub-stoichiometric conditions. A conveyor brings the waste through the primary chamber, and to the output shaft, where the burned waste falls down as slag. The flue gases are led through the boiler (convection unit) (5) and are cleaned in the fabric filter (8) after addition of activated carbon and lime.

Measurements with and without addition of 20 % by weight of bromine containing waste were carried out November $11^{\text{th}}-13^{\text{th}}$ 2003 by TÜV. The sampling points are positioned right into the inlet to the filter (8), and in the vertical outlet of the filter/inlet to stack (9).

Figure 8 Flow sheet for the Energos Plant at Ranheim



- 5. 2nd sto 5. Boiler
- Boiler
 Reactor
- 9. Chimney
- 3. Filter system

Steam system

6.

10. Control- / monitoring system

5.3.3 FREVAR Plant

FREVAR Incineration Plant is owned by Fredrikstad municipality. The plant incinerates approximately

78 000 tons waste annually, using two incineration furnaces. The plant produces 185 GWh steam per year, and has 99 % utilization of the produced energy (FREVAR, 2004).

The waste is fed into the feedershaft with a crane. From the shaft, the waste is fed in portions onto the fire grate. On the grate, the waste is dried, combusted and burned out. The movable grate takes the waste through the furnace to the outgoing shaft, into which the burned waste drops down as slag.

The flue gases are burned in a secondary combustion zone over the grate. The flue gas is cleaned in an electric precipitator, wet scrubber and a fabric filter. Active coal is added prior to the wet scrubber, and activated carbon and lime prior to the fabric filter.

The flue gases from the two furnaces are led in to the same duct before the scrubber, and let out trough a joint stack.

FREVAR also has a incinerator for hospital waste. The flue gas from this furnace is quenched and treated through a separate wet scrubber, before it is led in to one of the other furnaces for further combustion and cleaning. Annual control measurements at FREVAR are normally done with the hospital waste incinerator running.

Measurements of gaseous bromine and chlorine, and brominated, chlorinated and brominated/chlorinated dioxins were carried out parallel to the annual control measurements on November 5th 2003, by Kjelforeningen Norsk Energi.

A flow sheet of the plant is shown in Figure 9.

Figure 9 Sketch of the FREVAR incineration plant for household waste (FREVAR, 2004), with sampling points.



FREVAR FORBRENNINGSANLEGG - FA

6.Results

6.1 Operating conditions

The combustion conditions were normal during sampling and measurement at the Klemetsrud plant, with CO-levels of approximately 20-30 mg/Nm³. CO-levels during sampling at FREVAR Plant were approx. 50 mg/Nm³. At the Energos Plant, CO was not detected during measurements, which indicates a very effective combustion.

Some problems were experienced with the fabric filter at FREVAR Plant during the sampling period.

6.2 Measurements of gaseous bromine in flue gas before cleaning

Results of measurements of gaseous bromine in flue gas before cleaning are shown in Table 7.

Plant Addition of bromine containing waste		Gaseous bromine				
		Addition of bromine containing waste	HBr		Br ₂	
			mg/Nm ³	kg/hour	mg/Nm ³	kg/hour
Klemetsrud	No ad	dition	3,6	0,2	0,1	0,007
Plant	5 % b	y weight addition	6	0,4	0,3	0,02
	10 % by weight addition		40	2,5	2,3	0,1
FREVAR Plant	No addition		1,2	0,08	<0,5	<0,03
Energos	No ad	dition	< 2,15	< 0,014	< 2,15	< 0,014
Plant	20 %	by weight addition	97-200 ¹⁾	0,95-1,97	< 2,15	< 0,014

Table 7 Results from measurements in uncleaned flue gas with different addition of bromine containing waste

1) During the approx. 12 hour sampling period, the HBr-concentration in raw flue gas varied from approx. 97 to 200 mg/Nm³, with the highest level during the last sample.

The measurement results for gaseous bromine in uncleaned flue gas, with a normal waste composition, indicates a bromine level equal to, or slightly lower than what is normal for waste from households and small businesses (0,003-0,006 % bromine by weight).

Results from measurements during addition of bromine containing waste, shows a clear increase in the HBr-concentration in uncleaned flue gas, compared to results from measurements with no brominated waste.
At the Energos Plant, gaseous bromine was also measured after the filter. The concentration was $< 2,2 \text{ mg/Nm}^3$, which leads to a removal efficiency of >97 % for the filter.

6.3 Mass balance for bromine

Figure 10 shows a mass balance for bromine after addition of 10 % by weight bromine containing waste at Klemetsrud Plant.





From Figure 10, we can see that the mass balance of bromine from Klemetsrud Plant shows good correspondence between input and output flows in the plant.

At the Energos Plant, 20 % by weight of bromine containing waste was fed into the furnace.

Table 8 shows resulting bromine levels in input and output mass flows.

Mass flow	Amount (kg/hour)
Bromine in input waste (total input)	2,1
Bottom ash	0,045
Flue gas before filter	0,95-1,97
Flue gas after filter	0,014
Total output (excl. bromine in filterdust)	1,0-2,0

Table 8 Bromine in input and output mass flows, Energos Plant

During the approximately 12 hours of sampling and measurement, the bromine content in flue gas before filter varied between approximately 1-2 kgs/hr, with the highest level during the last measurement. According to Energos, it is very likely that the adsorption-/desorption-processes in the boiler system leads to a slow increase of HBr-level in flue gas. Bromine in filterdust is not measured.

6.4 Brominated, chlorinated and brominated/chlorinated dioxins

Concentrations in emissions to air (flue gas after cleaning)

Figure 11 shows resulting emissions of brominated, chlorinated and brominated/chlorinated dioxins with no addition of brominated waste, and with the addition of 5 %, 10 % and 20 % by weight of brominated waste respectively. The results are reported as the actual concentration levels, not as toxic equivalents.



Figure 11 Total emissions of the brominated, chlorinated and brominated/chlorinated dioxins analysed.

From Figure 11 one can see that the emissions of *chlorinated dioxins* (PCDDs+PCDFs, as actual concentrations, not toxic equivalents) from Klemetsrud Plant were approx. 1,5 ng/Nm³ with no addition of brominated waste, and approx. 0,5 ng/Nm³ with 5 % and 10 % by weight of bromine containing waste. At the FREVAR Plant, the emissions of chlorinated dioxins were approx. 1 ng/Nm³ (with no addition of bromine containing waste). The Energos Plant had an emission concentration of approx. 0,3 ng/Nm³ with the addition of 20 % bromine containing waste.

Further, one can see that the emission of *brominated dioxins* (PBDDs+PBDFs) was very low, both with and with no addition of bromine containing waste.

The emissions of *brominated/chlorinated dioxins* (ClxBryDDs+Clx BryDFs) with no addition of brominated waste (Klemetsrud and FRE-

VAR plants) were apparently higher than with addition of brominated waste (Klemetsrud and Energos plants).

Resulting concentrations of brominated/chlorinated dioxins (ClxBryDDs+ClxBryDFs) during measurements with addition of brominated waste, were less than half of the concentrations of chlorinated dioxins (PCDDs+PCDFs).

The emissions of *chlorinated dioxins (PCDDs/Fs)*, presented as nordic toxic equivalents, resulting from addition of brominated waste, is shown in Figure 12.





EU threshold value

From Figure 12, one can see that the emissions of chlorinated dioxins, presented as Nordic Toxic Equivalents, were approx. $0,03 \text{ ng/Nm}^3$ from the Klemetsrud Plant, both with addition of 5 % and 10 % by weight of bromine containing waste. From the Energos Plant, the concentration level was $0,006 \text{ ng/Nm}^3$ with addition of 20 % by weight of bromine containing waste.

The corresponding emission limit value in the EU-directive for waste incineration is $0,1 \text{ ng/Nm}^3$.

Uncertainty in sampling and analysis, variations in operating conditions and waste mixture, differences between laboratories with respect to methods of analysis (especially dioxins), makes comparison of results difficult. We may however draw the following main conclusions:

- Increasing the content of BFRs in the waste gave no significant increase in the emissions of chlorinated dioxins, or either brominated and chlorinated/brominated dioxins
- The emission level is highest for chlorinated dioxins, lower for chlorinated/brominated dioxins and lowest for brominated dioxins
- The emission levels for chlorinated dioxins, reported as Nordic toxic equivalents, are low compared to the emission threshold value in the

EU-directive for incineration of waste. The reported emission levels were 0,03 ng/Nm³ and 0,006 ng/Nm³ respectively for the Klemetsrud Plant (Oslo) and Energos Plant (Ranheim), and the EU threshold value is 0,1 ng/Nm³.

The emission measurement results indicate that the incineration efficiency and the operating conditions of the flue gas treatment systems are of greater importance to the resulting emission levels for dioxins, rather than the bromine content level.

Flue gas concentrations before and after cleaning

Measurements of dioxins in flue gas before and after cleaning were carried out with addition of a high proportion (10 % by weight) bromine containing waste, see Figure 13.

Figure 13 Concentrations of brominated, chlorinated and brominated/chlorinated dioxins analysed in raw flue gas/emission outlet from Klemetsrud Plant, resulting form addition of 10 % by weight of bromine containing waste.



Concentration of brominated/chlorinated dioxins before cleaning is significantly higher than the corresponding concentration of chlorinated dioxins. After cleaning, the brominated/chlorinated dioxins amounts to only 10-20 % of the total emissions of dioxin compounds. Dioxins removed from flue gas are found in the filter dust. The filter dust from incineration plants is treated as hazardous waste.

6.5 Brominated flame-retardants (BFRs)

Table 9 shows concentrations of BFRs in bottom ash, filter dust, water from the scrubber and in emissions to air from tests with addition of bromine containing plastics at Klemetsrud and Energos Plants.

		Level	of BFRs
	Unit	Klemetsrud Plant	Energos Plant
Bottom ash Filter dust	mg/kg mg/kg	0,034-0,1 0,04	<0,016
Scrubber water (untreated)	ng/l	0,01	-
cleaning)	ng/Nm ³	14-22	< 0

Table 9 Concentrations of BFRs from incineration tests with addition of bromine containing plastics at Klemetsrud and Energos Plants.

From table 9, one can see that the concentration of BFRs in flue gas from Klemetsrud Plant was 14-22 ng/Nm³. This equals 0,9-1,4 mg/hour and approximately 0,01 kg/year, assuming 8000 running hours/year at the same emission level. A Danish study (Miljøstyrelsen, 1999) estimates the total annual Danish emissions of BFRs from incineration to be < 0,04 tons. A report from the Norwegian National State Pollution Control Authority (SFT), estimates the national emissions from combustion in Norway to be < 0,01 tons/year (1998), i.e. < 10 kg/year.

At the Energos Plant (Ranheim) the reported concentration of BFRs in the flue gas was $<5 \text{ ng/Nm}^3$.

The concentration of BFRs in bottom ash from the tests at Klemetsrud Plant (Oslo) shows levels far below the emission limit value of 0,25 % by weight stated in the Hazardous Waste Directive.

DekaBDE and TBBPA (Tetrabrombisphenol A) are the dominating compounds of BFRs in the bottom ash from Klemetsrud Plant. In water from the scrubber and in the flue gas, dekaBEDE has the highest concentration level.

The amount of BFRs in the waste mixture used in the tests at Klemetsrud Plant was not analysed, but calculated/estimated to be approximately 30 kg/hr, based on a share of bromine containing plastics of 27 % by weight, and an assumed content of BFRs in the plastics of 12 % by weight.

Figure 14 shows input and output flows of brominated flameretardants at Klemetsrud Plant (Oslo) with 10 % by weight addition of brominated waste.



Figure 14 Input and output flows of brominated flame-retardants at Klemetsrud Plant (Oslo) with 10 % by weight addition of brominated waste.

The results indicates that the BFR-level in output flows amounts to less than 0,001 % by weight of the total BFRs in the waste mixture



Job Date

From

TNG Energy from Waste Facility, Eastern Creek, References Facilities 2016-10-26 Ahmet Erol, Martin Brunner

Background

TNG has been requested to provide further information on the capacity of the technology to process/handle the proposed volume (approx. 50%) of C&D waste. Ramboll has been asked to identify such plants and supply information on the operational experience of such plants.

Results

We acknowledge that it has not been possible to identify an EfW plant (neither with comparable nor with alternative technology) processing a documented input of 50% C&D waste. The main reason is the fact, that any EfW plant treating primarily preprocessed waste (as this is the case for TNG) receives these preprocessed waste streams from different sorting/pre-processing plants. Once waste has been pre-processed it "looses" its waste declaration/identification and cannot be tracked back to its origin. Therefore it is not possible neither to declare the initial origin of the waste nor the exact composition concerning C&D, C&I, etc.

Nevertheless when taking in to account the relevant aspects for the design of an EfW plant (mainly the physical and chemical waste composition) it is possible to demonstrate that TNG operates well within the range of comparable facilities, namely the listed reference plants.

Reference facilities

The information on reference facilities provided in February 2016 (attached) provides details on the capacity, technology type and fuel mix including the chemical analysis of the design fuel of these plants.

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The reference plants are treating big variation in the feedstock, which shows that the technology can be used on a broader level. There variation in the feedstock profile of all of the reference facilities combined is evidence to demonstrate a technological capacity to withstand a wide range of variance. The provided technology with moving grate technology and semi dry flue gas treatment is able to run with the waste composition of TNG.

Following several key design parameters are listed and discussed in relation to the design parameters of TNG.

Plant capacity

The mechanical throughput of TNG is comparable with the plant in Grossräschen (DE) and Ferrybridge (UK). While the plant capacity of TNG seems higher than these plants the thermal capacity (throughput x CV) – the most important design parameter – is identical. TNG therefore is in no way an exceptionally large plant

Calorific value

The calorific value defines the combustion characteristics of the waste. Generally it can be said that - except for very low CV below 8 MJ/kg - the higher the CV, the more difficult to maintain an ideal combustion process. With a CV of 12.3 MJ/kg TNG falls in the medium range between i.e. Knapsack with 15 MJ/kg or Riverside with 9.6 MJ/kg.

Chemical waste composition

Within the waste composition the most important parameters are:

- Moisture (limits the controlled ignition of the waste)
- Inert (ash) content (limits homogenous combustion and burnout)
- The larger of Chlorine or Sulphur content (is the limiting factor for the APC system)
- C/O ratio (high C/O ratio is an indicator for high plastic content which limits homogenous combustion and burnout)

For all these aspects TNG is well within the range of all the reference plants.

Summary

None of the listed reference facilities is an exact replica of the TNG fuel profile, however all relevant design parameters of TNG are well within comparable plants which are successfully in operation. As a result it can be said that the technology option pursued, being moving grate technology with semi dry flue gas treatment, was selected based on its capacity to handle a wide range of fuel types and variation of feed stock.



Appendix A Reference Facilities

Key Plant Parameters

Facility/Location	Country	Commission year	Capacity	Fuel mix	Furnace/Boiler	Furnace/Boiler Supplier Furnace/Boiler		Supplier APC
TNG	AU	-	4 x 276'250	C&I, C&D	Grate	HZI	Semi dry (lime)	-
Grossräschen	DE	2008	1 x 246'000	C&I, C&D	Grate	AEE*	Semi dry (lime)	LAB
Heringen	DE	2009	2 x 148'500	C&I, C&D, some MSW	Grate	AEE*	Semi dry (lime)	LAB
Premnitz	DE	2008	1 x 150'000	C&I, C&D	Grate	AEE*	Semi dry (lime)	Lühr
Hannover	DE	2005	2 x 140'000	C&I, C&D, some MSW	Grate	AEE*	Semi dry (lime)	LAB
Knapsack	DE	2009	2 x 150'000	C&I, C&D	Grate	AEE*	Semi dry (lime)	Lühr
Ferrybridge	UK	2015	2 x 256'500	C&I, C&D, some MSW, waste wood	Grate	HZI	Semi dry (lime)	HZI
Riverside	UK	2011	3 x 195'000	MSW, C&I	Grate	HZI	Semi dry (lime)	HZI

* up to 2010 HZI was part of the AEE Group



Appendix B Reference Facilities

Chemical Analysis Design Fuel

		TNG	Grossräschen	Heringen	Premnitz	Hannover	Knapsack	Ferrybridge	Riverside
Carbon (C)	%	31.44	35.20	n.a.	28.50	n.a.	n.a.	35.60	26.63
Hydrogen (H)	%	4.07	1.88	n.a.	3.96	n.a.	n.a.	5.20	3.78
Nitrogen (N)	%	0.26	3.80	n.a.	0.32	n.a.	n.a.	0.60	0.54
Sulphur(S)	%	0.43	0.37	n.a.	0.18	n.a.	< 0.8	0.20	0.10
Chloride (Cl)	%	0.88	0.70	n.a.	0.54	n.a.	1.20	0.50	0.70
Oxygen (O)	%	18.06	14.25	n.a.	19.50	n.a.	n.a.	25.10	17.79
Water (H2O)	%	23.38	25.00	n.a.	22.00	n.a.	18.00	20.00	30.76
Ash	%	21.49	18.80	n.a.	25.00	n.a.	19.00	12.80	19.70
Total	%	100.00	100.00	-	100.00	-	-	100.00	100.00
NCV	MJ/kg	12.30	12.50	12.6	13.00	13.5	15.00	13.50	9.60

Percentage of wood (estimation based on chemical analysis of waste)

		TNG	Grossräschen	Heringen	Premnitz	Hannover	Knapsack	Ferrybridge	Riverside
Wood	%	30.24	23.86	n.a.	32.65	n.a.	n.a.	42.03	29.79

Chloride range of fuel (average)

		TNG	Grossräschen	Heringen	Premnitz	Hannover	Knapsack	Ferrybridge	Riverside
CI	%	< 1	<1	n.a.	0.2-1.5	n.a.	0.5-1.7	<1	n.a.

information source for reference plants: HZI



Appendix C Reference Facilities

Fuel Mix

		Mixed C&D						Mixed C&I						SRF from MSW	MSW	
		paper/card	plastic	textile	glass	vegetation	poom	hazardous (hospital waste)		paper/card	plastic	textile	flock waste			
TNG	28.7%	х	х	х	х	х	х	-	71.3%	х	х	х	х	0.0%	-	-
Grossräschen *	9.8%	х	х	-	х	х	х	х	83.2%	х	х	х	х	7.0%	х	х
Heringen *	13.6%	х	х	х	-	х	х	х	62.4%	х	х	х	-	24.0%	х	-
Premnitz *	14.3%	х	х	х	-	х	х	х	57.0%	х	х	х	х	28.7%	х	-
Hannover *	9.0%	×	х	х	-	х	х	×	75.3%	х	х	×	х	15.7%	х	×
Knapsack	10.0%	х	х	х	-	х	х	х	90.0%	х	х	х	-	0.0%	х	-
Ferrybridge ¹⁾	10.0% ²⁾	n.a.	n.a.	n.a.	n.a.	n.a	х	n.a.	30.0%	n.a.	n.a.	n.a.	n.a.	60.0%	х	х
Riverside	n.a.	х	х	х	х	х	х	-	n.a.	х	х	х	-	n.a.	х	-

¹⁾ Design Waste

²⁾ Waste Wood

* values from the year 2014, see https://www.itad.de/information/abfallverwertungsanlagen

all other values given from the operators verbally

SRF Solid Recovered Fuel

MSW Municipal Solid Waste

n.a. not available





Job

Date From TNG Energy from Waste Facility, Eastern Creek, 850°C minimum operating temperature 2016-09-27 Ahmet Erol

850°C minimum operating temperature

The design of the proposed Energy from Waste (EfW) facility includes a secondary combustion chamber to optimise flow conditions and temperature profile and reduce CO, VOC and other organic pollutants emissions.

A minimum flue gas temperature of 850°C with a residence time of at least 2 seconds is kept at any time after the last injection of air (in this case the secondary air). The energy contained in the waste, the design of the secondary combustion chamber and the combustion control enable the plant to run at these conditions without any additional fuel or energy input.

Running an EfW facility at a temperature of 1100°C and a residence time of at least 2 seconds after secondary air injection is not possible with only the energy of the waste. The auxiliary burners (fuelled by gas or diesel) have to support the combustion to reach this temperature and residence time. This additional (fossil) energy input reduces the overall energy efficiency of the facility.

The requirement of 1100°C given by the IED is in Europe only applies for hazardous waste treatment facilities where rather small quantities of hazardous/chemical waste are treated and where (due to the limited size of the plant) energy efficiency is not a key issue.

Based on the waste analysis and the mixing described in the Project Definition Brief (chapter 2.3.1) a maximum chlorine content of the mixed waste of 1% can be assumed. Ramboll Hannemanns Allé 53 DK-2300 Copenhagen S Denmark

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Job Date

From

TNG Energy from Waste Facility, Eastern Creek, Treated Wood Waste 2016-08-30 Martin Brunner, Ahmet Erol

Treated wood waste

Treated wood waste (TWW) represents a large proportion of waste wood arising. A WRAP study on waste composition found that (including laminated and veneered wood) an average of 85% of the wood from the observed Civic Amenity sites was treated and 23% of the wood from the observed construction and demolition sites was treated.

TWW is defined as wood that has been treated with one or more of the following:

- Copper Chromium Arsenic (CCA)
- Copper Organics
- Creosote
- Light Organic Solvent Preservatives (LOSP)
- Micro-emulsion
- Paint / stain
- Varnish

The EfW **plant must burn waste** aligning with the relevant requirements of Australian and NSW Regulatory Framework.

According to NSW Energy from Waste Policy Statement, chapter 4 Energy recovery facilities, technical criteria the gas resulting from the process should be raised, after the last injection of combustion air to a minimum temperature of **850** °**C for two** seconds. If a waste has a content of more than 1% of halogenated organic substances, expressed as chlorine, the temperature should be raised to 1,100°C for at least 2 seconds after the last injection of air.

Certain wood wastes are treated with wood preservatives or coatings like listed above.

One of the main sources for organic chlorine is varnish containing Polychlorinated biphenyl (PCB).

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To get an idea of how much chlorine in treated wood can be, a simple calculation with following assumptions made:

Wood size:	0.0254 m x 0.0254 m x 1 m (1 inch x 1 inch)
Varnish thickness on wood:	60 μm
Specific weight wood	700 kg/m ³
Specific weight varnish	900 kg/m ³
Calculated varnish weight on treated wood	5.48 g
Calculated wood weight	452 g
Percentage of varnish on wood	1.2%

The chlorine content of PCB varies from 20% to 70% depending on PCB configuration¹.

With the assumptions and calculation above the possible chlorine content in treated wood for TNG is as follows.

The wood content in TNG design fuel is varying from 0 to 58.20% with an average of 30.24%. Assumption: 100% CRW incineration with 58.20% of waste wood and varnish contents 100% chlorine. The chlorine content for this unrealistic case of the incinerated waste would be about 0.71%.

More realistic is to calculate with the average waste wood content of 30.24% and chlorine content in varnish of 50%. For this case the chlorine content in incinerated waste would be about 0.19%.

<u>Conclusion</u>

Even if TNG would incinerate 100% CRW fraction with the highest waste wood content, it is unrealistic to reach the chlorine content of 1%. There is no reason at any time to raise the incineration temperature to 1,100°C for at least two seconds.

<u>Source</u>

Options and Risk Assessment for Treated Wood Waste, The Waste & Resources Action Programme

http://www2.wrap.org.uk/downloads/Options_and_Risk_Assessment_for_Treated_Wood_W aste.6ac4f667.2237.pdf

¹ Polychlorinated biphenyl (PCB) is an organic chlorine compound with the formula $C_{12}H_{10-x}Cl_{x}$. There are 209 configurations with 1 to 10 chlorine atoms.

https://en.wikipedia.org/wiki/Polychlorinated_biphenyl



BAT Evaluation Job Client DADI TNG NSW Memo no. 4 18/02/2016 Date To whom it may concern То From Martin Brunner Ian Malouf (DADI) Copy to Phill Andrew (Savills) Rachael Snape (Urbis) Geert Stryg (Ramboll)

1. Methodology to compare the TNG technology with the BAT requirements

Based on the long term experience of Waste-to-Energy in Europe the best available techniques (BAT) have been evaluated, defined and documented in the "Reference Document on the Best Available Techniques for Waste Incineration (August 2006)" (in short BREF).

Following a request by TNG Ramboll has compared the requirements as summarized as 68 basic requirements in Chapter 5.1 and 5.2 of the above document. Date 11.02.2016

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2. Results and conclusions

The detailed evaluation of each point is found in the attachment. In summary the results of the evaluation are:

- 1. Design: all requirements defined by BREF are fulfilled
- 2. Emissions: The expected emissions are within the required operational values given by BREF
- 3. Energy efficiency: The TNG facility exceeds the requirements of the BREF

It can therefore be concluded that the TNG facility fulfils the BREF requirements and therefore is BAT.



BAT	TNG	BAT fullfilled?
1. the selection of an installation design that is suited to the characteristics of the waste received, as described in 4.1.1 and 4.2.1 and 4.2.3	The thermal treatment is calculated for a calorific value of 8,5 MJ/kg- 16,5 MJ/kg. The plant is designed for a waste throughput of 25,35 t/h to 46,48 t/h. The waste type is mainly Chute Residual Waste (CRW) from the Genesis Facility, Commercial and Industrial (C&I), Construction and Demolition (C&D) and flock waste. A combination of water cooled grate and air cooled grate was chosen. The air cooled grate is characterized through: - suitable for waste with low to medium heat values - suitable for municipal and other heterogeneous solid wastes - can cope with sewage sludge and/or medical waste mixed with municipal waste - applied at most modern MSW installations The water cooled grate protects the grate against intense heat (LCV: 10 – 20 GJ/t) otherwise it has the same characteristics.	Yes
2. the maintenance of the site in a generally tidy and clean state, as described in 4.1.2	It is integral part of the contract; it is defined as one the operator's tasks.	Yes
3. to maintain all equipment in good working order, and to carry out maintenance inspections and preventative maintenance in order to achieve this	The Proposed Facility will be operated and maintained by a dedicated Operations and Maintenance team. For planning the major shutdown a short shutdown is carried out approx. 1 year in advance. A normal duration for such a short shutdown is generally a few days. The operator installs a computer-controlled program used to ensure ongoing maintenance of the plant components. The program records conducted maintenance jobs and systematic maintenance can be planned for the individual components. A maintenance plan will be established to determine weekly, monthly, annual or longer interval inspections, tests and maintenance activities which have to be performed.	Yes



ВАТ	TNG	BAT fullfilled?
 4. to establish and maintain quality controls over the waste input, according to the types of waste that may be received at the installation, as described in: 4.1.3.1 Establishing installation input limitations and identifying key risks, and 4.1.3.2 Communication with waste suppliers to improve incoming waste quality control, and 4.1.3.3 Controlling waste feed quality on the incinerator site, and 4.1.3.4 Checking, sampling and testing incoming wastes, and 4.1.3.5 Detectors for radioactive materials. 	A three point checking procedure will be on-site to ensure only permitted waste is unloaded into the waste bunker. Initial inspections of the vehicle loads are undertaken at the weighbridge to check the content of the incoming and its origin. All loads are recorded and monitored by on-site CCTV when entering the tipping hall and while the loads are tipped off. Suspect loads are either rejected from the facility or are analysed in a designated inspection area or on the tipping hall floor prior to being tipped into the waste bunker. If the analysis shows that the load contains unsuitable material the waste is reloaded to the vehicle and rejected. Detectors for radioactive material are not necessary.	Yes
such that the risk of potentially polluting released is minimised. In general it is BAT to store waste in areas that have sealed and resistant surfaces, with controlled and separated drainage as described in 4.1.4.1.	thermal value of the waste there is no need for controlled and separated drainage.	Yes
 6. to use techniques and procedures to restrict and manage waste storage times, as described in 4.1.4.2, in order to generally reduce the risk of releases from storage of waste/container deterioration, and of processing difficulties that may arise. In general it is BAT to: prevent the volumes of wastes stored from becoming too large for the storage provided in so far as is practicable, control and manage deliveries by communication with waste suppliers, etc. 	 The waste bunker has the capacity to store the amount of 5-7 days of waste delivery. The waste is continuously removed. It is ensured that no excessive amounts will arrive. 	Yes
 7. to minimise the release of odour (and other potential fugitive releases) from bulk waste storage areas (including tanks and bunkers, but excluding small volume wastes stored in containers) and waste pre-treatment areas by passing the extracted atmosphere to the incinerator for combustion (see 4.1.4.4). In addition it is also considered to be BAT to make provision for the control of odour (and other potential fugitive releases) when the incinerator is not available (e.g. during maintenance) by: a. avoiding waste storage overload, and/or b. extracting the relevant atmosphere via an alternative odour control system 	The incineration air supply (primary or secondary) is taken from the waste storage areas. By enclosing the waste storage areas and limiting the size of the entrances to the waste storage areas, a slight under pressure can be ensured on whole waste storage area.	
8. the segregation of the storage of wastes according to a risk assessment of their chemical and physical characteristics to allow safe storage and processing, as described in 4.1.4.5	The waste is delivered in defined fractions. Segregation of the storage of wastes is not necessary.	Yes



BAT	TNG	BAT fullfilled?
9. the clear labelling of wastes that are stored in containers such that they may continually be identified, as described in 4.1.4.6.	The waste is not stored in containers.	Not relevant
 10. the development of a plan for the prevention, detection and control (described in 4.1.4.7) of fire hazards at the installation, in particular for: waste storage and pre-treatment areas furnace loading areas electrical control systems bag house filters and static bed filters. It is generally BAT for the plan implemented to include the use of: a. automatic fire detection and warning systems, and b. the use of either a manual or automatic fire intervention and control system as required according to the risk assessment carried out. 	Fire detection and firefighting systems are installed at the critical areas of the plant (e.g. waste bunker, electrical rooms, feed hopper,). In case of fire detection the firefighting systems start automatically.	Yes
11. the mixing (e.g. using bunker crane mixing) or further pre-treatment (e.g. the blending of some liquid and pasty wastes, or the shredding of some solid wastes) of heterogeneous wastes to the degree required to meet the design specifications of the receiving installation (4.1.5.1). When considering the degree of use of mixing/pre-treatment it is of particular importance to consider the cross-media effects (e.g. energy consumption, noise, odour or other releases) of the more extensive pre-treatment's (e.g. shredding). Pre-treatment is most likely to be a requirement where the installation has been designed for a narrow specification, homogeneous waste.	The waste bunker has sufficient space and sufficient capacity for the mixing of different waste streams. The waste is mainly delivered as shredded fraction.	Yes
12. the use of the techniques described in 4.1.5.5 or 4.6.4 to, as far as practicably and economically viable, remove ferrous and non-ferrous recyclable metals for their recovery either: a. after incineration from the bottom ash residues, or b. where the waste is shredded (e.g. when used for certain combustion systems) from the shredded wastes before the incineration stage.	The recyclable material is removed before the waste is delivered to the waste incineration plant.	Yes
13. the provision of operators with a means to visually monitor, directly or using television screens or similar, waste storage and loading areas, as described in 4.1.6.1	A video system is installed to monitor waste reception, feeding and storage, furnace as well as main out streams and their loading areas.	Yes
14. the minimisation of the uncontrolled ingress of air into the combustion chamber via waste loading or other routes, as described in 4.1.6.4	To minimize the uncontrolled ingress of air into the combustion chamber, the feed hopper always has to be filled to a certain level during operation. The level of the feed hopper is checked by the DCS.	Yes



BAT	TNG	BAT fullfilled?
 15. the use of flow modelling which may assist in providing information for new plants or existing plants where concerns exist regarding the combustion or FGT performance (such as described in 4.2.2), and to provide information in order to: a. optimise furnace and boiler geometry so as to improve combustion performance, and b. optimise combustion air injection so as to improve combustion performance, and c. where SNCR or SCR is used, to optimise reagent injection points so as to improve the efficiency of NOx abatement whilst minimising the generation of nitrous oxide, ammonia and the consumption of reagent (see general sections on SCR and SNCR at 4.4.4.1 and 4.4.4.2). 	The incineration and the boiler layout are based on good experiences and are realised on several operating plants. The SNCR (injection of ammonia), the secondary combustion chamber and the FGT design is based on a fluid dynamics flow model.	Yes
16. in order to reduce overall emissions, to adopt operational regimes and implement procedures (e.g. continuous rather than batch operation, preventative maintenance systems) in order to minimise as far as practicable planned and unplanned shutdown and start-up operations, as described in 4.2.5	The waste feeding is continuously. Implementation of a process orientated automatic start up and shut down procedure to minimise the emissions in those operational cases. Mixture of the waste to get a homogeneous fuel and preventative maintenance system avoid unplanned shutdowns.	Yes
17. the identification of a combustion control philosophy, and the use of key combustion criteria and a combustion control system to monitor and maintain these criteria within appropriate boundary conditions, in order to maintain effective combustion performance, as described in 4.2.6. Techniques to consider for combustion control may include the use of infrared cameras (see 4.2.7), or others such as ultra-sound measurement or differential temperature control	The following information is part of the combustion control system: - grate temperatures at various positions - caloric value of the waste - thickness of waste layer on the grate (visual control) - furnace and flue gas temperature at various positions - CO-, O ₂ -, CO ₂ - and H ₂ O-measurements at various positions - steam production data (e.g. temperature, pressure) - openings in the combustion wall for visual observation by cameras - length and position of the fire in the furnace - emission data for combustion related substances	Yes



ВАТ	TNG	BAT fullfilled?
 18. the optimisation and control of combustion conditions by a combination of: a. the control of air (oxygen) supply, distribution and temperature, including gas and oxidant mixing b. the control of combustion temperature level and distribution, and c. the control of raw gas residence time. Appropriate techniques for securing these objectives are described in: 4.2.8 Optimisation of air supply stoichiometry 4.2.9 Primary air supply optimisation and distribution 4.2.11 Secondary air injection, optimisation and distribution 4.2.19 Optimisation of time, temperature, turbulence of gases in the combustion zone, and oxygen concentrations 4.2.4 Design to increase turbulence in the secondary combustion chamber 	For an optimal combustion the thermal processes are monitored and regulated by measurements of the furnance temperature and the content of CO and O2 in the flue gas. For a more detailed description see point 17. by means of combustion control the following elements are controlled: - Primary air quantity and distribution - Oxygen content of flue gas (stoichiometry) - Secondary air quantity and distribution - Injection of recirculating flue gas in order to increase turbulence in the	Yes
19. in general it is BAT to use those operating conditions (i.e. combustion temperatures residence times and turbulence) that are specified in Article 6 of Directive 2000/76. The use of operating conditions in excess of those that are required for efficient destruction of the waste should generally be avoided. The use of other operating conditions may also be BAT – if they provide for a similar or better level of overall environmental performance. For example, where the use of operational temperatures of below the 1100 °C (as specified for certain hazardous waste in 2000/76/EC) have been demonstrated to provide for a similar or better level of overall environmental performance, the use of such lower temperatures is considered to be BAT.	The temperature / time requirements of the post-combustion chamber are continuously monitored. See point 17.	Yes
20. the preheating of primary combustion air for low calorific value wastes, by using heat recovered within the installation, in conditions where this may lead to improved combustion performance (e.g. where low LCV/high moisture wastes are burned) as described in 4.2.10. In general this technique is not applicable to hazardous waste incinerators.	For high energy efficiency and optimal combustion conditions preheating of primary and secondary combustion air is realised by using low pressure steam and saturated steam from the boiler drum.	Yes
21. the use of auxiliary burner(s) for start-up and shut-down and for maintaining the required operational combustion temperatures (according to the waste concerned) at all times when unburned waste is in the combustion chamber, as described in 4.2.20	Only in the case that the temperature in the secondary combustion chamber drops below a minimum temperature of 850 °C oil or gas fired support burners automatically start operation. Otherwise, the burners remain in a standby position.	Yes



BAT	TNG	BAT fullfilled?
 22. the use of a combination of heat removal close to the furnace (e.g. the use of water walls in grate furnaces and/or secondary combustion chambers) and furnace insulation (e.g. refractory areas or other lined furnace walls) that, according to the NCV and corrosiveness of the waste incinerated, provides for: a. adequate heat retention in the furnace (low NCV wastes require higher retention of heat in the furnace) b. additional heat to be transferred for energy recovery (higher NCV wastes may allow/require heat removal from earlier furnace stages) The conditions under which the various techniques may be applicable are described in 4.2.22 and 4.3.12 	To protect the walls of the boiler against corrosion Inconel will be cladded in sections where the flue gas temperature exceeds 850 °C.	Yes
23. the use of furnace (including secondary combustion chambers etc.) dimensions that are large enough to provide for an effective combination of gas residence time and temperature such that combustion reactions may approach completion and result in low and stable CO and VOC emissions, as described in 4.2.23	The furnace dimensions are large enough to ensure that for 2 seconds residence time the flue gas has a temperature above 850 °C (for all operational conditions). Experiences of several plants have shown low values for CO and VOC. After the combustion chamber two empty passes are installed to enable a complete burnout. In addition to secondary air a part of the flue gas is recirculated and injected together with the secondary air to achieve a maximum turbulence and burnout as well as stable CO and VOC emissions.	Yes
 24. When gasification or pyrolysis is used, in order to avoid the generation of waste, it is BAT to: a. combine the gasification or pyrolysis stage with a subsequent combustion stage with energy recovery and flue-gas treatment that provides for operational emission levels to air within the BAT associated emission ranges specified in this BAT chapter, and/ or b. recover or supply for use of the substances (solid, liquid or gaseous) that are not combusted 	No gasification or pyrolysis.	Not relevant



BAT	TNG	BAT fullfilled?
25. in order to avoid operational problems that may be caused by higher temperature sticky fly ashes, to use a boiler design that allows gas temperatures to reduce sufficiently before the convective heat exchange bundles (e.g. the provision of sufficient empty passes within the furnace/boiler and/or water walls or other techniques that aid cooling), as described in 4.2.23 and 4.3.11. The actual temperature above which fouling is significant is waste type and boiler steam parameter dependent. In general for MSW it is usually 600 – 750 °C, lower for HW and higher for SS. Radiative heat exchangers, such as platten type super heaters, may be used at higher flue-gas temperatures than other designs (see 4.3.14).	At the first position in the horizontal pass a protection evaporator is installed. The calculated gas temperature before the protection evaporator is around 650°C. This evaporator protects the superheater in order to minimize fouling. Further a conservative spacing of the boiler tubes prevents clogging and resulting operational problems.	Yes
 26. the overall optimisation of installation energy efficiency and energy recovery, taking into account the techno-economic feasibility (with particular reference to the high corrosivity of the flue-gases that results from the incineration of many wastes e.g. chlorinated wastes), and the availability of users for the energy so recovered, as described in 4.3.1, and in general: a. to reduce energy losses with flue-gases, using a combination of the techniques described in 4.3.2 and 4.3.5 b. the use of a boiler to transfer the flue-gas energy for the production of electricity and/or supply of steam/heat with a thermal conversion efficiency of: i. for mixed municipal waste at least 80 % (ref. Table 3.46) ii. for pretreated municipal wastes (or similar waste) treated in fluidized bed furnaces, 80 to 90 % iii. for hazardous wastes giving rise to increased boiler corrosion risks (typically from chlorine/sulphur content), above 60 to 70 % iv. for other wastes conversion efficiency should generally be increased in the range 60 to 90 % c. for gasification and pyrolysis processes that are combined with a subsequent combustion stage, the use of a boiler with a thermal conversion efficiency of at least 80 %, or the use of a gas engine or other electrical generation technology 	The most important measures to reduce the energy loss are: a) reduce excess air b) recirculate flue gas c) reduce the flue gas temperature at the boiler exit of 145°C d) heat recovery by condensate preheating The thermal conversion of the boiler is 91,9%.	Yes
27. to secure where practicable, long-term base-load heat/steam supply contracts to large heat/steam users (see 4.3.1) so that a more regular demand for the recovered energy exists and therefore a larger proportion of the energy value of the incinerated waste may be used.	The average net electrical efficiency is 29,6%. Necessary measures have been foreseen for later export of heat.	Yes



BAT	TNG	BAT fullfilled?
 28. the location of new installations so that the use of the heat and/or steam generated in the boiler can be maximised through any combination of: a. electricity generation with heat or steam supply for use (i.e. use CHP) b. the supply of heat or steam for use in district heating distribution networks c. the supply of process steam for various, mainly industrial, uses (see examples in 4.3.18) d. the supply of heat or steam for use as the driving force for cooling/air conditioning systems Selection of a location for a new installation is a complex process involving many local factors (e.g. waste transport, availability of energy users, etc) which are addressed by IPPC Directive Article 9(4). The generation of electricity only may provide the most energy efficient option for the recovery of the energy from the waste in specific cases where local factors prevent heat/steam recovery. 	See point 27.	Yes
 29. in cases where electricity is generated, the optimisation of steam parameters (subject to user requirements for any heat and steam produced), including consideration of (see 4.3.8): a. the use of higher steam parameters to increase electrical generation, and b. the protection of boiler materials using suitably resistant materials (e.g. claddings or special boiler tube materials) The optimal parameters for an individual installation are highly dependent upon the corrosivity of the flue-gases and hence upon the waste composition. 	Depending on the flue gas composition and the waste conditions the steam parameters were determined to be 73barA/430°C. This superheated steam parameters ensure high energy efficiency. The net electrical efficiency is 29,6%.	Yes
30. the selection of a turbine suited to:a. the electricity and heat supply regime, as described in 4.3.7b. high electrical efficiency	A condensing turbine was chosen as there are no possibilities to supply heat to customers.	Yes
31. at new or upgrading installations, where electricity generation is the priority over heat supply, the minimisation of condenser pressure, as described in 4.3.9	The turbine exhaust pressure (100 mbar, 22°C) and the air cooled condenser are designed so that the high ambient temperatures of Sydney can be handled.	Yes



ВАТ	TNG	BAT fullfilled?
 32. the general minimisation of overall installation energy demand, including consideration of the following (see 4.3.6): a. for the performance level required, the selection of techniques with lower overall energy demand in preference to those with higher energy demand b. wherever possible, ordering flue-gas treatment systems in such a way that flue gas reheating is avoided (i.e. those with the highest operational temperature before those with lower operational temperatures) c. where SCR is used; i. to use heat exchangers to heat the SCR inlet flue-gas with the flue-gas energy at the SCR outlet ii. to generally select the SCR system that, for the performance level required (including availability/fouling and reduction efficiency), has the lower operating temperature d. where flue-gas reheating is necessary, the use of heat exchange systems to minimise flue-gas reheating energy demand e. avoiding the use of primary fuels by using self produced energy in preference to imported sources 	To minimise the overall energy consumption the following measures were taken: - SNCR instead of SCR to avoid the flue gas reheating for the catalytic reaction - efficient preheating of primary and secondary air - minimal condenser pressure - placing high temperature equipment before (upstream) lower temperature equipment - use of frequency controlled rotating equipment for those equipment parts which operate at variable speeds. - high efficient DCS system to minimise the measure failure SCR is not relevant.	Yes
33. where cooling systems are required, the selection of the steam condenser cooling system technical option that is best suited to the local environmental conditions, taking particular account of potential cross-media impacts, as described in 4.3.10	As result of the local conditions and to minimize the water consumption an air cooled condenser has been chosen.	Yes
34. the use of a combination of on-line and off-line boiler cleaning techniques to reduce dust residence and accumulation in the boiler, as described in 4.3.19	Online cleaning devices for all parts of the boiler are foreseen. Offline cleaning is carried out manually during revisions.	Yes
35. the use of an overall flue-gas treatment (FGT) system that, when combined with the installation as a whole, generally provides for the operational emission levels for releases to air associated with the use of BAT listed in Table 5.2	The emission guarantees are in accordance with the European Industrial Emission Directive. The expected operational emission levels are in line with the values given in Table 5.2.	Yes



ВАТ	TNG	BAT fullfilled?
 36. when selecting the overall FGT system, to take into account: a. the general factors described in 4.4.1.1 and 4.4.1.3 b. the potential impacts on energy consumption of the installation, as described in section 4.4.1.2 c. the additional overall-system compatibility issues that may arise when retrofitting existing installations (see 4.4.1.4) 	The following factors have been taken account for the selection of the FGT system: - type of waste, its composition and variation - type of combustion process, and its size - through recirculation of flue gases less flue gas flow and lower flue gas temperature - flue gas composition and fluctuations in the composition - target emission limit values - restrictions on discharge of aqueous effluents - availability of land and space - availability and cost of outlets for residues accumulated/recovered - minimize consumables	Yes
37. when selecting between wet / semi-wet / and dry FGT systems, to take into account the (non-exhaustive) general selection criteria given as an example in Table 5.3 [See at the end of this document]	 The Flue gas cleaning process is characterised by the following features: No effluent as necessary for wet FGT minimized consumables and residues Dry injection of Calcium Hydroxide (Ca(OH)2) and Powdered Activated Carbon (PAC) Separate injection of water for conditioning and reactivation of recycled lime particles Compact design Low manpower requirement 	Yes
38. to prevent the associated increased electrical consumption, to generally (i.e. unless there is a specific local driver) avoid the use of. two bag filters in one FGT line (as described in 4.4.2.2 and 4.4.2.3)	The installation has only one bag filter.	Yes



BAT	TNG	BAT fullfilled?
 39. the reduction of FGT reagent consumption and of FGT residue production in dry, semi-wet, and intermediate FGT systems by a suitable combination of: a. adjustment and control of the quantity of reagent(s) injected in order to meet the requirements for the treatment of the flue-gas such that the target final operational emission levels are met b. the use of the signal generated from fast response upstream and/or downstream monitors of raw HCI and/or SO2 levels (or other parameters that may prove useful for this purpose) for the optimisation of FGT reagent dosing rates, as described in 4.4.3.9 c. the re-circulation of a proportion of the FGT residues collected, as described in 4.4.3.7 The applicability and degree of use of the above techniques that represents BAT will vary according to, in particular: the waste characteristics and consequential flue-gas nature, the final emission level required, and technical experience from their practical use at the installation 	 a) The quantity of reagents injected is regularly checked by measuring the target final operational emission levels. Depending on the emission level the reagents are injected. b) Measuring devices are installed to check the raw gas c) A part of the residues are recirculated into the semi dry reactor which results in a high efficiency, lower consumption of absorbent, protection of filter bags and a higher operational safety. 	Yes
40. the use of primary (combustion related) NOx reduction measures to reduce NOx production, together with either SCR (4.4.4.1) or SNCR (4.4.4.2), according to the efficiency of flue-gas reduction required. In general SCR is considered BAT where higher NOx reduction efficiencies are required (i.e. raw flue-gas NOx levels are high) and where low final flue-gas emission concentrations of NOX are desired. One MS reported that technical difficulties have been experienced in some cases when retrofitting SNCR abatement systems to existing small MSW incineration installations, and that the cost effectiveness (i.e. NOX reduction per unit cost) of NOX abatement (e.g. SNCR) is lower at small MSWIs (i.e. those MSWIs of capacity <6 tonnes of waste/hour).	In the first pass of the boiler ammonia is injected into the flue gas stream. The results are well controlled and low NOx emissions which are below the levels given by the Industrial Emissions Directive. The most important factors for NOx reduction measures are: - O ₂ content - flue gas recirculation - optimized secondary air injection Neither retrofitting nor small installation, therefore not relevant.	Yes



BAT	TNG	BAT fullfilled?
 41. for the reduction of overall PCDD/F emissions to all environmental media, the use of: a. techniques for improving knowledge of and control of the waste, including in particular its combustion characteristics, using a suitable selection of techniques described in 4.1, and b. primary (combustion related) techniques (summarised in 4.4.5.1) to destroy PCDD/F in the waste and possible PCDD/F precursors, and c. the use of installation designs and operational controls that avoid those conditions (see 4.4.5.2) that may give rise to PCDD/F reformation or generation, in particular to avoid the abatement of dust in the temperature range of 250 – 400 °C. Some additional reduction of de-novo synthesis is reported where the dust abatement operational temperature has been further lowered from 250 to below 200 °C, and d. the use of a suitable combination of one or more of the following additional PCDD/F abatement measures: i. adsorption by the injection of activated carbon or other reagents at a suitable reagent dose rate, with bag filtration, as described in 4.4.5.6, or ii. adsorption using fixed beds with a suitable adsorbent replenishment rate, as described in 4.4.5.7, or iii. multi layer SCR, adequately sized to provide for PCDD/F control, as described in 4.4.5.3, or iv. the use of catalytic bag filters (but only where other provision is made for effective metallic and elemental Hg control), as described in 4.4.5.4 	For the reduction of PCDD/F emissions the following techniques are applied: - optimized secondary air injection - maximal gas burn out - minimization of fly ash deposits in the boiler - continuous boiler cleaning - no dedusting equipment where the gas temperature is above 200°C - SNCR	Yes
42. where wet scrubbers are used, to carry out an assessment of PCDD/F build up (memory effects) in the scrubber and adopt suitable measures to deal with this build up and prevent scrubber breakthrough releases. Particular consideration should be given to the possibility of memory effects during shut down and start-up periods.	No wet scrubbers are used.	Not relevant
43. if re-burn of FGT residues is applied, then suitable measures should be taken to avoid the re-circulation and accumulation of Hg in the installation	No re-burn of FGT residues is applied.	Not relevant



BAT	TNG	BAT fullfilled?
44. for the control of Hg emissions where wet scrubbers are applied as the only or main effective means of total Hg emission control: a. the use of a low pH first stage with the addition of specific reagents for ionic Hg removal (as described in 4.4.6.1, 4.4.6.6 and 4.4.6.5), in combination with the following additional measures for the abatement of metallic (elemental) Hg, as required in order to reduce final air emissions to within the BAT emission ranges given for total Hg b. activated carbon injection, as described in 4.4.6.2, or c. activated carbon or coke filters, as described in 4.4.6.7	No wet scrubbers are used.	Not relevant
45. for the control of Hg emissions where semi-wet and dry FGT systems are applied, the use of activated carbon or other effective adsorptive reagents for the adsorption of PCDD/F and Hg, as described in 4.4.6.2, with the reagent dose rate controlled so that final air emissions are within the BAT emission ranges given for Hg	For the control of Hg emissions an activated carbon injection into the semi dry reactor is applied. The reagent dose rate is controlled to ensure that the final air emissions are within the BAT ranges given for Hg.	Yes
46. the general optimisation of the re-circulation and re-use of waste water arising on the site within the installation, as described in 4.5.8, including for example, if of sufficient quality, the use of boiler drain water as a water supply for the wet scrubber in order to reduce scrubber water consumption by replacing scrubber feed-water (see 4.5.6)	To avoid additional water consumption a complete recirculation of condensate is installed. The boiler drain water is used for the bottom ash quenching.	Yes
47. the use of separate systems for the drainage, treatment and discharge of rainwater that falls on the site, including roof water, so that it does not mix with potential or actual contaminated waste water streams, as described in 4.5.9. Some such waste water streams may require only little or no treatment prior to their discharge, depending on contamination risk and local discharge factors	A separate system is used for the drainage, treatment and discharge of rainwater so that it does not mix with potential or actual contaminated waste water streams.	Yes



BAT	TNG	BAT fullfilled?
48. where wet flue-gas treatment is used: a. the use of on-site physico/chemical treatment of the scrubber effluents prior to their discharge from the site, as described in 4.5.11, and thereby to achieve, at the point of discharge from the effluent treatment plant (ETP), emission levels generally within the BAT associated operational emission level ranges that are identified in Table 5.4 [See at the end of this document] b. the separate treatment of the acid and alkaline waste water streams arising from the scrubber stages, as described in 4.5.13, when there are particular drivers for the additional reduction of releases to water that result, and/or where HCl and/or gypsum recovery is to be carried out c. the re-circulation of wet scrubber effluent within the scrubber system, and the use of the electrical conductivity (mS/cm) of the re-circulated water as a control measure, so as to reduce scrubber water consumption by replacing scrubber feed-water, as de- scribed in 4.5.4 d. the provision of storage/buffering capacity for scrubber effluents, to provide for a more stable waste water treatment process, as described in 4.5.11 f. when SNCR is used with wet scrubbing the ammonia levels in the effluent discharge may be reduced using ammonia stripping, as described in 4.5.12, and the recovered ammonia re-circulated for use as a NOX reduction reagent 	No wet flue gas treatment is used.	Not relevant



ВАТ	TNG	BAT fullfilled?
 49. the use of a suitable combination of the techniques and principles described in 4.6.1 for improving waste burnout to the extent that is required so as to achieve a TOC value in the ash residues of below 3 wt % and typically between 1 and 2 wt %, including in particular: a. the use of a combination of furnace design (see combustion technology selection in 4.2.1), furnace operation (see 4.2.17) and waste throughput rate (see 4.2.18) that provides sufficient agitation and residence time of the waste in the furnace at sufficiently high temperatures, including any ash burn-out areas b. the use of furnace designs that, as far as possible, physically retain the waste within the combustion chamber (e.g. narrow grate bar spacings for grates, rotary or static kilns for appreciably liquid wastes) to allow its combustion. The return of early grate riddlings to the combustion chamber for re-burn may provide a means to improve overall burn out where they contribute significantly to the deterioration of burnout (see 4.2.21) c. the use of techniques for mixing and pre-treatment of the waste, as described in BAT 11, according to the type(s) of waste received at the installation d. the optimisation and control of combustion conditions, including air (oxygen) supply and distribution, as described in BAT 18 	The expected value of TOC is < 1%. A combination of different techniques/measures is used to improve the waste burnout: - optimal combustion conditions with a classic combustion control system, - flow optimised secondary combustion chamber, - visual check of the fire on the grate, - optimal design of the boiler geometry, - optimal temperature distribution on the grate, - optimal waste distribution over the grate - adjustment of particular grate speed - ensuring that the plant is operated within its capacity (fire control diagram) - good mixing of the waste before feeding	Yes
50. the separate management of bottom ash from fly ash and other FGT residues, so as to avoid contamination of the bottom ash and thereby improve the potential for bottom ash recovery, as described in 4.6.2. Boiler ash may exhibit similar or very different levels of contamination to that seen in bottom ash (according to local operational, design and waste specific factors) – it is therefore also BAT to assess the levels of contaminants in the boiler ash, and to assess whether separation or mixing with bottom ash is appropriate. It is BAT to assess each separate solid waste stream that arises for its potential for recovery either alone or in combination.	There is a separate management of bottom ash from fly ash and FGT residues. The FGT residues will be stored in separate enclosed silos before being transported by sealed tankers to an appropriate offsite treatment facility. This complies with the hazardous waste legislation.	Yes
51. where a pre-dedusting stage (see 4.6.3 and 4.4.2.1) is in use, an assessment of the composition of the fly ash so collected should be carried out to assess whether it may be recovered, either directly or after treatment, rather than disposed of	No pre-dedusting stage is installed.	Not relevant
52. the separation of remaining ferrous and non-ferrous metals from bottom ash (see 4.6.4), as far as practicably and economically viable, for their recovery	A magnetic separator is placed above a conveyor to remove ferrous metals from bottom ash.	Yes



BAT	TNG	BAT fullfilled?
 53. the treatment of bottom ash (either on or off-site), by a suitable combination of: a. dry bottom ash treatment with or without ageing, as described in 4.6.6 and 4.6.7, or b. wet bottom ash treatment, with or without ageing, as described in 4.6.6 and 4.6.8, or c. thermal treatment, as described in 4.6.9 (for separate treatment) and 4.6.10 (for in-process thermal treatment) or d. screening and crushing (see 4.6.5) to the extent that is required to meet the specifications set for its use or at the receiving treatment or disposal site e.g. to achieve a leaching level for metals and salts that is in compliance with the local environmental conditions at the place of use. 	The bottom ash is treated according to b) before reuse or landfill	Yes
54. the treatment of FGT residues (on or off-site) to the extent required to meet the acceptance requirements for the waste management option selected for them, including consideration of the use of the FGT residue treatment techniques described in 4.6.11	The FGT residues will be stored in separate enclosed silos before being transported by sealed tankers to an appropriate offsite treatment facility. This complies with the hazardous waste legislation.	Yes
55. the implementation of noise reduction measures to meet local noise requirements (techniques are described in 4.7 and 3.6)	Noise reduction measures are installed at the turbine-generator, at the fans and at other critical plant sections.	Yes
56. apply environmental management. A number of environmental management techniques are determined as BAT. The scope (e.g. level of detail) and nature of the EMS (e.g. standardised or non-standardised) will generally be related to the nature, scale and complexity of the installation, and the range of environmental impacts it may have. BAT is to implement and adhere to an Environmental Management System (EMS) that incorporates, as appropriate to individual circumstances, the following features: (see Chapter 4.8) [Number of bullet points omitted]	The plant is operated by qualified personnel and the staff will be sent to relevant qualifying courses. To ensure that employees are aware of the relevant parts of the environmental permit, regular dialogue meetings are held with the employees. The plant is ISO 14001 and OHSA 18001 certified. Additional information can be found in the quality manual where environmental measurements, reports and factors such as environmental requirements, responsibilities and competencies of facilities and technical installations are described. The operator is responsible to install the environmental system.	Yes
57. the storage of all waste, (with the exception of wastes specifically prepared for storage or bulk items with low pollution potential e.g. furniture), on sealed surfaces with controlled drainage inside covered and walled buildings	The waste is stored in the waste bunker. The waste bunker has sealed surfaces.	Yes



BAT	TNG	BAT fullfilled?	
58. when waste is stockpiled (typically for later incineration) it should generally be baled (see Section 4.1.4.3) or otherwise prepared for such storage so that it may be stored in such a manner that risks of odour, vermin, litter, fire and leaching are effectively controlled.	The risks of odour, vermin, litter, fire and leaching are effectively controlled because the waste is stockpiled in a closed building. There is no influence to the environment (only the open delivery boxes).	Yes	
 59. to pre-treat the waste, in order to improve its homogeneity and therefore combustion characteristics and burn-out, by: a. mixing in the bunker (see 4.1.5.1), and b. the use of shredding or crushing for bulky wastes e.g. furniture (see 4.1.5.2) that are to be incinerated, to the extent that is beneficial according to the combustion system used. In general grates and rotary kilns (where used) require lower levels of pre-treatment (e.g. waste mixing with bulky waste crushing) whereas fluidized bed systems require greater waste selection and pre- treatment, usually including full shredding of the MSW. 	The pre-treat of waste in order to improve its homogeneity is a) mixing of waste in the bunker b) shredding of waste.	Yes	
60. the use of a grate design that incorporates sufficient cooling of the grate such that it permits the variation of the primary air supply for the main purpose of combustion control, rather than for the cooling of the grate itself. Air-cooled grates with well distributed air cooling flow are generally suitable for wastes of average NCV of up to approx 18 MJ/kg. Higher NCV wastes may require water (or other liquid) cooling in order to prevent the need for excessive primary air levels (i.e. levels that result in a greater air supply than the optimum for combustion control) to control grate temperature and length/position of fire on the grate (see section 4.2.14)	For the grate a combination of water cooled blocks and air cooled blocks is used. The calorific value is below 18 GJ/t (usually 10-11 GJ/t).	Yes	
61. the location of new installations so that the use of CHP and/or the heat and/or steam utilisation can be maximised, so as to generally exceed an overall total energy export level of 1.9 MWh/tonne of MSW (ref. Table 3.42), based on an average NCV of 2.9 MWh/tonne (ref. Table 2.11)	The thermal efficiency is above 79,3 % (based on calculation of R1). Currently no heat or steam export is planned (however the plant is designed for heat export in case of future possibilities)	Yes	



BAT	TNG	BAT fullfilled?
62. in situations where less than 1.9 MWh/tonne of MSW (based on an average NCV of 2.9 MWh /tonne) can be exported, the greater of: a. the generation of an annual average of 0.4 – 0.65 MWh electricity/tonne of MSW (based on an average NCV of 2.9 MWh/tonne (ref. Table 2.11) processed (ref. Table 3.40), with additional heat/steam supply as far as practicable in the local circum- stances, or b. the generation of at least the same amount of electricity from the waste as the annual average electricity demand of the entire installation, including (where used) on-site waste pre-treatment and on-site residue treatment operations (ref. Table 3.48)	The annual average production is 1,02 MWh electricity/tonne of waste.	Yes
63. to reduce average installation electrical demand (excluding pre-treatment or residue treatment) to be generally below 0.15 MWh/tonne of MSW processed (ref. Table 3.47 and section 4.3.6) based on an average NCV of 2.9 MWh/tonne of MSW (ref. Table 2.11)	The electrical demand of the of the plant is 0,123 MWh electricity/tonne of waste.	Yes
64. the storage of wastes:a. in enclosed hoppers or,b. on sealed surfaces with controlled drainage inside covered and walled buildings	The waste is stored in the waste bunker. The waste bunker has sealed surfaces and is inside a closed building.	Yes
65. when waste is stockpiled (typically for later incineration) it should generally be baled (see Section 4.1.4.3) or otherwise prepared for such storage so that it may be stored in such a manner that risks of odour, vermin, litter, fire and leaching are effectively controlled	see point 58.	
66. at new and existing installations, the generation of the greater of: a. an annual average of generally at least 0.6 – 1.0 MWh electricity/tonne of waste (based on an average NCV of 4.2 MWh/tonne), or b. the annual average electricity demand of the entire installation, including (where used) on-site waste pretreatment and on-site residue treatment operations	The annual average production is 1,02 MWh electricity/tonne of waste.	Yes



BAT	TNG	BAT fullfilled?
67. the location of new installations so that: a. as well as the 0.6 – 1.0 MWhe/ tonne of electricity generated, the heat and/or steam can also be utilised for CHP, so that in general an additional thermal export level of 0.5 – 1.25 MWh/tonne of waste (ref. section 3.5.4.3) can be achieved (based on an average NCV of 4.2 MWh/tonne), or b. where electricity is not generated, a thermal export level of 3 MWh/tonne of waste can be achieved (based on an average NCV of 4.2 MWh/tonne)	The annual average production is 1,02 MWh electricity/tonne of waste.	Yes
68. to reduce installation energy demand and to achieve an average installation electrical demand (excluding pretreatment or residue treatment) to generally below 0.2 MWh/tonne of waste processed (ref. Table 3.47 and section 4.3.6) based on an average NCV of 4.2 MWh/tonne of waste	The electrical demand of the plant is 0,123 MWh electricity/tonne of waste.	Yes





Job

Date From TNG Energy from Waste Facility, Eastern Creek, Differences between quoted design Ash outputs and operational ash outputs 2016-10-24 Ahmet Erol

Differences between quoted design Ash outputs and operational ash outputs

Introduction

TNG have been asked to provide an explanation for the differences between quoted design ash outputs and operational ash outputs from the reference facilities.

Factors influencing bottom ash quantity of EfW plants

The design of an EfW plant is based on the waste amount, the waste composition and the calorific value (CV) of the waste.

An important aspect of the waste composition is the ash content which is made up of the ash of combustible fuel (ash of wood, paper, etc.) as well as inert materials like metal, glass, stone, sand etc.

Depending on the nature of the waste the ash content varies. Even different waste streams with identical CV can have a variation of the ash content. The CV is mainly influenced by the relation between combustible, water and inert. A waste with high water but low ash content can have the same CV as a waste with high ash and low water content.

As a result there can be a certain variation (usually +/- 3-5% points) of the ash in real operation compared to the design value even if a plant is running at its designed throughput capacity.

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File: TNGWTE-141-015 Explanation design Ash and operational Ash outputs.docx Ver. 3

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Ash outputs

						Bottom Ash Dry %	
Facility/Location	LHV @ LPN MJ/kg	Design t/a	Fuel mix	bottom ash (wet) t/a	bottom ash Quantity (wet) %	Operation	Design
TNG	12.3	4 x 276'250	C&I, C&D	293'166	26.5%	-	21.49%
Grossräschen	12.5	1 x 246'000	C&I, C&D	68'729	27.9%	22.4%	18.8%
Knapsack	11-17	2 x 150'000	C&I, C&D	81'000	27.0%	21.6%	19.0%
Ferrybridge	8.5	2 x 256'500	C&I, C&D, some MSW, wood	57'830	11.3%	9.0%	12.8%
Riverside	9.6	3 x 195'000	MSW, C&I	146'250	25.0%	20.0%	19.7%
TIRME Mallorca	10	2 x 208'000	MSW, C&I, C&D, Hospital waste, sewage sludge, tyres	92'350	22.2%	17.8%	20.0%

The above table shows updated the design and operational values of several EfW plants. The operation and design bottom ash quantities are all within a variation range of 3-4%.

The ash quantities in case of TIRME Mallorca have been derived from the Environmental Master plan (design) and information received from the operator (operation). The data given in earlier memos was data from the initial design fuel and the reference sheet of the supplier which were obviously outdated.
Pacific Environment Limited

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Appendix D FUEL COMPOSITION

	Units	CRW	C&D	C&I	Floc waste	Paper	Glass	GO	AWT	MRF	Design
						Pulp	Recovery	Residual	Residual	Residual	Fuel Mix
Fuel Mix	%	23.37%	28.69%	16.84%	14.43%	4.81%	1.72%	2.06%	6.87%	1.20%	100
Compositional Analysis											
Paper/Card	%	4.30	14.05	22.44	3.93	78.40	62.00	30.00	21.05	38.54	16.75
Plastic Film	%	10.20	6.37	10.90	10.90	21.60	3.80	2.50	20.00	26.94	10.47
Dense Plastic	%	0.00	6.37	10.90	10.90	0.00	34.20	2.50	21.05	0.00	7.32
Textiles	%	5.30	0.00	12.89	0.18	0.00	0.00	0.00	10.53	0.00	4.16
Glass	%	0.00	0.00	1.81	0.00	0.00	0.00	4.00	0.00	8.50	0.49
Vegetation	%	8.30	0.00	1.70	0.00	0.00	0.00	35.00	3.16	0.00	3.16
Other combustibles	%	0.00	0.00	0.00	70.40	0.00	0.00	0.00	0.00	0.00	10.16
Metal	%	1.80	1.12	0.37	0.00	0.00	0.00	5.00	0.00	7.59	1.00
Fines	%	0.00	0.94	0.18	0.00	0.00	0.00	0.00	11.58	0.00	1.10
Wood	%	58.20	43.90	21.53	0.85	0.00	0.00	0.00	4.21	0.00	30.24
Combustibles	%	0.00	0.00	2.84	2.84	0.00	0.00	0.00	2.11	0.00	1.03
Non-Combustibles	%	4.50	0.00	0.00	0.00	0.00	0.00	21.00	1.05	0.03	1.56
Hazardous	%	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Gyprock	%	2.40	6.50	0.00	0.00	0.00	0.00	0.00	0.00	0.00	2.42
Other	%	5.00	20.75	14.44	0.00	0.00	0.00	0.00	5.26	18.40	10.14
Total	%	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00
Chemical Analysis											
Carbon (C)	%	31.34	27.02	35.00	29.65	42.90	41.01	16.98	38.96	32.63	31.44
Hydrogen (H)	%	4.21	3.51	4.29	3.80	5.84	4.63	2.12	4.98	4.84	4.07
Nitrogen (N)	%	0.34	0.06	0.59	0.18	0.00	0.00	0.12	0.47	0.00	0.26
Sulphur(S)	%	0.42	1.04	0.05	0.11	0.12	0.09	0.06	0.04	0.06	0.43
Chloride (Cl)	%	0.09	0.66	1.15	1.78	0.19	3.27	0.26	2.18	0.23	0.88
Oxygen (O)	%	21.11	21.50	17.50	7.04	24.64	26.69	12.58	13.77	12.11	18.06
Water (H2O)	%	28.47	21.51	21.68	22.62	22.58	20.81	36.20	18.40	15.20	23.38
Ash	%	14.03	24.70	19.74	34.82	3.73	3.50	31.68	21.20	34.93	21.49
Total	%	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00
NCV	MJ/kg	11.95	9.97	13.84	12.59	17.22	15.24	5.67	16.33	14.23	12.30

Table D-1: Proposed design fuel analysis, as received basis (Ramboll, 2016)

Table D-2: Fuel mix (Ramboll, 2016)

Fuel component		Minimum	Maximum
Nitrogen(N)		0.19	0.34
Sulphur(S)		0.31	0.57
Chloride (Cl)	%	0.64	0.96
Ash		13	26
Water(H ₂ O)		14	34
NCV	MJ/kg	8.5	16.5

ENERGY

1 RAMBOLL MEMO – DESIGN FUEL MIX





MEMO

Job Date

From

TNG Energy from Waste Facility, Eastern Creek, Differences Design fuel mix between UTDI (Nov 15) and Concept of Design (March 15) reports 2016-10-24 Martin Brunner, Ahmet Erol

Differences of the Design fuel mix between UTDI (Nov 15) and Concept of Design (March 15) reports

Introduction

EPA requested an explanation why there are such differences in the design fuel mix between Updated Technical Design Information (UTDI Nov 15) report and the Concept of Design (March 15) report and why it has not had an impact on the percentage make-up of the design fuel mix.

Explanation

Ramboll was asked to review the basis design fuel in the Fichtner Concept Design Report. As a basis for this review Ramboll was provided a comprehensive report with description and pictures of the waste streams treated at the Genesis plant and the products produced by the screening and sorting process. Based on Ramboll's experience the composition of every product and every waste stream to the future TNG facility was evaluated and redefined. Ramboll Hannemanns Allé 53 DK-2300 Copenhagen S Denmark

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Composition of waste streams by products

The following table shows the composition of the design fuel based on products (paper, plastic, etc.) in the different waste streams (CRW, C&D, etc.) according to the Concept Design Report (CDR).

		Fab	le 1: Propo	osed design	i fuel analy	sis, as rec	eived basis				
	LINUS.	ERW	CaD	CNI	Flocit: waste	Paper Pulp	Glass Recovery	ISO Residual	AWT Residual	MRF Residual	Design Fuel Mix
Fuel Mix		23.37%	28.69%	16.84%	14.43%	4.81%	1.72%	2.06%	6.87%	1.20%	100
				Compo	sitional Anal	ysis					-
Paper/Card	5	4.30	14.05	22.44	3.93	78.40	62.00	30.00	21.05	38.54	16.75
Plastic Film	- W-	10.20	6.37	10.90	10.90	21.60	3.80	2.50	20.00	26,94	10.47
Dense Plastic	54	0.00	6.37	10.90	10.90	0.00	34.20	2.50	21.05	0.08	7.32
Textiles		5.30	0.00	12.89	0.18	0,00	0.00	0.00	10.53	0.00	4.16
Glass		0.00	0.00	1.81	0.00	0.00	0.00	4.00	0.00	8.50	0.49
Vegetation	1.1	8.30	0.00	1.70	0.00	0,00	0.00	35.00	3.16	0.00	3.16
Other putrescibles	5	0.00	0.00	0.00	70,41	0,00	0,00	0.00	0.00	0.00	10.16
Metal	- 56	1.80	1.12	D.37	0,00	0.00	0.00	5.00	0.00	7,59	1.00
Fines		0.00	0.94	0.18	0.00	0.00	0.00	0.00	11.58	0.00	1.10
Wood	. N.	58.20	41.91	21.53	0.85	0.00	0.00	0.00	4.21	0.08	30.24
Combustibles		0.00	0.00	2.84	2.84	0.00	0.00	0.00	2.11	0.00	1.03
Non-Combustibles		6.90	6,50	0,00	0,00	0,00	0.00	21.00	1,05	0.03	3.98
Hazardous		0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Other	5	5.00	20.75	14.44	0.00	0.00	0.00	0.00	5.26	18.40	10.14
Total		100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00

Table 1 Fichtner Concept Design Report (March 2015), Design fuel analysis

Based on the description and pictures of the waste streams it was obvious, that one important product in the CRW and C&D waste (so called Gyprock, waste from gypsum building material) had not been considered. As gypsum contains sulphur (CaSO₄), which contributes to the SO₂ content of the flue gas, this waste stream was added to the list of products. To compensate for this change the fraction of non-combustible was reduced. The updated composition (changes shown in green) is shown in Table 2 below.

Ramboll revised	Material Sour	ces and Com	positions							
Fuel Mix	136000 23.37%	167000 28.69%	98000 16.84%	84000 14.43%	28000 4.81% e Paper Pulp	10000 1.72%	12000 2.06%	40000 6.87%	7000 1.20%	582000 100%
	CRW	C&D	C&I	Flock Waste		Glass Residual	GO Residual	AWT Residual	MRF Residual	Design Fuel Mix
Paper/card	4.30%	14.05%	22.44%	3.93%	78.40%	62.00%	30.00%	21.05%	38.54%	16.75%
Plastic film	10.20%	6.37%	10.90%	10.90%	21.60%	3.80%	2.50%	20.00%	26.94%	10.47%
Dense plastic	0.00%	6.37%	10.90%	10.90%	0.00%	34.20%	2.50%	21.05%	0.00%	7.32%
Textiles	5,30%	0.00%	12,89%	0.18%	0,00%	0.00%	0.00%	10.53%	0.00%	4.16%
Glass	0.00%	0.00%	1.81%	0.00%	0.00%	0.00%	4.00%	0.00%	8.50%	0.49%
Vegetation	8.30%	0.00%	1.70%	0.00%	0.00%	0.00%	35.00%	3.16%	0.00%	3.16%
Other combustibles	0.00%	0.00%	0.00%	70.41%	0.00%	0.00%	0.00%	0.00%	0.00%	10.16%
Metal	1.80%	1.12%	0.37%	0.00%	0.00%	0.00%	5.00%	0.00%	7.59%	1.00%
Fines	0.00%	0.94%	0.18%	0.00%	0.00%	0.00%	0.00%	11.58%	0.00%	1.10%
Wood	58.20%	43.91%	21.53%	0.85%	0.00%	0.00%	0.00%	4.21%	0.00%	30.24%
Combustibles	0.00%	0.00%	2,84%	2.84%	0.00%	0.00%	0.00%	2.11%	0.00%	1.03%
Non-combustibles	4.50%	0.00%	0.00%	0.00%	0.00%	0.00%	21.00%	1.05%	0.03%	1.56%
Hazardous	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Gyprock	2.40%	6.50%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	2.43%
Other	5.00%	20.75%	14.44%	D.00%	0.00%	0.00%	0.00%	5.26%	18.40%	10.14%
Total	100.00%	100.00%	100.00%	100.01%	100.00%	100.00%	100.00%	100.00%	100.00%	100.00%

Table 2 Ramboll Updated Technical Design Information (UTDI Nov 15), Design fuel mix revised



Composition of waste streams chemical analysis

The following table shows the initial composition of the design fuel according to the CDR.

	Units	CRW	CED	CAI	Floc waste	Paper Pulp	Elass Recovery	GO Residual	AWT Residual	MHF Residual	Design Fuel Mix
					Chemical	Analysis			_		
Carbon (C)	%	37.37	38.90	40.05	23.44	35.31	40.32	18.53	38.81	30.87	35.83
Hydrogen (H)	%	4.78	5.02	5,40	3,30	5.11	5,61	2.50	5.37	4.53	4.76
Nitrogen (N)	%	0.90	0.71	0.95	0.90	0.37	0.42	0.50	0.85	0.34	0.80
Sulphur(S)	96	0.12	0.12	0,12	0.12	0.12	0.11	0.18	0.17	0.12	0.12
Chloride (CI)	%	0.17	0.66	1.09	1.03	0.39	2,35	0.39	1,73	0.41	0.75
Oxygen (O)	9/6	28.21	28.28	24,86	11.84	24.57	22.87	14.30	16.68	15.78	23.81
Water (H2O)	9%	17.12	13.81	18.57	50.00	25.00	19,57	31.65	23.77	21.96	22.39
Ash	%	11.33	12.50	8.97	9.36	9.14	8.75	31.95	12.63	25.97	11.53
Total	%	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00
NCV	MJ/kg	13.68	12.08	13.46	8.46	13.22	16.13	6.31	14.86	10.09	12.34

Table 3: Fichtner Concept Design Report (March 2015), Chemical Analysis

In order to reassess the design fuel, every single product was analysed and its chemical composition redefined based on a database of comparable products. The chemical composition of each product as used for the assessment is shown in the appendix of this memo. The chemical composition of each waste stream was then calculated based on the chemical composition of its products. The results of the redefinition are shown in the table below.

	Units	CRW	CSD	681	Floc waste	Paper Pulp	Glass Recovery	GO Residual	AWT Residual	MRF Residual	Design Fuel Mix
					Chemica	Analysis					
Carbon (C)	%	31.34	27.02	35	29.65	42.9	41.01	16.98	38.96	32.63	31.44
Hydrogen (H)	%	4.21	3.51	4.29	3.8	5.84	4.63	2.12	4.98	4.84	4.07
Nitrogen (N)	%	0.34	0.06	0.59	0.18	0	0	0.12	0.47	.0	0.26
Sulphur(S)	%	0.42	1.04	0.05	0.11	0.12	0.09	0.06	0.04	0.06	0.43
Chloride (CI)	%	0.09	0.66	1,15	1,78	0.19	3.27	0.26	2,18	0.23	0.88
Oxygen (0)	9/0	21.11	21.5	17.5	7.04	24.64	26.69	12.58	13.77	12.11	18.06
Water (H2O)	96	28.47	21.51	21.68	22,62	22.58	20,81	36.2	18.4	15.2	23.38
Ash	9%	14.03	24.7	19.74	34.82	3,73	3.5	31.68	21.2	34.93	21.49
Total	%	100	100	100	100	100	100	100	100	100	100
NCV	MJ/kg	11.95	9.97	13.84	12.59	17.22	15.24	5.67	16.33	14.23	12.3

Table 4: Ramboll Updated Technical Design Information (UTDI Nov 15), Chemical Analysis revised

The following tables summarize the overall chemical analysis in the Fichtner CDR (March 2015) and the Ramboll UTDI (Nov 2015).

		c	н	0	N	5	d	Ash	H ₂ O	NCV
Concept of Design (March 15) report		35.83%	4.76%	23.81%	0.80%	0.12%	0.75%	11.53%	22.39%	12.34 MJ/kg
UTDI (Nov 15) report	100.0%	31,4%	4.1%	18.1%	0.3%	0.4%	0.9%	21.5%	23.4%	12.30 MJ/kg

Table 5: Comparison of Fichtner CDR and Ramboll UTDI

The major change is the increased ash content. This is mainly a result of increased inert content in C&D, C&I and floc waste. In C&D the inert is mainly coming from adherent mineral substances like gypsum or mortar. The ash in floc waste is made up from paints, dirt, fine metals and rust.

As a result of the higher ash content the percentage of combustible (mainly C, H, O) is reduced accordingly. There further is an increase of the sulphur and chloride content. Both are important parameters for the design of the air pollution control equipment and need to be chosen carefully.

The decrease of oxygen is more than proportional and a result of the chemical reassessment of the different products. As oxygen has a negative impact on the calorific value, the reduction of oxygen results in a constant overall CV – irrespective of the decrease of combustible.

3/5



Appendix

Chemical analysis and CV calculation of each waste stream.

CHW		r		0	N	c	n	Ach	H.O	NOV
Danasland	4 2054	35.35	4.15/	31 49/	0.0%	.0.29/	0.0%	4.19/	35.0%	12.15 681/60
Paper/caru Diagtic film	4.30%	33.376	4,170	0.0%	0.0%	0.2%	0.0%	4.1.20	13.9%	25.50 641/kg
Donse plastic	0.00%	48.2%	A 7%	21.1%	0.0%	0.0%	9.5%	2.0%	14.0%	18 58 AN/kg
Textiles	5 30%	40.270	4.720	10.5%	4 392	0.0%	0.0%	2.0%	25.0%	19.36 MU/kg
Glass	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MI/kg
Vegetation	8 30%	9.8%	1.3%	7.5%	0.3%	0.0%	0.0%	1.0%	80.0%	1.92 Mi/kg
Other combustibles	0.00%	20.4%	2.4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Metal	1.80%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Fines	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Wood	58.20%	33.0%	4.2%	30.4%	0.1%	0.1%	0.0%	2.8%	29.3%	11.50 MJ/kg
Combustibles	0.00%	20.4%	2.4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Non-combustibles	4.50%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Hazardous	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Gyprock	2.40%	0.0%	0.0%	36.8%	0.0%	15.2%	0.0%	28.0%	20.0%	-2.87 MI/kg
Other	5.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	90.0%	10.0%	-0.24 MJ/kg
UTDL (Nov 15) report	100.0%	31.3%	4.7%	21.1%	0.3%	0.8%	0.1%	14.0%	29.5%	11.95 MI/kg
Constant of Day ins (March 15) suggest	300.00	37.45	A TW	70.70	0.000		0.36	11.00		17 CE MILLIN
concept of Design (Warch 15) report	100.0%	37.4%	4.074	25,670	0.9%	1.15	0,270	40.00	3 Aug	13'90 MD/KB
									2.5	
CAD		c	н	0	N	5	d	Ash	H ₂ O	NCV
Paper/card	14.05%	35.3%	4.1%	31.4%	0.0%	0.2%	0.0%	4.1%	25.0%	12.15 MJ/kg
Plastic film	6.37%	70.7%	12.1%	0.0%	0.0%	0.0%	0.9%	2.6%	13.8%	35.59 MU/kg
Dense plastic	6.37%	48.2%	4.7%	21.1%	0.0%	0.0%	9.5%	2,6%	14.0%	18.58 MJ/kg
Textiles	0.00%	48.8%	4.1%	10.5%	4.3%	0.0%	0.0%	7.5%	25.0%	19.36 MJ/kg
Glass	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MU/kg
Vegetation	0,00%	9.8%	1.3%	7.5%	0.3%	0.0%	0.0%	1,0%	80.0%	1.92 MU/kg
Other combustibles	0.00%	20.4%	2.4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Metal	1.12%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Fines	0.94%	0.0%	D.0%	0.0%	0.0%	0.0%	0.0%	100,0%	0.0%	0.00 MJ/kg
Wood	43,91%	33.0%	4.2%	30.4%	0.1%	0.1%	0.0%	2.8%	29.3%	11.50 MJ/kg
Combustibles	0.00%	20,4%	2,4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Non-combustibles	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Hazardous	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MU/kg
Gyprock	6.50%	0.0%	0,0%	35.8%	0.0%	15.2%	0.0%	28.0%	20,0%	-2.87 MJ/kg
Other	20,75%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	90.0%	10.0%	-0.24 MJ/kg
UTDI (Nov 15) report	100,0%	27.0 -	3,5%	21.5%	0.1%	II.(0%6	0.7%	24.7%	21.5%	9.97 MJ/kg
Concept of Design (March 15) report	100.0%	100	5,0%	28.3%	0.7%	0.1%	0.7W.		-0.00	12.08 MJ/kg
	-	_				-				
(8)		c	н	0	N	5	d	Ash	H.O	NCV
Paper/card	22.44%	35.4%	4.1%	31.4%	0.0%	0.2%	0.0%	4.1%	25.0%	12.15 MJ/kg
Plastic film	10 90%	70.7%	12 1%	0.0%	0.0%	0.0%	0.9%	2.6%	13.8%	35.59 MI/kg
Dense plastic	10 90%	48.2%	4.7%	21.1%	0.0%	0.0%	9.5%	2.6%	14.0%	18.58 MI/kg
Textiles	12.89%	48.8%	4.1%	10.5%	4.3%	0.0%	0.0%	7.5%	25.0%	19.36 MI/kg
Glass	1.81%	0.0%	D.0%	0.0%	0.0%	0.0%	0.0%	100.0%		and a real of
Vegetation	1,70%	9.8%	1 396	7 580					0.0%	0.00 MI/ke
Other combustibles	0.00%	10 Capitor		1.370	0.3%	0.0%	0.0%	1.0%	0.0%	0.00 MJ/kg 1.92 MJ/kg
	0,0010	20.4%	2.4%	4.4%	0.3%	0.0%	0.0%	1.0%	0.0% 80.0% 25.0%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg
Metal	0.37%	20.4%	2.4%	4.4%	0.3%	0.0%	0.0%	1.0% 46.5% 100.0%	0.0% 80.0% 25.0%	0.00 MI/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg
Metal	0.37%	20.4%	2.4% 0.0%	4.4% 0.0%	0.3% 0.2% 0.0% 0.0%	0.0%	0.0%	1.0% 46.5% 100.0%	0.0% 80.0% 25.0% 0.0%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg
Metal Fines Wood	0.37% 0.18% 21.53%	20.4% 0.0% 0.0% 33.0%	2.4% 0.0% 0.0% 4.2%	4.4% 0.0% 0.0% 30.4%	0.3% 0.2% 0.0% 0.0% 0.0%	0.0% 0.2% 0.0% 0.0% 0.1%	0.0% 0.9% 0.0% 0.0%	1.0% 46.5% 100.0% 100.0% 2.8%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3%	0.00 MI/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 11.50 MJ/kg
Metal Fines Wood Combustibles	0.37% 0.18% 21.53% 2.84%	20.4% 0.0% 0.0% 33.0% 20.4%	2.4% 0.0% 0.0% 4.2% 2.4%	4.4% 0.0% 0.0% 30.4% 4.4%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2%	0.0% 0.2% 0.0% 0.0% 0.1% 0.2%	0.0% 0.9% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 100.0% 2.8% 46.5%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 25.0%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 11.50 MJ/kg 8.30 MJ/kg
Metal Fines Wood Combustibles Non-combustibles	0.37% 0.18% 21.53% 2.84% 0.00%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0%	7.3% 4.4% 0.0% 0.0% 30.4% 4.4%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0%	0.0% 0.2% 0.0% 0.1% 0.2% 0.2%	0.0% 0.9% 0.0% 0.0% 0.0% 0.9%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 11.50 MJ/kg 8.30 MJ/kg 0.00 MJ/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous	0.37% 0.18% 21.53% 2.84% 0.00% 0.00%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0% 0.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0%	4.4% 0.0% 0.0% 30.4% 4.4% 0.0%	0.3% 0.2% 0.0% 0.1% 0.2% 0.2% 0.0%	0.0% 0.2% 0.0% 0.1% 0.2% 0.2% 0.0%	0.0% 0.9% 0.0% 0.0% 0.9% 0.9% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0%	0.00 Mi/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 11.50 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Goyprock	0.37% 0.18% 21.53% 2.84% 0.00% 0.00%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0%	4.4% 0.0% 0.0% 30.4% 4.4% 0.0% 0.0% 36.8%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0%	0.0% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 15.2%	0.0% 0.9% 0.0% 0.0% 0.9% 0.9% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0%	0.00 Mu/kg 1.92 Mu/kg 8.30 Mu/kg 0.00 Mu/kg 11.50 Mu/kg 8.30 Mu/kg 0.00 Mu/kg 0.00 Mu/kg 0.00 Mu/kg 7.87 Mu/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprock Other	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0%	4.4% 0.0% 0.0% 30.4% 4.4% 0.0% 0.0% 36.8% 0.0%	0.3% 0.2% 0.0% 0.1% 0.1% 0.2% 0.0% 0.0%	0.0% 0.2% 0.0% 0.1% 0.2% 0.0% 0.0%	0.0% 0.9% 0.0% 0.0% 0.0% 0.9% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 10.0%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 11.50 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.287 MJ/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprock Other UTDI (Nov 15) report	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0%	7.3% 4.4% 0.0% 30.4% 4.4% 0.0% 0.0% 36.8% 0.0% 17.5%	0.3% 0.2% 0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.2% 0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 15.2% 0.0% 0.1%	0.0% 0.9% 0.0% 0.0% 0.9% 0.9% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0% 19.7%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 10.0% 21.7%	0.00 Mu/kg 1.92 Mu/kg 8.30 Mu/kg 0.00 Mu/kg 11.50 Mu/kg 8.30 Mu/kg 0.00 Mu/kg 0.00 Mu/kg 0.00 Mu/kg 0.00 Mu/kg -2.87 Mu/kg 13.84 Mu/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprosk Gyprosk Other UTDI (Nov 15) report	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14,44% 100.0%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0%	2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 4.3%	7.3% 4.4% 0.0% 30.4% 4.4% 0.0% 0.0% 36.8% 0.0% 17.5%	0.3% 0.2% 0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0%	0.0% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.9% 0.0% 0.0% 0.9% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0% 49.7%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 25.0% 0.0% 20.0% 10.0% 21.7%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.287 MJ/kg 13.84 MJ/kg 13.84 MJ/kg
Metal Fines Combustibles Non-combustibles Hatardous Gyprock UTDI (Nov 15) report Concept of Design (March 18) report	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0% 40.1	2.3% 2.4% 0.0% 0.0% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 4.3% 5.4%	7.3% 4.4% 0.0% 30.4% 4.4% 0.0% 36.8% 0.0% 17.5%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.6% 1.0%	0.0% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.1% 0.1%	0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0% 19.7%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 10.0% 21.7%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 1.50 MJ/kg 0.00 MJ/kg 0.00 MJ/kg -2.87 MJ/kg -0.24 MJ/kg 13.84 MJ/kg 13.46 MJ/kg
Metal Finas Combustibles Non-combustibles Hatardous Gyprock Other UTDI (Nov 15) report Cloncept of Design (March 15) report	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0% 40.1	2.4% 2.4% 0.0% 0.0% 2.4% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 5.4%	4.4% 0.0% 0.0% 30.4% 4.4% 0.0% 36.8% 0.0% 17.5%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.1% 0.1%	0.0% 0.9% 0.0% 0.0% 0.9% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 20.0% 20.0% 20.0% 21.7%	0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg 0.00 MJ/kg 11.50 MJ/kg 11.50 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 13.84 MJ/kg 13.84 MJ/kg
Metal Fines Combustibles Non-combustibles Non-combustibles Hazardous Gypred UTDI (Nov 15) report UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste	0.37% 0.18% 21.53% 21.53% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0% 40.1	2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 4.3% 5.4%	4.3% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 14.3% 0	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.6% 1.0%	0.0% 0.2% 0.0% 0.1% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.1% 0.1% 0.1%	0.0% 0.9% 0.0% 0.0% 0.9% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0% 49.7% 49.7% 49.7%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 25.0% 0.0% 20.0% 20.0% 20.0% 10.0% 21.7%	0 00 Mi/kg 1.92 Mi/kg 8.30 Mi/kg 0.00 Mi/kg 11.50 Mi/kg 11.50 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 13.84 Mi/kg 13.84 Mi/kg 13.84 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprodus Gyprodus Other UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 10.00% 14.44% 100.0% 100.0%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0% 40.1 C 35.3%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 4.3% 5.4% H 4.1%	4.4% 0.0% 0.0% 30.4% 4.4% 0.0% 36.8% 0.0% 17.5% 14.9%	0.3% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.2% 0.0% 0.0% 0.1% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 28.0% 90.0% 19.7% 1.0% Ash 4.1%	0.0% 80.0% 25.0% 0.0% 0.0% 25.0% 25.0% 0.0% 20.0% 10.0% 21.7% 24.7% 25.0%	0 00 M/kg 1 92 M/kg 8 30 M/kg 0 00 M/kg 0 00 M/kg 11.50 M/kg 8 30 M/kg 0 00 M/kg 0 00 M/kg 13.50 M/kg 13.84 M/kg 13.84 M/kg NCV
Metal Finas Wood Combustibles Hazardous Gyprock Other UTDI (Nov 15) report Concept of Design (March 13) report Flock Waste Paper/card Plastic film	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0% 100.0%	20.4% 0.0% 0.0% 20.4% 20.4% 0.0% 0.0% 0.0% 35.0% 4g.(====================================	2.4% 2.4% 0.0% 0.0% 2.4% 0.0% 0.0% 0.0% 0.0% 5.4% 5.4% H 4.1% 12.1%	7 35% 4 44% 0.0% 0.0% 30.4% 4 4% 0.0% 30.6% 36.8% 0.0% 17.5% 14.9% 0.0% 14.9%	0.3% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.9% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 28.0% 90.0% 100.0% 100.0% 28.0% 90.0% 100.0% 4.1% 4.1% 2.6%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 10.0% 20.0% 10.0% 21.7% 20.0% 11.8%	0 00 M/kg 1 92 M/kg 8.30 M/kg 0.00 M/kg 0.00 M/kg 0.00 M/kg 0.00 M/kg 0.00 M/kg 11.50 M/kg 0.00 M/kg 13.46 M/kg 13.46 M/kg 13.46 M/kg
Metal Fines Wood Combustibles Non-combustibles Haardous Gyprot UTDI (Nov 15) report UTDI (Nov 15) report UTDI (Nov 15) report Flock Waste Paper/card Plastic film Dense plastic.	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0% 3.93% 10.90% 10.90%	20.4% 0.0% 33.0% 20.4% 20.4% 0.0% 0.0% 0.0% 0.0% 35.0% 40.1% 5.3% 70.7% 48.2%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 5.4% H 4.1% 12.1% 4.7%	7 35% 4 43% 0.0% 0.0% 30.4% 4 43% 0.0% 0.0% 17.5% 14.3% 0.0% 17.5% 14.3%	0.3% 0.2% 0.0% 0.0% 0.0% 0.2% 0.2% 0.2% 0.2	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.9% 0.0% 0.0% 0.0% 0.9% 0.9% 0.0% 0.0	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0% 100.0% 28.0% 90.0% 19.7% 4.5% 2.6%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 25.0% 0.0% 20.0% 20.0% 21.7% 20.0% 21.7% 23.9% 25.0% 13.8% 14.0%	0 00 M//kg 1 92 M//kg 8 30 M//kg 0 00 M//kg 0 00 M//kg 0 00 M//kg 0 00 M//kg 0 00 M//kg 0 00 M//kg 13.84 M//kg 13.84 M//kg 13.84 M//kg 15.59 M//kg 18.58 M//kg
Metal Fines Wood Combustibles Non-combustibles Haardous Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 1.09.0% 10.90% 0.19% 0.19% 0.19%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 4.2% 48.2% 48.8%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 6.0% 5.4% H H 4.1% 5.4% 4.2% 4.2% 0.0%	7 35% 7 4 44% 0.0% 0.0% 30.4% 4 4% 0.0% 30.4% 4 4% 0.0% 17.5% 0.0% 17.5% 0.0% 17.5% 0.0% 17.5% 0.0% 17.5% 0.0% 0	0.3% 0.2% 0.0% 0.0% 0.0% 0.2% 0.2% 0.2% 0.2	0.0% 0.2% 0.0% 0.0% 0.0% 0.0% 0.2% 0.0% 0.2% 0.0% 0.0	0.0% 0.9% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 28.0% 90.0% 19.7% 4.3% 2.6% 2.6% 7.5% 2.6%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 21.7% 21.7% 24.0% 25.0% 13.8% 14.0% 25.0%	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 11.50 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.559 Mi/kg 19.36 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprack Other UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic. Textiles Glass	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 10.00% 100.0% 10.00% 10.90% 0.18% 0.05%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0	2.4% 2.4% 0.0% 4.2% 2.4% 2.4% 0.0% 0.0% 0.0% 0.0% 5.4% 5.4% H 4.1% 12.1% 4.7% 4.2%	7 35% 7 4.4% 0.0% 0.0% 30.4% 4.4% 0.0% 0.0% 36.8% 0.0% 17.5% 14.9% 0.0% 14.9% 0.0% 14.9% 0.0% 14.9% 0.0% 14.9% 0.0% 14.9% 0.0% 14.4% 0.0% 0.0% 14.4% 0.0%	0.3% 0.2% 0.0% 0.0% 0.3% 0.3% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.2% 0.0% 0.0	0.0% 0.9% 0.9% 0.0% 0.0% 0.0% 0.0% 0.9% 0.0% 0.0	1.0% 46.5% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 25.0% 0.0% 0.0% 20.0% 21.7% 21.7% 23.3% 10.0% 21.7% 25.0% 13.8% 14.0% 25.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 11:50 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13:46 Mi/kg 13:46 Mi/kg 13:46 Mi/kg 13:46 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13:55 Mi/kg 13:55 Mi/kg 13:55 Mi/kg 13:55 Mi/kg 0 00 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprodus Gyprodus Other UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Vegetation	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 14.44% 100.0% 100.0% 10.90% 10.90% 10.90% 10.90% 10.90% 10.90% 0.00% 0.00%	20.4% 0.0% 0.0% 20.4% 20.4% 0.0% 0.0% 0.0% 35.0% 40.7% 48.2% 48.2% 48.2% 48.2% 48.2% 9.8%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 5.4% H 4.3% 5.4% H 4.1% 12.1% 4.7% 4.2% 0.0% 0.0%	7 35% 7 4 48 0.0% 0.0% 30.4% 4 48 0.0% 0.0% 36.8% 0.0% 17.5% 13.9% 0.0% 21.1% 10.5% 0.0% 21.1% 10.5% 0.0%	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.6% 1.10% N 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.9% 0.0% 0.0% 0.0% 0.9% 0.9% 0.0% 0.0	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 46.5% 46.5% 28.0% 90.0% 19.7% 46.5% 28.0% 90.0% 100.0% 28.0% 100.0%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 29.3% 25.0% 0.0% 0.0% 20.0% 20.0% 20.0% 21.7% 22.0% 10.0% 21.7% 23.8% 14.0% 25.0% 13.8% 14.0% 80.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13 84 Mi/kg 13 84 Mi/kg 13 84 Mi/kg 13 9 Mi/kg 13 9 Mi/kg 19 36 Mi/kg 0 00 Mi/kg 19 25 9 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gypred UTDI (Nov 15) report Concept of Design (March 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles	0.37% 0.18% 21.53% 2.84% 0.00% 0.00% 0.00% 0.00% 100.0% 100.0% 3.93% 10.90% 0.18% 0.00% 0.18% 0.00% 0.18%	20.4% 0.0%	2.4% 2.4% 0.0% 0.0% 2.4% 2.4% 2.4% 0.0% 0.0% 5.4% H 4.1% 5.4% H 4.1% 5.4% 4.1% 5.4%	7 35% 7 4 48 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 14.9% 0 0 11.4% 0.0% 11.4% 0.0% 11.5% 0.0%	0.3% 0.2% 0.0% 0.0% 0.3% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.2% 0.0% 0.1% 0.1% 0.1% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.9% 0.9%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 90.0% 90.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 25.0% 0.0% 20.0% 10.0% 21.7% 23.3% 25.0% 10.0% 25.0% 10.0% 25.0% 0.0% 80.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 8.30 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 0.00 Mi/kg 13.50 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.55 Mi/kg 13.55 Mi/kg 19.36 Mi/kg 0.00 Mi/kg 19.36 Mi/kg 0.00 Mi/kg 19.36 Mi/kg 0.00 Mi/kg 19.36 Mi/kg 0.00 Mi/kg 19.36 Mi/kg 0.00 Mi/kg 19.36 Mi/kg 0.00 Mi
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprot UTDI (Nov 15) report UTDI (Nov 15) report UTDI (Nov 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Ukgetation Other combustibles	0.37% 0.18% 0.18% 0.18% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00%	20.4% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 0.0% 35.0% 40.0% 35.3% 70.7% 48.2% 48.8% 9.8% 0.0% 5.0% 5.3%	2.4% 2.4% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 4.3% 4.3% 4.3% 4.3% 4.3% 4.3% 4.1% 4.1% 4.1% 6.0% 1.3% 6.0%	7 35% 7 4 48 4 48 0.0% 0.0% 30.4% 4 4% 0.0% 0.0% 17 5% 14.9% 0.0% 17 5% 14.9% 0.0% 21.1% 10.5% 21.1% 0.0% 0	0.3% 0.2% 0.0% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.6% 1.1% N 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.0% 0.9% 0.0% 0.0% 0.0% 0.9% 0.0% 0.9% 0.0% 0.0	1.0% 46.5% 100.0% 2.8% 46.5% 46.5% 100.0% 28.0% 90.0% 100.0% 19.7% 100.0% 19.7% 100.0% 100.0%	0.0% 80.0% 25.0% 0.0% 0.0% 29.3% 29.3% 20.0% 0.0% 0.0% 20.0%	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 1 50 Mi/kg 1 50 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 1 50 Mi/kg 1 3 46 Mi/kg 0 00 Mi/kg 0 0 Mi/k
Metal Fines Wood Combustibles Hazardous Gyprock Other UTDI (Nov 15) report Corcept of Design (March 13) resort Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles Metal Fines	0.37% 0.18% 0.18% 0.48% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.08% 0.08% 0.08% 0.00% 0.18%	20.4% 0.0% 33.0% 33.0% 20.4% 0.0% 0.0% 0.0% 35.0% 40.0% 5.3% 70.7% 48.2% 48.2% 48.8% 0.0% 5.3% 70.7% 20.4% 0.0%	2.4% 2.4% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 4.2% 0.0% 4.2% 0.0% 4.3% 5.4% H 4.1% 12.1% 4.1% 12.1% 4.1% 12.1% 4.1% 1.3% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0	7 35% 7 4 48 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 0.0% 17.5% 0.0% 17.5% 0.0% 11.4% 0.0% 0.5% 0.0%	0.3% 0.2% 0.0% 0.0% 0.0% 0.2% 0.0% 0.0% 0.0	0.0% 0.2% 0.0% 0.1% 0.1% 0.1% 0.0% 0.0% 0.0% 0.0	0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 2.8% 46.5% 100.0% 2.8% 2.8% 3.0% 90.0% 4.1% 2.6% 7.5% 7.5% 7.5% 7.5% 7.5% 7.5% 7.5% 7.5	0.0% 80.0% 25.0% 0.0% 0.0% 0.0% 25.0% 0.0% 20.0% 21.0% 21.7% 23.0% 25.0% 25.0% 25.0% 25.0% 25.0% 0.0% 80.0% 0.0% 80.0% 25.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 11.50 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 0 00 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Hazardous Gypreduc UTDI (Nov 15) report Concept of Design (March 15) report Concept of Design (March 15) report Plack Waste Paper/card Plastic film Dense plastic. Textiles Glass Vegetation Other combustibles Metal Fines Wood	0.37% 0.18% 0.18% 0.18% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.8%	20.4% 0.0% 0.0% 33.0% 20.4% 20.4% 0.0% 0.0% 0.0% 35.0%	2.3% 2.4% 0.0% 0.0% 2.4% 2.4% 0.0% 0.0% 0.0% 5.4% F H 4.3% 12.1% 4.3% 12.1% 4.7% 4.7% 4.7% 4.7% 4.7% 4.7% 4.7% 4.7	7 35% 7 4 48 4 48 0.0% 0.0% 30.4% 4 4% 0.0% 0.0% 17 5% 16 36 8% 0.0% 17 5% 10 5% 0.0% 0.0% 10.0% 0.0%	0.3% 0.2% 0.2% 0.0% 0.1% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.2% 0.2% 0.0% 0.1% 0.1% 0.1% 0.7% 0.0% 0.0% 0.0% 0.0% 0.1% 0.1% 0.0% 0.0	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0%	0.0% 80.0% 25.0% 0.0% 0.0% 25.0% 25.0% 0.0% 0.0% 20.0% 20.0% 20.0% 20.0% 20.0% 20.0% 20.0% 20.0% 20.0% 25.0% 25.0% 25.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 0 00 Mi/kg
Metal Fines Wood Combustibles Non-combustibles Haardous Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Treatiles Glass Vegetation Other combustibles Metal Fines Wood	0.37% 0.18% 0.18% 0.15% 0.4% 0.00% 0	20.4% 20.4% 0.0% 0.0% 20.4% 0.0	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 5.4% 1.3% 4.3% 4.3% 4.3% 4.3% 4.3% 4.3% 12.3% 4.2% 4.2% 0.0% 0.0% 0.0% 0.0% 0.0% 4.2% 4.2% 4.2% 4.2% 4.2% 4.2% 4.2% 4.2	2 35% 2 4.4% 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 14.9% 0 0 31.4% 0.0% 21.1% 10.5% 10.5% 0.0% 21.1% 0.0% 0.	0.3% 0.2% 0.2% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0% 100.0% 100.0% 100.0% 19.7% 100.0% 19.7% 4.1% 2.6% 2.6% 7.5% 100.0% 1.0% 46.5%	0.0% 80.0% 25.0% 0.0% 29.3% 25.0% 0.0% 0.0% 0.0% 20.0% 21.7% 21.7% 21.7% 21.7% 21.7% 21.7% 22.0% 25.0% 25.0% 0.0% 0.0% 0.0% 0.0% 0.0% 25.0% 20.0% 25.0% 20.0% 25.0	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 0 00 Mi/kg 0 0
Metal Fines Wood Combustibles Non-combustibles Harardous Gyprot UTDI (Nov 15) report Concept of Design (March 15) report Concept of Design (March 15) report Rock Waste Paper/card Plastic film Denne plastic Textiles Glass Vegetation Other combustibles Metal Fines Wood Combustibles Non-combustibles	0.37% 0.18% 2.84% 0.00% 0.00% 0.00% 14.44% 100.0% 100.0% 10.90% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00%	20.4% 0.0% 0.0% 20.4% 20.4% 20.4% 0.0% 0.0% 0.0% 35.0% 35.0% 35.0% 40.0% 55.3% 70.7% 48.2%48.2% 49.2% 49.2% 49.2%49.2% 49.2% 49.2% 49.2%49.2% 49.2% 49.2%40.2% 40.2% 40.2%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 5.4% 5.4% 7.4% 7.4% 12.1% 4.3% 12.1% 4.7% 4.2% 0.0% 0.0% 0.0% 0.0% 0.0%	7 35% 7 4 48 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 175% 0.0	0.3% 0.2% 0.2% 0.0% 0.3% 0.3% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.2% 0.0% 0.1% 0.1% 0.1% 0.1% 0.0% 0.0% 0.1% 0.0% 0.1% 0.1	0.0% 0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 1.0% 100.0% 100.0% 100.0% 2.8% 46.5% 100.0	0.0% 80.0% 25.0% 0.0% 0.0% 0.0% 29.3% 25.0% 0.0% 20.0% 21.7% 21.7% 23.0% 21.7% 23.0% 25.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 0 00
Metal Fines Wood Combustibles Non-combustibles Haardous Gyprotu UTDI (Nov 15) report UTDI (Nov 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Usegetation Other combustibles Metal Fines Wood Combustibles Non-combustibles	0.37% 0.18% 0.18% 0.18% 0.00%	20.4% 20.4% 0.0% 0.0% 20.4% 20.4% 0.0% 0.0% 35.0% dC_1 C 35.3% dC_1 C 35.3% dC_1 0.0%	2.4% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 5.4% 4.1% 5.4% 4.1% 12.1% 4.1% 1.2% 4.1% 0.0% 5.4% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0	7 35% 7 36% 7 4 48% 0.0% 0.0% 0.0% 0.0% 0.0% 17 5% 14.9% 0.0% 11 5% 0.0% 21.1% 0.0% 0.0% 21.1% 0.0% 0.	0.3% 0.2% 0.2% 0.0% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.2% 0.0% 0.1% 0.1% 0.1% 0.1% 0.0% 0.0% 0.0	0.0% 0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0%	0.0% 80.0% 25.0% 0.0% 25.0% 25.0% 25.0% 25.0% 25.0% 20.0% 20.0% 21.7% 21.7% 21.7% 21.7% 21.7% 21.0% 25	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13.84 Mi/kg 13.84 Mi/kg 13.84 Mi/kg 13.84 Mi/kg 13.95 Mi/kg 0 00 Mi/kg 0
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprock UTDI (Nov 15) report Concept of Design (March 15) report Concept of Design (March 15) report Flock Waste Paper/card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles Metal Fines Wood Combustibles Non-combustibles Hazardous	0.37% 0.18% 0.18% 0.18% 0.18% 0.46% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00%	20.4% 0.0% 0.0% 20.4% 20.4% 0.0% 0.0% 0.0% 0.0% 35.0% 40.1% 25.3% 70.7% 48.2% 48.8% 0.0% 9.8% 20.4% 0.0% 0.0% 0.0% 0.0%	2.4% 2.4% 0.0% 0.0% 4.2% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 5.4% H H H H H H H H H H	7 35% 7 4 48 4 48 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 14.9% 0.0% 13.4% 0.0% 13.4% 0.0% 0.0% 13.4% 0.0%	0.3% 0.2% 0.2% 0.0% 0.3% 0.3% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.2% 0.0% 0.1% 0.1% 0.1% 0.1% 0.1% 0.1% 0.1	0.0% 0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 1.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 1.0%	0.0% 80.0% 80.0% 80.0% 25.0% 0.0% 0.0% 0.0% 29.3% 29.3% 20.0% 20.0% 20.0% 20.0% 20.0% 20.0% 25.0% 13.8% 14.0% 25.0% 25.0% 25.0% 25.0% 0.0% 0.0% 25.0% 0.0% 0.0% 25.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0 00 Mi/kg 1 92 Mi/kg 3 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 13.46 Mi/kg 0 00 Mi/kg 0 0
Metal Final Gombustibles Non-combustibles Non-combustibles Hazardous Gypreduc UTDI (Nov 15) report Concept of Design (March 15) report Concept of Design (March 15) report Plack Waste Paper/card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles Non-combustibles Non-combustibles Nazardous Gyproduc	0.37% 0.18% 0.18% 0.18% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00%	20.4% 20.4% 0.0% 0.0% 20.4% 0.0% 0.0% 33.0% 32.0% 35.0% dt1,	2.3% 2.4% 0.0% 0.0% 2.4% 0.0% 0.0% 0.0% 5.4% 5.4% 12.1% 4.3% 12.1% 4.3% 12.1% 4.7% 4.3% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	7 25% 7 25% 7 4 43% 0.0% 0.0% 30.4% 4 43% 0.0% 17 5% 13 5% 10 5% 0.0% 10 5% 10 5	0.3% 0.2% 0.2% 0.0% 0.1% 0.1% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.2% 0.0% 0.0% 0.1% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.0% 0.9% 0.0% 0.0% 0.0% 0.0% 0.0%	1.0% 46.5% 100.0% 2.8% 46.5% 100.0% 28.0% 90.0% 100.0% 28.0% 90.0% 100.0% 28.0% 2.6% 2.6% 2.6% 2.6% 2.6% 2.6% 100.0% 100.0% 100.0% 2.8% 46.5%	0.0% 80.0% 25.0% 0.0% 25.0% 25.0% 25.0% 25.0% 20.0% 21.7% 24.7% 25.0% 21.7% 25.0% 21.7% 25.0% 20.0% 20	0 00 Mi/kg 1 92 Mi/kg 8 30 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 1 150 Mi/kg 0 00 Mi/kg 0 00 Mi/kg 1 3 46 Mi/kg 1 3 46 Mi/kg 1 3 46 Mi/kg 1 3 46 Mi/kg 0 00 Mi/k
Metal Fines Wood Combustibles Non-combustibles Hazardous Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) resort Flock Waste Paper/card Plastic film Dense plastic Flock Waste Raper/card Plastic film Dense plastic Glass Uegetation Other combustibles More downstribles Non-combustibles Non-combustibles Hazardous Gyprock Other Other	0.37% 0.18% 0.18% 0.18% 0.00%	20.4% 0.0% 0.0% 33.0% 20.4% 0.0% 0.0% 0.0% 0.0% 35.0% 40.1 55.3% 40.0% 20.4% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	2.4% 2.4% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 5.4% H H 4.1% 5.4% H H 4.1% 12.1% 4.7% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0	7 35% 7 4 48 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 17.5% 14.4% 0.0% 1.1.5% 14.4% 0.0%	0.3% 0.2% 0.0% 0.0% 0.0% 0.2% 0.0% 0.0% 0.0	0.0% 0.0% 0.2% 0.0% 0.3% 0.0% 0.3% 0.0% 0.3% 0.0% 0.3% 0.0% 0.1% 0.1% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.9% 0.9% 0.0% 0.0% 0.9% 0.9% 0.0% 0.0	1.0% 46.5% 100.0% 100.0% 2.8% 46.5% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 100.0% 1.0% 4.1% 1.0% 46.5% 100.0% 2.8% 90.0% 34.2%	0.0% 80.0% 25.0% 0.0% 29.3% 29.3% 25.0% 0.0% 20.0% 20.0% 21.7% 21.7% 21.7% 21.7% 21.7% 21.7% 21.7% 21.7% 25.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0 00 M/kg 1 92 M/kg 8 30 M/kg 0 00 M/kg 0 00 M/kg 0 00 M/kg 0 00 M/kg 11.50 M/kg 13.46 M/kg 13.46 M/kg 13.46 M/kg 13.46 M/kg 0 00 M/kg 19.36 M/kg 0 00 M/kg 1.50 M/kg 1.50 M/kg 1.50 M/kg 1.50 M/kg 1.50 M/kg 1.50 M/kg 0.00



Paper Mag C N C N C N C N C N C N C N C N C N Diss M Dece plate 0.00 0.20 <td< th=""><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th><th></th></td<>											
Periora Field 33.6 43.8 34.0 50.6 0.75 43.8 20.75 43.8 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 39.9 40.75 10.9 40.75 <t< td=""><td>Paper Pulp</td><td></td><td>C</td><td>H</td><td>0</td><td>N</td><td>s</td><td>D</td><td>Ash</td><td>H,O</td><td>NCV</td></t<>	Paper Pulp		C	H	0	N	s	D	Ash	H,O	NCV
Instriction 12.06 2.25 12.15 4.05 4.05 4.05 2.05 12.85 13.85 <t< td=""><td>Paper/rard</td><td>78.40%</td><td>35 3%</td><td>4.1%</td><td>31.4%</td><td>0.0%</td><td>0.2%</td><td>0.0%</td><td>4.1%</td><td>25.0%</td><td>12.15 MI/kg</td></t<>	Paper/rard	78.40%	35 3%	4.1%	31.4%	0.0%	0.2%	0.0%	4.1%	25.0%	12.15 MI/kg
Open pisses	Plastic film	21.60%	70.7%	12.1%	0.0%	0.0%	0.0%	0.9%	2.6%	13.8%	35.59 MI/kg
Tenten 0.000 0.85 0.15 0.00	Dense plastic	0.00%	48.2%	4.7%	21.1%	0.0%	0.0%	9.5%	2.6%	14.0%	18.58 MJ/ke
Gene DDN DDN <thdn< th=""> <thdn< th=""></thdn<></thdn<>	Textiles	0.00%	48.8%	4.1%	10.5%	4.3%	0.0%	0.0%	7.5%	25.0%	19.36 MI/ke
Wegative 0.005 0.316 0.36 0.05	Glass	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Other anisolation	Vegetation	0.00%	9.8%	1.3%	7.5%	0.3%	0.0%	0.0%	1.0%	80.0%	1.92 MJ/kg
Note ODE ODE <td>Other combustibles</td> <td>0.00%</td> <td>20.4%</td> <td>2.4%</td> <td>4.4%</td> <td>0.2%</td> <td>0.2%</td> <td>0.9%</td> <td>46.5%</td> <td>25.0%</td> <td>8.30 MJ/kg</td>	Other combustibles	0.00%	20.4%	2.4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
ime 0.006 0	Metal	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MI/ke
Wine and Dots Diff.	Fines	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 Mi/kg
Generation 0.000 2.00 2.00 0.000	Wood	0.00%	33.0%	4.2%	30.4%	0.1%	0.1%	0.0%	2.8%	29.3%	11.50 MI/ke
Ner-sensentistis 0.000	Combustibles	0.00%	20.4%	2.4%	4 4%	0.2%	0.2%	0.9%	46.5%	25.0%	8 30 Mi/kg
Number barehols Opp	Non-combustibles	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MI/kg
Operator ODD OD	Harardous	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 Mi/kg
Differ 0.000 <t< td=""><td>Gynrock</td><td>0.00%</td><td>0.0%</td><td>0.0%</td><td>36 844</td><td>0.0%</td><td>15 7%</td><td>0.0%</td><td>28.0%</td><td>20.0%</td><td>-2.97 MI/kg</td></t<>	Gynrock	0.00%	0.0%	0.0%	36 844	0.0%	15 7%	0.0%	28.0%	20.0%	-2.97 MI/kg
UDD UDD <thudd< th=""> <thudd< th=""> <thudd< th=""></thudd<></thudd<></thudd<>	Other	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	90.0%	10.0%	-0.24 MI/kg
Out Description Description <thdescription< th=""> <thdes< td=""><td>Utrai las</td><td>100.0076</td><td>0.076</td><td>0.036</td><td>0.076</td><td>0.0%</td><td>0.076</td><td>0,074</td><td>30.076</td><td>20.074</td><td>-Dian Wu/ng</td></thdes<></thdescription<>	Utrai las	100.0076	0.076	0.036	0.076	0.0%	0.076	0,074	30.076	20.074	-Dian Wu/ng
Spenner Monet 31 areyof 10005 1000 1000 00000 0000 0000	UTDI (Nov 15) report	100.0%	42,9%	5.8%	24.6%	0.0%	0.1%	0.2%	3.7%	22.6%	17.21 MJ/kg
Enservice C N O N S O No Processor Preperior 62.000 65.75 2.15 0.05 0.055 0.055 1.05 1.05 0.055 <t< td=""><td>Concept of Design (March 15) report</td><td>100.0%</td><td>Manager.</td><td>5,1%</td><td>24.6%</td><td>0.4%</td><td>0.1%</td><td>0.4%</td><td>000</td><td>25.0%</td><td>13.22 MJ/kg</td></t<>	Concept of Design (March 15) report	100.0%	Manager.	5,1%	24.6%	0.4%	0.1%	0.4%	000	25.0%	13.22 MJ/kg
Bites Residual C B O N S O Auh Hyo 12 by V/V/V/V/V/V/V/V/V/V/V/V/V/V/V/V/V/V/V/											
Description Paper/ant 12:00 13:10	Glass Residual	1 1	r.		0	N	5	n	Ash	8.0	NCV
Privatic Fin 1.28 0.28 1.28 0.08 0.08 0.08 0.08 0.08 0.08 1.28 1.18 1.28 1.18 1.28 1.18 1.28	Baner/card	63.00%	20 210	4.152	21.49/	0.0%	0.7%	0.0%	4.1%	76,787	10 15 AM/Am
Drose plateix 2.0% 2.1% 2.0% 0.0%	Paper/caru Diactia film	2 909/	22.3/0	4.170	0.0%	0.0%	0.2%	0.0%	7.1%	13.9%	25 50 MU/kg
Jobis Textistic Jobis	Plastic mm	3.80%	70,7%	12.1%	0,0%	0.0%	0.0%	0.9%	2.0%	13.6%	30.09 Mu/kg
facture 0.00x d.B. 0.00x 0.00x <t< td=""><td>Dense plastic</td><td>34.20%</td><td>48.2%</td><td>4./%</td><td>21.1%</td><td>0.0%</td><td>0.0%</td><td>9.5%</td><td>2,6%</td><td>14.0%</td><td>18.58 MJ/kg</td></t<>	Dense plastic	34.20%	48.2%	4./%	21.1%	0.0%	0.0%	9.5%	2,6%	14.0%	18.58 MJ/kg
Open A OPA OPA<	Textiles	0.00%	48.8%	4,1%	10.5%	4.3%	0.0%	0.0%	7.5%	25.0%	19.36 MU/kg
Vegetation 0.001 2.8 Ph 1.95 0.75	Glass	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Other embaulties 0.004 2.4% 4.8% 0.07% 0.0% <td>Vegetation</td> <td>0.00%</td> <td>9.8%</td> <td>1.3%</td> <td>7.5%</td> <td>0.3%</td> <td>0.0%</td> <td>0.0%</td> <td>1.0%</td> <td>80.0%</td> <td>1.92 MJ/kg</td>	Vegetation	0.00%	9.8%	1.3%	7.5%	0.3%	0.0%	0.0%	1.0%	80.0%	1.92 MJ/kg
Metal 0.00% 0.0% 0.0% 0.0% 0.0% 0.0% 0.00	Other combustibles	0.00%	20,4%	2,4%	4.4%	0.2%	0,2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Image 0.00% 0.0% 0.0% 0.0% 0.0% 0.00% 0.00% 0.00 0.00 0.00% 0.0%	Metal	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Wead 0.00% 31.0% 4.2% 0.9.4% 0.1% 0.2% 0.7%	Fines	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Combustifies 0.007 2.01% 2.9% 4.43% 0.23% 0.2% 4.5% 4.50% 4.5% 4.5% 4.5% 4.5% 4.5% 4.5% 4.5% 0.05% 0.	Wood	0.00%	33.0%	4.2%	30.4%	0.1%	0.1%	0.0%	2.8%	29.3%	11.50 MJ/kg
Nen-combustible 0.0%	Combustibles	0.00%	20.4%	2.4%	4,4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Haardees 0.0%	Non-combustibles	0.00%	0.0%	0,0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Opposit ODS ODS <thods< th=""> <thods< <="" td=""><td>Hazardous</td><td>0.00%</td><td>0.0%</td><td>0,0%</td><td>0.0%</td><td>0.0%</td><td>0.0%</td><td>0,0%</td><td>100.0%</td><td>0,0%</td><td>0.00 MI/ke</td></thods<></thods<>	Hazardous	0.00%	0.0%	0,0%	0.0%	0.0%	0.0%	0,0%	100.0%	0,0%	0.00 MI/ke
Option Option<	Gunrack	0.00%	0.0%	0.0%	36.9%	0.0%	15 380	0.0%	28.08/	20.0%	2 87 Mil/kg
Other Other <th< td=""><td>Gyprock</td><td>0.00%</td><td>0.0%</td><td>0.0%</td><td>0,000</td><td>0.0%</td><td>0.00/</td><td>0.0%</td><td>90.0%</td><td>10.0%</td><td>-0.24 MU/rg</td></th<>	Gyprock	0.00%	0.0%	0.0%	0,000	0.0%	0.00/	0.0%	90.0%	10.0%	-0.24 MU/rg
VID UTO VID VID <td>Other</td> <td>0.00%</td> <td>0.0%</td> <td>U.07%</td> <td>0,0%</td> <td>0.0%</td> <td>0.0%</td> <td>U.U%</td> <td>90.0%</td> <td>10.0%</td> <td>-0.24 MU/kg</td>	Other	0.00%	0.0%	U.07%	0,0%	0.0%	0.0%	U.U%	90.0%	10.0%	-0.24 MU/kg
Concert of Design (March 15) report 10005 8.0.75 5.766 0.4% 0.4% 0.3% 2.4% 1.998 15.11 MU GD Residual C H O N S D A MO NO Properioral 2.506 4.325 4.11% 0.01% 0.02% 0.01% 4.15% 2.506 1.325.5 (MU)	UTDI (Nov 15) report	100.0%	41.0%	4.5%	26.7%	0.0%	0.1%	3.3%	3.5%	20.8%	15.24 MJ/kg
BC Residual C H O N S G Ath H ₀ O NOC Bised Film 2.007 55.3% 4.1% 21.5 M/0 0.07% 0.2% 0.05% 4.1% 25.0% 1.25.0% <td>Concept of Design (March 15) report</td> <td>100.0%</td> <td>40,3%</td> <td>5.6%</td> <td>22.9%</td> <td>0.4%</td> <td>0.1%</td> <td>2.4%</td> <td>1.00</td> <td>19.6%</td> <td>16.13 MJ/Kg</td>	Concept of Design (March 15) report	100.0%	40,3%	5.6%	22.9%	0.4%	0.1%	2.4%	1.00	19.6%	16.13 MJ/Kg
C H O N S C Ath H ₀ O NOV. Plants film 2.50% 0.57% 1.21% 0.0% 0.0% 0.0% 2.5% 1.21% 0.0% 0.0% 0.0% 0.0% 0.0% 1.21% 0.0% <td>1</td> <td></td> <td>(c) (c) (c) (c) (c) (c) (c) (c) (c) (c)</td> <td></td> <td></td> <td></td> <td>2000</td> <td></td> <td></td> <td>-</td> <td></td>	1		(c)				2000			-	
Description 20:00 33:3% 0.1% 0.2% 0.0% 4.1% 7.0% 2.3.5% Description 2.50% 3.5.3% 4.1% 1.1.5% 0.0% 0.0% 4.1% 2.50% 3.2.3.5% 0.0% 0.0% 4.1% 1.1.5% 0.0% 0.0% 4.1% 1.1.5% 0.0%	60 Peridual				0	N	c	a	Ach	HO	NOV
Paper (ard 30.0% 33.3% 41.% 31.4% 0.0% 0.0% 0.0% 0.0% 2.6% 12.1% 55.99 M/J Derive plastic 2.35% 42.7% 42.7% 21.1% 0.0% 0.0% 0.0% 2.6% 1.40% 1.35% 43.7% Derive plastic 2.35% 44.2% 4.7% 21.1% 0.0% 0.0% 0.0% 2.6% 1.40% 1.35% 43.7% 0.0% 0.0% 0.0% 2.6% 1.40% 0.0% <td0< td=""><td>SO Residual</td><td></td><td>L</td><td>17</td><td>0</td><td>IN</td><td>3</td><td>u</td><td>Asn</td><td>1120</td><td>NCV</td></td0<>	SO Residual		L	17	0	IN	3	u	Asn	1120	NCV
Pieste film 2.50% 4.27% 4.7% 2.11% 0.0% 0.0% 0.9% 2.6% 1.38% 3.559 4.43% 1.00% 0.0% 0.0% 0.0% 2.6% 1.38% 3.559 4.16% 1.00% 0.0% <th0.0%< th=""> <th0.0%<< td=""><td>Paper/card</td><td>30.00%</td><td>35.3%</td><td>4.1%</td><td>31.4%</td><td>0.0%</td><td>0.2%</td><td>0.0%</td><td>4.1%</td><td>25.0%</td><td>12.15 MJ/kg</td></th0.0%<<></th0.0%<>	Paper/card	30.00%	35.3%	4.1%	31.4%	0.0%	0.2%	0.0%	4.1%	25.0%	12.15 MJ/kg
Dense plastic 2.5% 44.7% 21.1% 0.0% 3.9% 2.9% 1.4.0% 1.38.8 1.39.5% 0.0% 0.0% 7.5% 0.3% 0.0% <th0.0%< th=""> 0.0%</th0.0%<>	Plastic film	2.50%	70,7%	12.1%	0.0%	0.0%	0.0%	0.9%	2.6%	13.8%	35.59 MJ/kg
Teatline 0.00% 48.8% 4.1% 10.5% 4.3% 0.0%	Dense plastic	2.50%	48,2%	4.7%	21.1%	0.0%	0.0%	9.5%	2.6%	14.0%	18.58 MJ/kg
Gless 4.00% 0.0% <th0.0%< th=""> 0.0% 0.0% <t< td=""><td>Textiles</td><td>0.00%</td><td>48.8%</td><td>4,1%</td><td>10.5%</td><td>4.3%</td><td>0.0%</td><td>0.0%</td><td>7.5%</td><td>25.0%</td><td>19.36 MJ/kg</td></t<></th0.0%<>	Textiles	0.00%	48.8%	4,1%	10.5%	4.3%	0.0%	0.0%	7.5%	25.0%	19.36 MJ/kg
Vegration 93:80 9.38 1.3% 7.5% 0.3% 0.0% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.0%	Glass	4.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Other combusible 0.00% 2.4% 4.4% 0.2% 0.2% 0.0% 10.0% 0.0% <td>Vegetation</td> <td>35.00%</td> <td>9.8%</td> <td>1.3%</td> <td>7.5%</td> <td>0.3%</td> <td>0.0%</td> <td>0.0%</td> <td>1.0%</td> <td>80.0%</td> <td>1.92 MJ/kg</td>	Vegetation	35.00%	9.8%	1.3%	7.5%	0.3%	0.0%	0.0%	1.0%	80.0%	1.92 MJ/kg
Metal 5.00% 0.0% <	Other combustibles	0.00%	20.4%	2.4%	4.4%	0.2%	0.2%	0.9%	46.5%	25.0%	8.30 MJ/kg
Fines 0.00% 0.0% <	Metal	5.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Word 0.09% 23.0% 4.2% 30.4% 0.1% 0.1% 0.0% 2.8% 23.8% 11.50 M/J Non-combustibles 20.09% 0.05% 0.07%	Fines	0.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MJ/kg
Combustibles 20.0% 20.4% 24.% 24.% 0.2% 0.2% 0.9% 46.5% 25.0% B.30 M/J Non-combustibles 21.0% 0.0%	Wood	0.00%	33.0%	4.7%	30.4%	D 1%	0.1%	0.0%	2.8%	29.3%	11.50 MI/kg
Non-combattles 20.0% 0.0%	Comhustibles	0.00%	20.4%	2.4%	1 194	0.7%	0.2%	0.9%	46.5%	25.0%	8 30 MI/kg
Non-standous 1.00% 0.0%	Non-combustibles	21.00%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 MI/kg
Initiations DOIN	Hon-combusciones	0.00%	0.0%	0,0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 10/16
Opyrock OUM OUM <thoum< th=""> <thoum< <="" td=""><td></td><td>12-11/28</td><td>0.076</td><td>0.076</td><td>0,0%</td><td>0.0%</td><td>0.0%</td><td>0.0%</td><td>100.0%</td><td>0.0%</td><td>0.00 WU/Kg</td></thoum<></thoum<>		12-11/28	0.076	0.076	0,0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.00 WU/Kg
Uther 0.00% 0.0% <th0.0%< th=""> 0.0% 0.0% <!--</td--><td>Hazardous</td><td>0.000/</td><td>0.00/</td><td>0.00/</td><td>30.8%</td><td>0.0%</td><td>17,220</td><td>0.0%</td><td>28.0%</td><td>20.0%</td><td>-2.87 WU/KE</td></th0.0%<>	Hazardous	0.000/	0.00/	0.00/	30.8%	0.0%	17,220	0.0%	28.0%	20.0%	-2.87 WU/KE
UTOI (Nov 15) report 100.0% 17.0% 2.1% 12.6% 0.1% 0.1% 0.3% 31.7% 36.7% 5.67 MJ/ Concept of Design (Morch 15) report 100.0% 18.5% 2.5% 14.3% 0.5% 0.2% 0.4% 33.0% 31.7% 56.37 M F.41 MJ/ AWT Residual C H O N S C Ash H_0O NV Paper/ard 21.05% 35.3% 41.1% 31.4% 0.0% 0.0% 41.1% 12.15 MJ/ NV Paper/ard 21.05% 48.2% 4.7% 12.1% 0.0% 0.0% 0.0% 2.5% 13.6% 13.5% 10.5% 10.5%	Gyprock	0.00%	0.0%	0.0%	0.00	0.004	0.000	0.00/	00.011	10.004	0.74 14/8-
Concept of Design (Morch 15) report 100.0% 18.5% 2.5% 14.3% 0.5% 0.2% 0.4% 32.0% 31.7% 5.31 M/r AMYT Residual C H O N S C Ash H/O NCV Paper/card 21.05% 35.3% 4.1% 31.4% 0.0% 0.0% 0.0% 4.1% 25.0% 12.15 M/r Dense plastic film 20.05% 48.2% 4.7% 21.1% 0.0% 0.0% 0.0% 2.5% 13.8% 35.59 M/r Genes plastic film 20.5% 48.2% 4.7% 21.1% 0.0%	Gyprock Other	0.00%	0.0% 0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	90.0%	10.0%	-0.24 MJ/kg
AWT Residual C H O N S C Ash H;O NCV Paper(rard person plastic 21.05%, d82.5% 21.05%, d82.5% 41.1% d1.05%, d82.5% 31.4%, d1.05% 0.0% d0.0% 0.0% d0.0% 0.0%, d0.0% 0.0%, d0.0% 25.5% d1.05%, d1.05%	Gyprock Other JTDI (Nov 15) report	0.00% 0.00% 100.0%	0.0% 0.0% 17.0%	0.0% 0.0% 2.1%	0.0%	0.0% 0.1%	0.0% 0.1%	0.0%	90.0% 31.7%	10.0% 36.2%	-0.24 MJ/kg 5.67 MJ/kg
AVT Residual C H O N S C Ash H ₂ O NCV Paper/Lard 21.05% 85.3% 41.5% 31.4% 0.07% 0.2% 0.0% 4.1% 25.0% 121.18.M/V Plastic fill 20.05% 48.2% 4.7% 21.1% 0.0% 0.0% 0.9% 2.5% 13.8% 135.59 M/V Textiles 10.5% 48.2% 4.7% 21.1% 0.0% 0.0% 0.0% 1.0% 135.8 M/V 135.8 M/V Vegtation 3.16% 9.8% 1.1% 10.5% 0.0% 0.0% 1.0% 1.00% 0.0% 1.00% 0.0% 1.00% 1.00 Model 1.25 M/V 1.20 M/V 1.26 M/V 1.25 M/V 1.00 M/V	Gyprock Other UTDI (Nov 15) report Ioncept of Design (March 15) report	0.00% 0.00% 100.0% 100.0%	0.0% 0.0% 17.0% 18.5%	0.0% 0.0% 2.1% 2.5%	0.0% 12.6% 14.3%	0.0% 0.1% 0.5%	0.0% 0.1% 0.2%	0.0% 0.3% 0.4%	90.0% 31.7% 32.0%	10.0% 36.2% 31.7%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg
Paper/card 21.05% 35.3% 4.1% 31.4% 0.0% 0.2% 0.0% 4.1% 25.0% 12.15 M// Plastic film 20.00% 70.7% 22.1% 0.0% 0.0% 0.0% 4.1% 13.8% 35.59 M// Dense plastic film 20.00% 62.3% 47.1% 21.1% 0.0% 0.0% 0.0% 2.6% 13.8% 35.59 M// Textiles 10.53% 48.8% 4.1% 10.5% 4.3% 0.0% <td>Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) report</td> <td>0.00% 0.00% 100.0% 100.0%</td> <td>0.0% 0.0% 17.0% 18.5%</td> <td>0.0% 0.0% 2.1% 2.5%</td> <td>0.0% 12.6% 14.3%</td> <td>0.0% 0.1% 0.5%</td> <td>0.0% 0.1% 0.2%</td> <td>0.0% 0.3% 0.4%</td> <td>90.0% 31.7% 32.0%</td> <td>10.0% 36.2% 31.7%</td> <td>-0.24 MJ/kg 5.67 MJ/kg 6,31 MJ/kg</td>	Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) report	0.00% 0.00% 100.0% 100.0%	0.0% 0.0% 17.0% 18.5%	0.0% 0.0% 2.1% 2.5%	0.0% 12.6% 14.3%	0.0% 0.1% 0.5%	0.0% 0.1% 0.2%	0.0% 0.3% 0.4%	90.0% 31.7% 32.0%	10.0% 36.2% 31.7%	-0.24 MJ/kg 5.67 MJ/kg 6,31 MJ/kg
Particle 21.00% 23.3% 4.1% 0.0% 0.0% 0.0% 1.1% 23.0% 1.1.1% 1.0.0% 0.0% 0.0% 2.0% 1.1.1% 1.0.0% 0.0% 0.0% 0.0% 2.0% 1.1.8% 1.0.0% 0	Hazaroous Gyprock Other UtTDI (Nov.15) report Concept of Design (March 15) report	0.00% 0.00% 100.0% 100.0%	0.0% 0.0% 17.0% 18.5%	0.0% 0.0% 2.1% 2.5%	0.0% 12.6% 14.3%	0.0% 0.1% 0.5%	0.0% 0.1% 0.2%	0.0% 0.3% 0.4%	90.0% 31.7% 32.0%	10.0% 36.2% 31.7%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg
Dense plastic COV/R 22.1.8 0.078 0.078 0.078 0.078 0.078 0.078 0.078 0.078 0.078 0.078 0.078 2.578 11.858 11.858 11.858 11.858 11.858 11.858 11.858 11.858 11.858 11.858 10.076 0.	Hazaroous Gyprock Other UTDI (Nov 15) report Concept of Design (March 15) report AWT Residual	0.00% 0.00% 100.0% 100.0%	0.0% 0.0% 17.0% 18.5% C	0.0% 0.0% 2.1% 2.5% H	0.0% 12.6% 14.3%	0.0% 0.1% 0.5%	0.0% 0.1% 0.2% S	0.0% 0.3% 0.4%	90.0% 31.7% 32.0% Ash	10,0% 36,2% 31,7% H ₂ O	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV
Johnson Plastic 21.02% 48.2% 4.1% 0.0% 0.0% 0.0% 75% 2.6% 14.0% 16.558 Glass 0.00% 0.0% </td <td>Hazardous Gyprock UTDI (Nov 15) report Concept of Design (March 15) report NWT Residual Paper/Card Nexts cr</td> <td>0.00% 0.00% 100.0% 100.0% 21.05%</td> <td>0.0% 0.0% 17.0% 18.5% C 35.3%</td> <td>0.0% 0.0% 2.1% 2.5% H 4.1%</td> <td>0.0% 12.6% 14.3%</td> <td>0.0% 0.1% 0.5% N 0.0%</td> <td>0.0% 0.1% 0.2% \$ 0.2%</td> <td>0.0% 0.3% 0.4%</td> <td>90.0% 31.7% 32.0% Ash 4.1%</td> <td>10,0% 36,2% 31,7% H₂O 25,0%</td> <td>-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg</td>	Hazardous Gyprock UTDI (Nov 15) report Concept of Design (March 15) report NWT Residual Paper/Card Nexts cr	0.00% 0.00% 100.0% 100.0% 21.05%	0.0% 0.0% 17.0% 18.5% C 35.3%	0.0% 0.0% 2.1% 2.5% H 4.1%	0.0% 12.6% 14.3%	0.0% 0.1% 0.5% N 0.0%	0.0% 0.1% 0.2% \$ 0.2%	0.0% 0.3% 0.4%	90.0% 31.7% 32.0% Ash 4.1%	10,0% 36,2% 31,7% H ₂ O 25,0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg
rextites 10.53% 4.85% 0.05% 0.05% 0.05% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.000	Gyprock Gyprock UTDI (Nov 15) report Concept-of Design (Mirch 15) report AWT Residual Paper/Card Plastic film	0.00% 0.00% 100.0% 100.0% 20.0% 21.05% 20.00%	0.0% 0.0% 17.0% 18.5% C 35.3% 70.7%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1%	0.0% 12.6% 14.3% 0 31.4% 0.0%	0.0% 0.1% 0.5% N 0.0% 0.0%	0.0% 0.1% 0.2% \$ 0.2% 0.0%	0.0% 0.3% 0.4%	90.0% 31.7% 32.0% Ash 4.1% 2.5%	10.0% 36.2% 31.7% H ₂ O 25.0% 13.8%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg
Glass 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.07% 0.00% 0.07% 1.000 0.000 0.000 0.000 0.07% <th< td=""><td>Hazardous Gyprock Other UTDI (Nov 15) report Concept of Dasign (March 15) report AWT Residual Paper/card Plastic film Dense plastic</td><td>0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05%</td><td>0.0% 0.0% 17.0% 18.5% 2 35.3% 70.7% 48.2%</td><td>0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7%</td><td>0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1%</td><td>0.0% 0.1% 0.5% N 0.0% 0.0% 0.0%</td><td>0.0% 0.1% 0.2% 5 0.2% 0.0% 0.0%</td><td>0.0% 0.3% 0.4% 0.0% 0.9% 9.5%</td><td>90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6%</td><td>10.0% 36.2% 31.7% H20 25.0% 13.8% 14.0%</td><td>-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg 18.58 MJ/kg</td></th<>	Hazardous Gyprock Other UTDI (Nov 15) report Concept of Dasign (March 15) report AWT Residual Paper/card Plastic film Dense plastic	0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05%	0.0% 0.0% 17.0% 18.5% 2 35.3% 70.7% 48.2%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7%	0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1%	0.0% 0.1% 0.5% N 0.0% 0.0% 0.0%	0.0% 0.1% 0.2% 5 0.2% 0.0% 0.0%	0.0% 0.3% 0.4% 0.0% 0.9% 9.5%	90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6%	10.0% 36.2% 31.7% H20 25.0% 13.8% 14.0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg 18.58 MJ/kg
vegetation 3.16% 9.8% 1.3% 7.5% 0.3% 0.0% 0.1% 80.0% 1.19.4 Other combustibles 0.00% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.00% 0.0% <td>Gyprock Gyprock UTDI (Nov 15) report Concept of Design (Mirch 15) report AWT Residual AWT Residual Paper/card Plastic film Dense plastic Textiles</td> <td>0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05% 10.53%</td> <td>0.0% 0.0% 17.0% 18.5% C 35.3% 70.7% 48.2% 48.8%</td> <td>0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7% 4.1%</td> <td>0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1% 10.5%</td> <td>0.0% 0.1% 0.5% N 0.0% 0.0% 0.0% 4.3%</td> <td>0.0% 0.1% 0.2% 5 0.2% 0.0% 0.0% 0.0%</td> <td>0.0% 0.3% 0.4% 0.4% 0.0% 0.9% 9.5% 0.0%</td> <td>90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 7.5%</td> <td>10.0% 36.2% 31.7% H20 25.0% 13.8% 14.0% 25.0%</td> <td>-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg 18.58 MJ/kg 19.36 MJ/kg</td>	Gyprock Gyprock UTDI (Nov 15) report Concept of Design (Mirch 15) report AWT Residual AWT Residual Paper/card Plastic film Dense plastic Textiles	0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05% 10.53%	0.0% 0.0% 17.0% 18.5% C 35.3% 70.7% 48.2% 48.8%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7% 4.1%	0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1% 10.5%	0.0% 0.1% 0.5% N 0.0% 0.0% 0.0% 4.3%	0.0% 0.1% 0.2% 5 0.2% 0.0% 0.0% 0.0%	0.0% 0.3% 0.4% 0.4% 0.0% 0.9% 9.5% 0.0%	90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 7.5%	10.0% 36.2% 31.7% H20 25.0% 13.8% 14.0% 25.0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg 18.58 MJ/kg 19.36 MJ/kg
Other combustibles 0.00% 20.4% 2.4% 4.4% 0.2% 0.2% 0.9% 4.45 5% 25.0% 3.30 MJ/I Need 0.076	Autorous Gyprock Other UTDI (Nov 15) report Compt of Disign (Mirch 15) report AWT Residual Paper/Card Plastic film Dense plastic Textiles Glass	0.00% 0.00% 100.0% 20.0% 21.05% 20.00% 21.05% 10.53% 0.00%	0.0% 0.0% 17.0% 18.5% 0.5% 70.7% 48.2% 48.8% 0.0%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7% 4.1% 0.0%	0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1% 10.5% 0.0%	0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 4.3% 0.0%	0.0% 0.1% 0.2% 5 0.2% 0.0% 0.0% 0.0% 0.0%	0.0% 0.3% 0.4% 0.0% 0.0% 0.9% 9.5% 0.0% 0.0%	90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 7.5% 100.0%	10.0% 36.2% 31.7% H20 25.0% 13.8% 14.0% 25.0% 0.0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg 12.15 MJ/kg 35.59 MJ/kg 18.58 MJ/kg 19.36 MJ/kg 0.00 MJ/kg
Metal 0.0% <t< td=""><td>Hazirobis Gyprock Other UTDI (Nov 15) report Eonopt-of Design (Mirch 15) report AWT Residual Paper/card Plastic film Dense plastic Textiles Glass Vegetation</td><td>0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05% 20.00% 21.053% 0.00% 3.16%</td><td>0.0% 0.0% 17.0% 18.5% 0.35.3% 70.7% 48.2% 48.8% 0.0% 9.8%</td><td>0.0% 0.0% 2.1% 2.5% H 4.1% 4.7% 4.7% 4.7% 0.0% 1.3%</td><td>0.0% 12.6% 14.3% 0.0% 21.1% 0.0% 21.1% 10.5% 0.0% 7.5%</td><td>0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 4.3% 0.0% 0.3%</td><td>0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0%</td><td>0.0% 0.3% 0.4% 0.0% 0.9% 9.5% 0.0% 0.0%</td><td>90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 7.5% 100.0% 1.0%</td><td>10.0% 36.2% 31.7% H₂O 25.0% 13.8% 14.0% 25.0% 0.0% 80.0%</td><td>-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg 12.15 MJ/kg 13.59 MJ/kg 19.36 MJ/kg 0.00 MJ/kg 1.92 MJ/kg</td></t<>	Hazirobis Gyprock Other UTDI (Nov 15) report Eonopt-of Design (Mirch 15) report AWT Residual Paper/card Plastic film Dense plastic Textiles Glass Vegetation	0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05% 20.00% 21.053% 0.00% 3.16%	0.0% 0.0% 17.0% 18.5% 0.35.3% 70.7% 48.2% 48.8% 0.0% 9.8%	0.0% 0.0% 2.1% 2.5% H 4.1% 4.7% 4.7% 4.7% 0.0% 1.3%	0.0% 12.6% 14.3% 0.0% 21.1% 0.0% 21.1% 10.5% 0.0% 7.5%	0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 4.3% 0.0% 0.3%	0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.3% 0.4% 0.0% 0.9% 9.5% 0.0% 0.0%	90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 7.5% 100.0% 1.0%	10.0% 36.2% 31.7% H ₂ O 25.0% 13.8% 14.0% 25.0% 0.0% 80.0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg 12.15 MJ/kg 13.59 MJ/kg 19.36 MJ/kg 0.00 MJ/kg 1.92 MJ/kg
Fines 11.58% 0.0% 0.0% 0.0% 0.0% 0.0% 0.00% 0.0% 0.00% 0.0% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.00% 0.0% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.2% 0.0% 0.00% 0.00% 0.0%	Autroous Gyprock Other UTDI (Nov 15) report Concept of Design (Mirch 15) report AWT Residual Paper/Card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles	0.00% 0.00% 100.0% 100.0% 21.05% 20.00% 21.05% 20.00% 21.05% 3.16% 0.00%	0.0% 0.0% 17.0% 18.5% 0.0% 35.3% 70.7% 48.2% 48.2% 48.8% 0.0% 9.8% 20.4%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 4.7% 4.7% 4.7% 1.3% 1.3% 2.4%	0.0% 12.6% 14.3% 0.0% 21.1% 10.0% 21.1% 10.5% 0.0% 7.5% 4.4%	0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 0.0% 0.0% 0.3% 0.3% 0.2%	0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.3% 0.4% 0.0% 0.9% 9.5% 0.0% 0.0% 0.0% 0.0%	90,0% 31.7% 32,0% Ash 4.1% 2.6% 2.6% 7.5% 100.0% 1.0% 45.5%	10,0% 36,2% 31,7% 25,0% 13,8% 14,0% 25,0% 0,0% 80,0% 25,0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg NCV 12.15 MJ/kg 35.59 MJ/kg 19.36 MJ/kg 0.00 MJ/kg 1.92 MJ/kg 8.30 MJ/kg
Wood 4.21% 33.0% 4.2% 30.4% 0.1% 0.1% 0.0% 2.8% 23.8% 13.50 M/J Combustibles 2.113 2.04% 4.4% 0.2% 0.2% 0.9% 2.5% 2.5% 33.0 M/J Non-combustibles 1.05% 0.0%	Autorous Gyprock Other UTDI (Nov 15) report Eonoptiol Design (Murch 15) report AWT Residual Paper/card Plastic film Dense plastic Glass Vegetation Other combustibles Metal	0.00% 0.00% 100.0% 20.00% 21.05% 20.00% 21.05% 10.53% 0.00% 3.16% 3.16%	0.0% 0.0% 17.0% 18.5% C 35.3% 70.7% 48.2% 48.8% 0.0% 9.8% 20.4% 0.0%	0.0% 0.0% 2.1% 2.5% H 4.1% 4.1% 4.7% 4.1% 0.0% 1.3% 2.4% 0.0%	0.0% 12.6% 14.3% 0 31.4% 0.0% 21.1% 10.5% 0.0% 7.5% 4.4% 0.0%	0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 4.3% 0.0% 0.3% 0.3% 0.2% 0.0%	0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.3% 0.4% 0.0% 0.9% 9.5% 0.0% 0.0% 0.0% 0.0% 0.0%	90,0% 31,7% 32,0% Ash 4,1% 2,6% 2,6% 2,6% 7,5% 100,0% 46,5% 100,0%	10.0% 36.2% 31.7% 25.0% 13.8% 14.0% 25.0% 0.0% 80.0% 25.0% 0.0%	-0.24 MU/kg 5.67 MU/kg 6.31 MU/kg 12.15 MU/kg 35.59 MU/kg 19.36 MU/kg 0.00 MU/kg 0.00 MU/kg 0.00 MU/kg
Combustibles 21.113 20.4% 2.4% 4.4% 0.2% 0.2% 0.9% 4.65% 25.0% 8.30 M// Non-combustibles 1.05% 0.0%	Autorous Gyprock Other UTDI (Nov 15) report Conorpt-of Dasign (Mirch 15) report AWT Residual Paper/card Plastic film Dense plastic Textiles Glass Vegetation Other combustibles Metal Fines	0.00% 0.00% 100.0% 21.05% 20.00% 21.05% 20.00% 21.05% 3.16% 0.00% 3.16% 0.00% 11.58%	0.0% 0.0% 17.0% 18.5% C 35.3% 70.7% 48.2% 48.2% 0.0% 9.8% 20.4% 20.4% 0.0%	0.0% 0.0% 2.1% 2.5% H 4.1% 4.1% 4.1% 4.7% 4.2% 0.0% 0.0% 0.0%	0.0% 12.6% 14.3% 31.4% 0.0% 21.1% 10.5% 0.0% 7.5% 4.4% 0.0% 0.0%	0.0% 0.1% 0.5% 0.0% 0.0% 0.0% 0.0% 0.0% 0.3% 0.2% 0.3% 0.0%	0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.3% 0.4% 0.0% 0.9% 0.9% 0.0% 0.0% 0.0% 0.0%	90,0% 31.7% 32,0% Ash 4.1% 2.6% 2.6% 7.5% 100.0% 100.0% 100,0%	10.0% 36.2% 31.7% 4.20 25.0% 13.8% 14.0% 25.0% 0.0% 80.0% 80.0% 0.0%	-0.24 MJ/kg 5.67 MJ/kg 5.67 MJ/kg 5.31 MJ/kg 12.15 MJ/kg 15.59 MJ/kg 19.36 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg
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Hazardous 0.00% 0.0%	Autrobus Gyprock Other UTDI (Nov 15) report Concept of Dasign (March 15) report AWT Residual Paper/card Plastic film Dense plastic Textiles Glass Vegetaion Other combustibles Metal Fines Wood Combustibles	0.00% 0.00% 100.0% 21.05% 20.0% 21.05% 0.00% 3.16% 0.00% 3.16% 0.00% 11.58% 4.21%	0.0% 0.0% 17.0% 18.5% 20.3% 70.7% 48.2% 0.0% 48.8% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0%	0.0% 0.0% 2.1% 2.5% H 4.1% 12.1% 12.1% 4.7% 0.0% 1.3% 0.0% 0.0% 0.0% 0.0% 2.4%	0.0% 0.0% 12.8% 14.3% 0.0% 21.1% 10.5% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 30.4%	0.0% 0.1% 0.5% 0.5% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.1% 0.2% 0.2% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0	0.0% 0.3% 0.4% 0.0% 0.9% 9.5% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0.0% 0	90.0% 31.7% 32.0% Ash 4.1% 2.6% 2.6% 2.6% 7.5% 100.0% 1.0% 46.5%	10.0% 36.2% 31.7% 4.0 25.0% 14.0% 25.0% 0.0% 80.0% 80.0% 0.0% 25.0% 25.0%	-0.24 MJ/kg 5.67 MJ/kg 6.31 MJ/kg 12.15 MJ/kg 13.559 MJ/kg 19.36 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg 0.00 MJ/kg
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21292C TNG EfW Local Air Quality Assessment Revision 5.docx Job Number 21292C | AQU-NS-001-21292C

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Appendix E SENSITIVE RECEPTORS

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Sensitive Receptor	Easting (m)	Northing (m)
James Erskine Primary School	296748	6257187
Eskrine Park High School	296709	6256992
Clairgate Public School	296299	6258187
Minchinbury Public School	299287	6259084
Pinegrove Memorial Park Lawn Cemetery	300567	6258692
Sunny Patch Preparation School & Long Day Care Centre	297153	6258266
Eastern Creek Public School	301201	6259319
St Agnes Catholic High School	300761	6259894
All Areas Family Day Care Pty	299581	6258986
Maria Hawey Child Care Centre	299370	6259272
Jiminey Cricket Long Day Care	298562	6259310
White Bunny Child Care Centre	299792	6259530
LITTLESMARTIES	296419	6258212
Kidz Fun Factory	298128	6259445
Industrial facility	297743	6259085
Industrial facility	298017	6259102
Industrial facility	298262	6259157
Industrial facility	298362	6259444
Industrial facility	298106	6259473
Industrial facility	297650	6259598
Industrial facility	297391	6259845
Industrial facility	297425	6259607
Industrial facility	297528	6259706
Industrial facility	297827	6259711
Industrial facility	297923	6259624
Industrial facility	298057	6259589
Industrial facility	298165	6259576
Industrial facility	298169	6259723
Industrial facility	297988	6259754
Industrial facility	297855	6259871
Industrial facility	298473	6259809
Industrial facility	298254	6259912
Industrial facility	297964	6259979
Industrial facility	297807	6260039
Industrial facility	299645	6258440
Industrial facility	299645	6258037
Industrial facility	299709	6257886
Industrial facility	299541	6257851
Industrial facility	299441	6258055
Industrial facility	299490	6257405
Industrial facility	299906	6257425
Industrial facility	300157	6257390
Industrial facility	300263	6257339
Industrial facility	300447	6257583

Table E-1: Sensitive receptor locations

Pacific	Environment
Limited	

Sensitive Receptor	Easting (m)	Northing (m)
Industrial facility	300228	6257651
Industrial facility	300560	6257928
Industrial facility	300633	6257735
Industrial facility	300948	6257833
Industrial facility	300802	6257591
Industrial facility	300633	6257403
Industrial facility	300755	6257374
Industrial facility	301037	6257567
Industrial facility	301057	6257410
Industrial facility	301003	6257186
Industrial facility	300950	6257066
Industrial facility	300910	6256975
Industrial facility	300682	6257126
Industrial facility	300691	6257026
Industrial facility	300830	6257241
Industrial facility	300436	6257299
Industrial facility	299601	6257064
Industrial facility	299490	6256891
Industrial facility	299689	6256705
Industrial facility	299501	6256224
Industrial facility	300008	6256426
Industrial facility	300219	6256526
Industrial facility	300529	6256577
Industrial facility	300899	6256202
Industrial facility	300786	6255839
Industrial facility	301006	6255854
Industrial facility	298652	6255402
Industrial facility	298508	6255389
Industrial facility	298584	6255037
Industrial facility	296204	6256521
Industrial facility	296614	6256526
Industrial facility	296388	6256355
Industrial facility	296643	6256280
Industrial facility	296700	6256087
Industrial facility	296946	6256040
Industrial facility	296598	6255723
Industrial facility	296410	6255743
Industrial facility	296055	6255881

Pacific Environment Limited

Appendix F

FIVE YEAR ANALYSIS OF METEOROLOGY

As specified in the EPA's Approved Methods for the Modelling and Assessment of Air Pollutants in New South Wales five years of data are required to be reviewed so that a representative year of meteorological conditions can be selected.

Annual and seasonal wind roses for Horsley Park Equestrian Centre have been prepared for 2009 through 2013 and are shown below. All five years of data collected at Horsley Park Equestrian Centre show a similar pattern both annually and seasonally. There are some minor differences which are discussed below.

On an annual basis the prevailing wind directions originate from all directions of the compass, with fewer winds experienced from the northeast and north-northeast.

During summer the prevailing winds are dominated by flows originating from the eastern and southeastern quadrants.

Conversely, the months of winter are dominated by wind from the south-western and north-western quadrants. Almost no winds are experienced from the north-eastern and north-north-eastern directions across all years of data examined.

The wind distribution patterns for autumn and spring are less consistent and present a transition of summer to winter and vice versa across all years.

The percentage of calms is fairly consistent across all years and ranged between 14.2 % for 2009 and 24.5% for 2013.

Further analysis was conducted for the five years of data. The long term trend of monthly average temperature and monthly average wind speed is also shown below.

A strong seasonal trend in monthly average temperatures is evidenced with the highest temperatures experienced during the summer months of December, January and February and the lowest temperatures during the winter months of June, July and August. 2009 and 2013 are shown to experience higher monthly average temperatures across most months. Generally speaking, the monthly average temperatures at Horsley Park Equestrian Centre do not vary significantly from year to year.

There is no strong relationship between the time of year and the monthly average wind speed. Generally speaking, the monthly average wind speeds are less during the months of autumn. Both 2009 and 2010 measured the highest winds speed across the five years investigated. The lowest wind speed was recorded in 2013.

From this analysis, in addition to the consistent wind distribution patterns experienced discussed above it is considered that 2013 is a typical year and is therefore deemed a representative year for dispersion modelling.



Annual Calms = 14.2%

Annual and Seasonal Windroses Horsley Park Equestrian Centre 2009







Autumn Calms = 15.4%

Ν

Summer Calms = 10.1%





Spring Calms = 13.8%



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Annual Calms = 16.2%



s

Summer Calms = 15.6%



Annual and Seasonal Windroses Horsley Park Equestrian Centre

2010

Wind speed (m/s) >0.5 - 1.5

> >1.5 - 3 >3 - 4.5

> >4.5 - 6

>6 - 7.5 >7.5



Ν

s



Winter Calms = 18.6%

NNW NNE NW NE WNW ENE w Е 1% wsw ESE sw SE ssw SSE s

Spring Calms = 19.5%





Annual Calms = 20.8%

Annual and Seasonal Windroses Horsley Park Equestrian Centre 2011





Autumn Calms = 17.5%









Calms = 25.6%

Figure F-3: Annual and seasonal wind roses for Horsley Park Equestrian Centre (2011)

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Calms = 25.5%



Annual Calms = 23.7%

Ν

NNE

NE

SE

SSE

ENE

Е

ESE

NNW

NW

sw

ssw

WNW

w

wsw

Annual and Seasonal Windroses Horsley Park Equestrian Centre 2012







s



NE

SE

ENE

E

ESE





21292C TNG EfW Local Air Quality Assessment Revision 5.docx Job Number 21292C | AQU-NS-001-21292C



Annual Calms = 24.5%

Annual and Seasonal Windroses Horsley Park Equestrian Centre 2013





s

Summer Calms = 19.7% SE

SSE

sw

SSW









Figure F-5: Annual and seasonal wind roses for Horsley Park Equestrian Centre (2013)





Figure F-6: Monthly average temperature at Horsley Park Equestrian Centre (2009 - 2013)











Appendix G SUMMARY OF IN-STACK CONCENTRATION ESTIMATES

Table G-1: In-stack concentrations

-

Deremeter	In Stack Concentration (mg/m³)						
Parameter	Normal Conditions	Upset Conditions					
Benzoic Acid	1.00E-01	1.00E+00					
Hexa-decanoic Acid	3.70E-02	3.70E-01					
Ethyl Benzoic Acid	3.50E-02	3.50E-01					
Toluene	3.00E-02	3.00E-01					
Phthalate	2.00E-02	2.00E-01					
Dichloro-methane	2.00E-02	2.00E-01					
Acetone (propanone)	1.80E-02	1.80E-01					
Tetra-decanoic Acid	1.50E-02	1.50E-01					
Benzene	1.50E-02	1.50E-01					
Acetonitrile	1.40E-02	1.40E-01					
Xylene	1.00E-02	1.00E-01					
Trichloro-phenol	9.00E-03	9.00E-02					
Methyl-hexane	6.00E-03	6.00E-02					
Trichloro-ethylene	5.00E-03	5.00E-02					
Heptane	5.00E-03	5.00E-02					
PM10	1.00E+00	1.50E+02					
PM2.5	1.00E+00	1.50E+02					
HCI	9.00E+00	9.00E+01					
HF	4.00E+00	4.00E+01					
H2S	5.00E+00	5.00E+01					
SO2	2.70E+01	2.70E+02					
NO2	1.88E+02	1.88E+03					
СО	2.30E+01	2.30E+02					
NH3	2.00E+00	2.00E+01					
Нд	4.00E-03	1.30E-02					
Cd	9.00E-03	9.00E-02					
TI	1.00E-03	9.00E-03					
Ве	7.00E-06	5.25E-04					
Ag	3.40E-04	2.55E-02					
Zn	3.70E-02	5.09E+00					
Sn	3.33E-03	2.50E-01					
Мо	2.20E-05	2.63E-03					
Se	2.12E-03	2.12E-02					
As	4.00E-03	4.00E-02					
Sb	1.48E-02	1.48E-01					
Cr	4.67E-02	4.67E-01					
Pb	1.72E-01	1.72E+00					
Ni	2.10E-02	2.08E-01					
Cu	1.63E-02	2.45E-01					
Со	4.00E-03	4.00E-02					



Deremeter	In Stack Concentration (mg/m ³)							
Palameter	Normal Conditions	Upset Conditions						
Mn	3.65E-02	4.58E-01						
V	1.00E-03	1.50E-02						
PCDD/F	1.00E-08	5.00E-07						
PCBs	1.60E-08	1.60E-07						
НСВ	8.21E-06	8.21E-05						
PAHs	5.00E-04	5.00E-03						



Appendix H PERFORMANCE SPECIFICATIONS FOR EMERGENCY DIESEL GENERATORS



EPA Tier 2 Exhaust Emission **Compliance Statement** C3000 D5e 40CFR 60 Subpart IIII **50 Hz Diesel Generator Set**

Compliance Information:

The engine used in this generator set complies with Tier 2 emissions limit of U.S. EPA New Source Performance Standards for stationary emergency engines under the provisions of 40 CFR 60 Subpart IIII when tested per ISO8178 D2.

Engine Manufacturer: EPA Certificate Number: Effective Date: Date Issued: EPA Engine Family (Cummins Emissions Family): Cummins Inc ECEXL060,AAD 05/20/2013 05/20/2013 ECEXL060.AAD-017

Engine Information:

Cummins Inc. QSK78-G15 Model: Engine Nameplate HP: 3403 (Standby) 4 Cycle, VEE 18 Cylinder Diesel Type: Aspiration: Turbocharged and Aftercooled **Emission Control Device:** Turbocharged and Aftercooled

6.69 in. (170 mm) Bore: Stroke: 7.48 in. (190 mm) Displacement: Compression Ratio: 15.5:1

4735 cu. in. (77.6 liters)

Diesel Fuel Emission Limits	Gram	s per BH	Grams per kWm-hr			
D2 Cycle Exhaust Emissions	NOx NMHC	<u>co</u>	PM	NOx NMHC	<u>co</u>	PM
Test Results - Diesel Fuel (300-4000 ppm Sulfur)	3.30	1.50	0.22	4.40	2.00	0.29
EPA Emissions Limit	3.50	3.70	0.22	4.70	5.00	0.30

Test Methods: EPA/CARB Nonroad emissions recorded per 40CFR89 (ref. ISO8178-1) and weighted at load points prescribed in Subpart E, Appendix A for Constant Speed Engines (ref. ISO8178-4, D2)

Diesel Fuel Specifications: 40-48 Cetane Number, ASTM D975 No. 2-D.

Reference Conditions: Air Inlet Temperature: 25°C (77°F), Fuel Inlet Temperature: 40°C (104°F). Barometric Pressure: 100 kPa (29.53 in Hg), Humidity: 10.7 g/kg (75 grains H2O/lb) of dry air, required for NOx correction, Restrictions: Intake Restriction set to a maximum allowable limit for clean filter; Exhaust Back Pressure set to a maximum allowable limit.

Tests conducted using alternate test methods, instrumentation, fuel or reference conditions can yield different results. Engine operation with excessive air intake or exhaust restriction beyond published maximum limits, or with improper maintenance, may result in elevated emission levels.

Cummins Power Generation

Data and Specifications Subject to Change Without Notice

EPA-1259b

Inhaltsverzeichnis Contents

	Genset	Marine	0&G	Rail	C & I		
Application	x						
Engine model	20V4000G	63L 6ETC			-		
Application group	3B	3E	3F	3G	ЗН		
Emission Stage/Optimisation	fuel-optimised						
Test cycle	D2 + 110%	r.					
Data Set	XZ5965410	XZ59654100282					
Fuel sulphur content [ppm]	5						

Inhalt content	Notiz Note	Seite Page	Buchstabe/Revision change index
Emissions Daten Blatt (EDS) emission Data Sheet (EDS)	O2 gem. O2 meas.	2	a/b
Emissions Daten Blatt (EDS) emission Data Sheet (EDS)	O2 5% O2 5%	3	a/b/c
Emissions Daten Blatt (EDS) emission Data Sheet (EDS)	O2 15% O2 15%	4	b/c
Not to exceed Werte Not to exceed values	O2 gem. O2 meas.	5	a/b
Not to exceed Werte Not to exceed values	O2 5% O2 5%	6	a/b/c
Not to exceed Werte Not to exceed values	O2 15% O2 15%	7	b/c

					7		Benennung/Title
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH	Emissionsdatenblatt Emission Data Sheet	
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.
а	5% O2 und NTE-Datenblätter hinzu	27.05.15	Khakhol	Bearbeiter/Drawn by	15.05.2012	Koehler	
-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0464
Buchstabe/ Revision	Änderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept.	TKF	Veser	ED3 4000 0461

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Seite 1 von 7

Change index a D									
Motordaten									
engine data	Geneet	Marino	086	Pail	681	1			
Application	x	Marine	040	Itan	001				
Engine model	20V4000G	63L 6ETC							
Application group	3B	3E	3F	3G	3H	1			
Emission Stage/Optimisation	fuel-optimis	ed				1			
Test cycle	D2 + 110%					1			
fuel sulphur content [ppm]	5					1			
mg/mN ³ values base on	measured					1			
residual oxygen value of [%]	mouourou								
Motor Robemissionen*									
Engine raw emissions*									
Cycle point	[-]	n1	n2	n3	n4	n5	n6	n7	n8
Power (P/PN)	[-]	1,1	1	0,75	0.50	0,25	0,10		
Power	[kW]	2850	2590	2138	1425	712	-1		
Speed (n/nN)	[-]	1	1	1	1	1			
Speed	[rpm]	1500	1500	1500	1500	1500			
Exhaust temperature	1901	500	409	490	442	265			
after turbine	[0]	509	490	402	443	305			
Exhaust massflow	[kg/h]	13744	12196	10390	7963	5965			
Exhaust back pressure	[mbar]	27	21	14	7	3			
NOx	[g/kWh]	8,3	10,3	10,9	9,7	7,0			
	[mg/mN ³]	2581	3274	3338	2556	1205			
co	[g/kWh]	0,8	0,6	0,4	0,4	0,9			
	[mg/mN ³]	223	177	108	87	152			
HC	[g/kWh]	0,11	0,12	0,16	0,21	0,42			
	[mg/mN ³]	31	36	45	52	68			
02	[%]	8,4	8,4	8,8	10,1	12,9			
Particulate measured	[g/kWh]	-	-	-	-	-			
	[mg/mN ³]	-	~	-					
Particulate calculated	[g/kWh]	0,05	0,04	0,05	0,06	0,13			
	[mg/mN ³]	12	12	12	13	19			-
Dust (only TA-Luft)	[mg/mN ³]	-	-		-	-			
FSN	[-]	0,2	0,2	0,2	0,3	0,6			
NO/NO2**	[-]	-	-	-	-	-			
CO2	[g/kWh]	604,1	589,9	591,3	612,2	689,0			
002	[mg/mN ³]	175320	175277	169783	151727	112002			
502	[g/kWh]	0,002	0,002	0,002	0,002	0,002			
002	[mg/mN ³]	0,6	0,6	0,5	0,5	0,4			

Emission data measurement procedures are consistent with the respective emission evaluation process. Noncertified engines are measured to sales data (TVU/TEN) standard conditions.

These boundary conditions might not be representative for detailed dimensioning of exhaust gas aftertreatment, in this case it is recommended to contact the responsible department for more information.

Measurements are subject to variation. The nominal emission data shown is subject to instrumentation, measurement, facility, and engine toengine variations.

All data applies to an engine in new condition. Over extended operating time deterioration may occur which might have an impact on emission. Exhaust temperature depends on engine ambient conditions

** No standard test. To be measured on demand.

				[mtu]	7		Benennung/Title Emissionsdatenblatt	
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH	Emission Data Sheet		
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.	
а	5% O2 und NTE-Datenblätter hinzu	27.05.15	Khakhol	Bearbeiter/Drawn by	15.05.2012	Koehler		
-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0461	
Buchstabe/ Revision	Anderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept. TKF Veser		Veser	ED3 4000 040 I	
Vers.2.0								

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Seite 2 von 7

Revision Change indexabc									
Motordaten									
engine data						_			
1:377	Genset	Marine	0&G	Rail	C & I				
Application	x]			
Engine model	20V4000G	63L 6ETC		ан С]			
Application group	3B	3E	3F	3G	3H				
Emission Stage/Optimisation	fuel-optimis	sed							
Test cycle	D2 + 110%)							
fuel sulphur content [ppm]	5]			
mg/mN ³ values base on	5								
residual oxygen value of [%]	Ŭ								
Motor Pohomicsionan*									
Engine rewernissions*									
Cycle point	[-]	n1	n2	n3	n4	n5	n6	n7	n8
Power (P/PN)	[-]	1,1	1	0,75	0,50	0,25	0,10	10000	
Power	[KW]	2850	2590	2138	1425	712			
Speed (n/nN)	[-]	1	1	1	1	1			
Speed	[rpm]	1500	1500	1500	1500	1500			
Exhaust temperature	1901	500	400	400	442	205			
after turbine		509	498	482	443	305			
Exhaust massflow	[kg/h]	13744	12196	10390	7963	5965			
Exhaust back pressure	[mbar]	27	21	14	7	3			
NOv	[g/kWh]	8,3	10,3	10,9	9,7	7,0			
NOX	[mg/mN ³]	3267	4150	4371	3741	2367			
20	[g/kWh]	0,8	0,6	0,4	0,4	0,9			
0	[mg/mN ³]	283	224	142	128	298			
110	[g/kWh]	0,11	0,12	0,16	0,21	0,42			
HC	[mg/mN ³]	39	46	59	76	133			
02	[%]	5,0	5,0	5,0	5,0	5,0			
5 K 14	[g/kWh]	-		-	-				
Particulate measured	[mg/mN ³]	-	-	-	-	-			
-	[g/kWh]	0,05	0,04	0,05	0,06	0,13			
Particulate calculated	[mg/mN ³]	16	15	16	20	38			
Dust (only TA-Luft)	[mg/mN ³]	-		-	-	-			
FSN	[-]	0.2	0.2	0.2	0.3	0.6			
NO/NO2**	[-]	-	-,	-	-	-			
	[a/kWh]	604.1	589.9	591.3	612.2	689.0			
CO2	[ma/mN ³]	221936	222178	222343	222043	219919			
4-10-200	[a/kWh]	0.002	0.002	0.002	0.002	0.002			-
SO2	[ma/mN ³¹	0,002	0,002	0,002	0.7	0,002			
	[[IIIg/IIIN]]	0,7	0,7	0,7	0,7	0,7	-		

* Emission data measurement procedures are consistent with the respective emission evaluation process. Noncertified engines are measured to sales data (TVU/TEN) standard conditions.

These boundary conditions might not be representative for detailed dimensioning of exhaust gas aftertreatment, in this case it is recommended to contact the responsible department for more information. Measurements are subject to variation. The nominal emission data shown is subject to instrumentation, measurement, facility, and engine-to-

engine variations.

All data applies to an engine in new condition. Over extended operating time deterioration may occur which might have an impact on emission. Exhaust temperature depends on engine ambient conditions.

** No standard test. To be measured on demand.

				mtu	7		Benennung/Title Emissionsdatenblatt	
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrichshafen GmbH			Emission Data Sheet	
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.	
а	5% O2 und NTE-Datenblätter hinzu	27.05.15	Khakhol	Bearbeiter/Drawn by	15.05.2012	Koehler		
-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0461	
Buchstabe/ Revision	Änderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept.	TKF	Veser	EDS 4000 0481	
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Seite 3 von 7

Motordaten									
engine data									
ongino data	Genset	Marine	0 & G	Rail	C & I	1			
Application	x					1			
Engine model	20V4000G	63L 6ETC				1			
Application group	3B	3E	3F	3G	3H	1			
Emission Stage/Optimisation	fuel-optimis	ed				1			
Test cycle	D2 + 110%					1			
fuel sulphur content [ppm]	5]			
mg/mN ³ values base on	15]			
residual oxygen value of [%]	10								
Motor Rohemissioner*									
Engine raw emissions*									
Cycle point	[-]	n1	n2	n3	n4	n5	n6	n7	n8
Power (P/PN)	[-]	1.1	1	0.75	0.50	0.25	0.10		.10
Power	[kW]	2850	2590	2138	1425	712	0,10	-	
Speed (n/nN)	[-]	1	1	1	1420	1			
Speed	[rpm]	1500	1500	1500	1500	1500			
Exhaust temperature	[[piii]	1000	1000	1000	1000	1000			
after turbine	[°C]	509	498	482	443	365			
Exhaust massflow	[kg/h]	13744	12196	10390	7963	5965			
Exhaust back pressure	[mbar]	27	21	14	7	3			
NOv	[g/kWh]	8,3	10,3	10,9	9,7	7,0			
NOX	[mg/mN ³]	1225	1556	1639	1403	887			
<u>~~</u>	[g/kWh]	0,8	0,6	0,4	0,4	0,9			
0	[mg/mN ³]	106	84	53	48	112			
110	[g/kWh]	0,11	0,12	0,16	0,21	0,42			
пс	[mg/mN ³]	15	17	22	29	50			
02	[%]	15,0	15,0	15,0	15,0	15,0			
Destinuiste mensued	[g/kWh]	-	-	-	-	-			
Particulate measured	[mg/mN ³]	-	-	-	-	~			
Beatless de la contra de la contr	[g/kWh]	0,05	0,04	0,05	0,06	0,13			
Particulate calculated	[mg/mN ³]	6	6	6	7	14			
Dust (only TA-Luft)	[mg/mN ³]	-	-	-	-	-			
FSN	[-]	0,2	0,2	0,2	0,3	0,6			
NO/NO2**	[-]	-		-	-	-			
	[g/kWh]	604,1	589.9	591,3	612.2	689,0			-
02	[ma/mN ³]	83226	83317	83379	83266	82470			
nens noc				~~~ ~					
	[ng/kWh]	0.002	0.002	0.002	0.002	0.002			

* Emission data measurement procedures are consistent with the respective emission evaluation process. Noncertified engines are measured to sales data (TVU/TEN) standard conditions.

These boundary conditions might not be representative for detailed dimensioning of exhaust gas aftertreatment, in this case it is recommended to contact the responsible department for more information.

Measurements are subject to variation. The nominal emission data shown is subject to instrumentation, measurement, facility, and engine-toengine variations.

All data applies to an engine in new condition. Over extended operating time deterioration may occur which might have an impact on emission. Exhaust temperature depends on engine ambient conditions.

** No standard test. To be measured on demand.

				mtu	7		Benennung/Title Emissionsdatenblatt	
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH	Emission Data Sheet		
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а	5% O2 und NTE-Datenblätter hinzu	27.05.15	Khakhol	Bearbeiter/Drawn by	15.05.2012	Koehler]	
-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0461	
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Seite 4 von 7

Application	Gancat	Marino	086	Pail	180				
ADDILAUDU	V	Warme	040	IXan	- Car				
Engine model	20\/4000G6	3L 6ETC							
Application group	38	3E 0210	3F	3G	3H				
Emission Stage/Ontimisation	fuel ontimise		51	50	511				
Test cycle	D2 + 110%	Ju -							
fuel sulphur content [ppm]	5								
mg/mN ³ values base on	5								
residual oxygen value of [%]	measured								
Not to exceed Werte*									
not to exceed values"				•					
Cycle point	[-]	n1	n2	n3	n4	n5	n6	n7	n8
Power (P/PN)	[-]	1	0,75	0,50	0,25				
Power	[KVV]	2590	2138	1425	/12				
Speed (n/nN)	[-]	1	1	1	1				
Speed	[rpm]	1500	1500	1500	1500				
Exhaust back pressure	[mbar]	12.4	14	12.0	3				
NOx	[g/kvvn]	13,4	14,2	12,6	10,5				
	[mg/mN ^s]	4256	4339	3323	1808				
со	[g/kvVh]	1,0	0,6	0,7	1,9				-
	[mg/mN ^s]	301	184	166	304				
HC	[g/kvvn]	0,21	0,27	0,40	0,83				
02	[mg/mN*j	61	//	99	135			-	
02	[%]	8,4	8,8	10,1	12,9				
Particulate calculated	[g/kvvn]	0,07	0,08	0,09	0,20				
	cording the given v	alue in this EL	DS						
 Exhaust gas back pressure acc Fuel according to EN 590 or US Coolant and Lubricants accordin The nominal emissions data show operating points and thus cannot type of exhaust gas aftertreatmer information. Field emission test data are not g procedures, and instrumentation. exhaust restriction beyond publis MTU Friedrichshafen GmbH has r 	S EPA 40CFR89 ng MTU Fuels and wn is subject to in: be used to comp nt that may be ins guaranteed to the: Over time deterior hed maximum lim made efforts to er hout obligation or	d Lubricants S instrumentation, are to EPA reg italled on the e se levels. Actu oration may oc- nits, or with im usure that the i itability. No li	pecification , measurement gulations which ingine, therefor al field test res cur which may proper mainter nformation in tr ability for any c	t, facility and e n use values b e it is suggest sults may vary have an impa nance, may rei his data sheet errors, facts o	engine to engin ased on a weig ed that the eng due to test situ ct on emission sults in elevate is accurate, b opinions is ac	e variations. E hted cycle. E gine manufact e conditions, i levels. Engine d emission le ut reserves th iccented Cust	missions data missions data urer be contac nstallation, fue e operation wit cels. e right to amer omers must se	a is based on s a may vary dep ted directly for el specification h excessive ai nd specification	ingle ending on the further , test r intake or ns and es as to the

							Emissionsdatenblatt		
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH		Emission Data Sneet		
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.		
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-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0461		
Buchstabe/ Revision	Änderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept.	TKF	Veser	ED3 4000 0481		
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Seite 5 von 7

engine data									
	Genset	Marine	0 & G	Rail	C & I				
Application	x								
Engine model	20V4000G63	BL 6ETC							
Application group	3B	3E	3F	3G	3H				
mission Stage/Optimisation	fuel-optimise	d							
est cycle	D2 + 110%				Î				
uel sulphur content [ppm]	5								
ng/mN ³ values base on	5								
esidual oxygen value of [%]	Ŭ.								
lot to exceed Werte*									
of to exceed values*									
	[1-1	n1	n2	n3	n4	n5	n6	n7	n8
ower (P/PN)	[-]	1	0.75	0.50	0.25	110	110		10
ower	[F]	2590	2138	1425	712				
peed (n/nN)	[-]	1	1	1	1			-	1
peed	[rpm]	1500	1500	1500	1500			-	
xhaust back pressure	[mbar]	21	14	7	3			-	
	[g/kWh]	13.4	14.2	12.6	10.5			1	
Ox	[mg/mN ³]	5395	5683	4863	3550			+	
со	[g/kWh]	1,0	0,6	0,7	1,9				
	[mg/mN ³]	381	241	242	597			1	1
	[a/kWh]	0.21	0.27	0.40	0.83			+	
	[mg/mN ³]	78	100	145	265			-	
)2	[%]	5,0	5,0	5.0	5.0				
	[g/kWh]	0,07	0,08	0.09	0.20				
articulate calculated	10 1	1.1.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2			0,20		1		
Calculated values are not proven missions data measurement pro he NOx, CO, HC and PM emissi onditions: Ambient air pressure 1 bar	[mg/mN ³] by tests and the acedures are const ion data tabulated	23 refore the accu istent with tho I here were tak	26 uracy cannot b use described i ten from a sing	29 e guaranteed. n the applicab le new engine	57 57 le rules and st under the test	andards. t conditions s	hown above a	nd are valid for	the following
Calculated values are not proven Emissions data measurement pro- the NOx, CO, HC and PM emissi conditions: Ambient air pressure 1 bar Air intake temperature approx. 2 Rel. Humidity 30%-60% New Engine New standard- air filter Exhaust gas back pressure acc Fuel according to EN 590 or US Coolant and Lubricants accordin the nominal emissions data show opperating points and thus cannot uppe of exhaust gas aftertreatmen information. Teid emission test data are not g procedures, and instrumentation. xxhaust restriction beyond publish ITU Friedrichshafen GmbH has n formation without notice and with uitability of this product for their at the nominat	[Img/mNP] by tests and the cecdures are cons ion data tabulated 55°C ording the given v EPA 40CFR89 gg MTU Fuels and m is subject to in be used to compu- t that may be ins' uaranteed to these uaranteed to these need maximum lim nade efforts to en heut obligation or application. No re	23 refore the accr isistent with tho here were tak alue in this EE Lubricants Sp strumentation, are to EPA reg talled on the e we levels. Actur ration may occ its, or with im sure that the ii isponsibility for	26 uracy cannot b use described i teen from a sing DS DS DS DS Decification measurement nulations which may be a sur- proper mainten nformation in ti nformation in ti ability for any e r any loss as a	29 e guaranteed. n the applicab le new engine , facility and e use values ba e it is suggest ults may vary have an impac ance, may res nis data sheet terrors, facts or result of any p	le rules and st under the tesi ngine to engin sed on a weig ed that the en- due to test sit to n emission utlts in elevate is accurate, b opinions is ac	e variations. I t conditions s hted cycle. I gine manufac e conditions, levels. Engin d emission le ut reserves th coperted. Cust reliance on a	Emissions dat Emissions dat Emissions dat Lutrer be conta installation, fi e operation w vels. e right to ame omers must so	nd are valid for ta is based on a ta may vary dep ccted directly fo uel specification ith excessive a end specificatio attisfy themselv ontained in this	the following bending on the further h, test ir intake or ns and ese as to the c data sheet

				mtu	7		Benennung/Title Emissionsdatenblatt	
с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH	Emission Data Sheet		
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.	
а	5% O2 und NTE-Datenblätter hinzu	27.05.15	Khakhol	Bearbeiter/Drawn by	15.05.2012	Koehler]	
-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDE 4000 0464	
Buchstabe/ Revision	Anderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept.	TKF	Veser	EDS 4000 0461	

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Seite 6 von 7

Change index b C						
Motordaten						
engine data						
	Genset	Marine	0&G	Rail	C & I	
Application	x					
Engine model	20V4000G63	3L 6ETC			00	
Application group	3B	3E	3F	3G	3H	
Emission Stage/Optimisation	fuel-optimise	ed				
Test cycle	D2 + 110%					
fuel sulphur content [ppm]	5					
mg/mN ^a values base on residual ovvigen value of (%).						

not to exceed values

Cycle point	[-]	n1	n2	n3	n4	n5	n6	n7	n8
Power (P/PN)	[-]	1	0,75	0,50	0,25				
Power	[kW]	2590	2138	1425	712				
Speed (n/nN)	[-]	1	1	1	1				
Speed	[rpm]	1500	1500	1500	1500				
Exhaust back pressure	[mbar]	21	14	7	3				
Nov	[g/kWh]	13,4	14,2	12,6	10,5				
NOX	[mg/mN ³]	2023	2131	1824	1331				
	[g/kWh]	1,0	0,6	0,7	1,9				
0	[mg/mN ³]	143	90	91	224				
10	[g/kWh]	0,21	0,27	0,40	0,83				
HC	[mg/mN ³]	29	38	54	99				
O2	[%]	15,0	15,0	15,0	15,0				
Destioulate coloulated	[g/kWh]	0,07	0,08	0,09	0,20				
Particulate calculated	[mg/mN ³]	9	10	11	21				
* Calculated values are not prov	en by tests and the	erefore the acc	uracy cannot	be guaranteed.					

Emissions data measurement procedures are consistent with those described in the applicable rules and standards. The NOx, CO, HC and PM emission data tabulated here were taken from a single new engine under the test conditions shown above and are valid for the following

conditions Ambient air pressure 1 bar

Air intake temperature approx. 25°C
 Rel. Humidity 30%-60%

New Engine

New standard- air filter

Exhaust gas back pressure according the given value in this EDS
 Fuel according to EN 590 or US EPA 40CFR89
 Coolant and Lubricants according MTU Fuels and Lubricants Specification

The nominal emissions data shown is subject to instrumentation, measurement, facility and engine to engine variations. Emissions data is based on single operating points and thus cannot be used to compare to EPA regulations which use values based on a weighted cycle. Emissions data may vary depending on the type of exhaust gas aftertreatment that may be installed on the engine, therefore it is suggested that the engine manufacturer be contacted directly for further information.

Field emission test data are not guaranteed to these levels. Actual field test results may vary due to test site conditions, installation, fuel specification, test procedures, and instrumentation. Over time deterioration may occur which may have an impact on emission levels. Engine operation with excessive air intake or exhaust restriction beyond published maximum limits, or with improper maintenance, may results in elevated emission levels.

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с	Fehler korrigiert	09.08.16	Khakhol	MTU Friedrich	shafen GmbH	Emission Data Sheet		
b	Seite 4 und 7 hinzugefügt	20.06.16	Khakhol		Datum/Date	Name/Name	Zeichnungs-Nr./Drawing No.	
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-	Freigabe	15.05.12	Link	Geprüft/Checked	15.05.2012	Koehler	EDS 4000 0464	
Buchstabe/ Revision	Anderung Modifikation	Datum Date	Name Name	OrgEinheit/Dept.	TKF	Veser	ED3 4000 0461	

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21292C TNG EfW Local Air Quality Assessment Revision 5.docx Job Number 21292C | AQU-NS-001-21292C

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Appendix I

SUMMARY OF EMISSIONS PERFORMANCE

I.1 SUMMARY OF EMISSIONS PERFORMANCE REPORTED IN WSP (2000)



I.2 CADMIUM AND METALS EMISSIONS PERFORMANCE IN HZI PLANTS

Heavy metal emissions HZI plants with semi-dry FGT in UK

			Plant A		Pla	nt B	Plant C	Average	EU	
Metal	Symbol	Line 1	Line 2	Line 3	Line 1	Line 2	Line 3		WID	
Mercury	Hg	0.0015	0.0004	0.0002	0.004	0.003	0.0017	0.002	< 0.05	
Cadmium	Cd	0.00270	0.00085	0.00111	0.009	0.001	0.004			
Thallium	TI	0.00005	0.00003	0.00002	0.000	0.000	0.0009	1000	1.00	
Sum Cd+Tl	Cd + TI	0.00275	0.00087	0.00113	0.009	0.001	0.0049	0.003	< 0.05	
Arsenic	As	0.0006	0.0003	0.0004	0.003	0.000	0.0013			
Antimony	Sb	0.0148	0.0047	0.0047	0.007	0.001	0.0026			
Chromium	Cr	0.0179	0.0115	0.0399	0.014	0.002	0.0467			
Cobalt	Co	0.0003	0.0002	0.0001	0.003	0.000	0.0006			
Copper	Cu	0.0085	0.0085	0.0263	0.051	0.001	0.0049			
Lead	Pb	0.0452	0.0137	0.0170	0.172	0.002	0.0094			
Manganese	Mn	0.0084	0.0041	0.0037	0.095	0.005	0.0051			
Nickel	Ni	0.0118	0.0058	0.0041	0.006	0.002	0.0208			
Vanadium	V	0.0003	0.0002	0.0004	0.003	0.000	0.0004	100 C		
Sum heavy metal	As-V	0.11	0.049	0.097	0.35	0.015	0.092	0.12	< 0.5	

all data in mg/m³ at STP and referred to 11% O₂ dry

28.4.2014 / HZI / Fy
I.3 SAMPLE CEMS REPORT FROM RIVERSIDE

Riverside Resource Recovery emission report – February 2014

The following charts summarise the emission data for the Riverside Resource Recovery facility. The charts show the MAXIMUM readings taken during the month.



<u>Why do we control and monitor Particulates (dust)?</u> Particulates is the term used to describe tiny particles in the air, made up of a complex mixture of soot, organic and inorganic materials having a particle size less than or equal to 10 microns diameter (10 microns is equal to one hundredth part of a millimetre). Particulates is one of the eight substances for which the government has established an air quality standard as part of



Why do we control and monitor Oxides of Nitrogen (NOx)? NOx includes various compounds, but is usually used to group two gases; nitrogen dioxide (NO₂) and nitric oxide (NO). These can be formed naturally, but are also formed from man-made processes like fuel combustion or biomass burning. There are a number of health and environmental issues attributed to NOx, including smog, acid rain, and possibly global warming.



Why do we control and monitor Carbon Monoxide?

Carbon monoxide is both a common naturally occurring chemical and is manufactured by man. It is a colourless, odourless poisonous gas. Carbon monoxide is one of the eight substances for which the government has established an air quality standard as part of its national Air Quality Strategy.

Carbon monoxide can cause harmful health effects by reducing oxygen delivery to the body's organs and tissues.



Why do we control and monitor Total Organic Carbon (TOC)?

Total Organic Carbon (TOC) consists of a wide range of organic compounds including Volatile Organic Compounds (VOCs). VOCs are numerous, varied and found everywhere. VOCs are of general concern because of their ability to react with other pollutants (such as nitrogen oxides) in the lower atmosphere to form ozone. High concentrations of ozone at ground level can harm human health, damage crops and affect materials such as rubber. Some VOCs may be directly harmful to human health, contribute to global warming or destroy stratospheric ozone needed to shield the earth's surface from harmful ultra violet radiation.





Why do we control and monitor Sulphur Dioxide and Hydrogen Chloride? Both gases dissolve in water to form strong acids and thus can contribute to the formation of acid rain. Acid rain is environmentally damaging to crops, soils and waters.



Why do we control and monitor Ammonia?

Although in wide-use in several industries, ammonia is both caustic and hazardous. It is a colourless gas with a characteristic pungent odour.

Ammonia, unlike the other species monitored, is not a product from the incineration of waste but is actually introduced into the furnace. Under the right conditions, ammonia is able to reduce oxides of nitrogen found in the flue gas by the chemical process Selective Non-Catalytic Reduction (SNCR) to nitrogen and water vapour which are both non-hazardous.

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Appendix J DETAILED MODELLING PREDICTIONS

To inform a Human Health Risk Assessment, modelling predictions are presented in Table I.1 for the discrete receptors described in Appendix C (particularly sensitive receptors such as schools and childcare centres).

There are too many residential receptors in the suburbs of Minchinbury and Erskine Park to provide individual modelling predictions for each, however modelling predictions for the closest residential receptors are shown in Table I.1 at the discrete locations shown in Figure J-1. Note that the cells highlighted grey represent the sensitive receptors with the highest concentration of those assessed.



Figure J-1: Receptor locations shown in result tables

J.1.1 Normal Operations

				0							5 51					
			1 hour	10- minute	1 hour	1 hour	1 hour	15- minute	1 hour	1 hour	1 hour	1 hour	1 hour	1 hour	1 hour	1 hour
Sensitive Receptor	Easting (m)	Northing (m)	NO ₂	SO ₂		PM10 / PM2.5	H ₂ S	СО		HCI	Cd	Hg	Dioxins / Furans	TOC (as benzene)	NH₃	PAH (as benzo(a) pyrene)
			µg/m³	μg,	/m³	µg/m³	µg∕m³	mg	/m³	mg/m³	mg/m³	mg/m³	mg/m³	mg/m³	mg/m³	mg/m³
			100 th	100 th		100 th	99th	100 th		100 th	99.9 th	99.9 th	99.9 th	99.9 th	99.9 th	99.9 th
Maximum (outside site boundary)			5.20E+01	1.10E+01	7.40E+00	2.70E-01	3.60E-01	8.30E-03	6.30E-03	2.50E-03	2.00E-06	8.70E-07	2.20E-12	3.30E-06	4.40E-04	1.10E-07
James Erskine Primary School	296748	6257187	1.4E+01	2.9E+00	2.0E+00	7.5E-02	3.6E-01	2.2E-03	1.7E-03	6.7E-04	6.6E-07	2.9E-07	7.4E-13	1.1E-06	1.5E-04	3.7E-08
Eskrine Park High School	296709	6256992	1.4E+01	3.0E+00	2.1E+00	7.6E-02	3.6E-01	2.4E-03	1.8E-03	6.9E-04	6.4E-07	2.8E-07	7.1E-13	1.1E-06	1.4E-04	3.6E-08
Clairgate Public School	296299	6258187	2.0E+01	4.0E+00	2.8E+00	1.0E-01	3.6E-01	3.2E-03	2.4E-03	9.3E-04	5.6E-07	2.5E-07	6.2E-13	9.3E-07	1.2E-04	3.1E-08
Minchinbury Public School	299287	6259084	4.6E+01	9.6E+00	6.7E+00	2.5E-01	3.6E-01	7.5E-03	5.7E-03	2.2E-03	1.8E-06	7.8E-07	2.0E-12	2.9E-06	3.9E-04	9.8E-08
Pinegrove Memorial Park Lawn Cemetery	300567	6258692	4.1E+01	8.4E+00	5.9E+00	2.2E-01	3.6E-01	6.6E-03	5.0E-03	2.0E-03	1.3E-06	5.8E-07	1.5E-12	2.2E-06	2.9E-04	7.3E-08
Sunny Patch Preparation School & Long Day Care Centre	297153	6258266	2.7E+01	5.6E+00	3.9E+00	1.5E-01	3.6E-01	4.5E-03	3.4E-03	1.3E-03	8.8E-07	3.9E-07	9.8E-13	1.5E-06	2.0E-04	4.9E-08
Eastern Creek Public School	301201	6259319	3.6E+01	7.3E+00	5.1E+00	1.9E-01	3.6E-01	5.7E-03	4.3E-03	1.7E-03	1.1E-06	4.7E-07	1.2E-12	1.8E-06	2.4E-04	5.9E-08
St Agnes Catholic High School	300761	6259894	3.1E+01	6.4E+00	4.5E+00	1.7E-01	3.6E-01	5.1E-03	3.9E-03	1.5E-03	1.2E-06	5.3E-07	1.3E-12	2.0E-06	2.6E-04	6.6E-08
All Areas Family Day Care Pty	299581	6258986	5.1E+01	1.0E+01	7.3E+00	2.7E-01	3.6E-01	8.3E-03	6.3E-03	2.4E-03	1.7E-06	7.4E-07	1.9E-12	2.8E-06	3.7E-04	9.3E-08
Maria Hawey Child Care Centre	299370	6259272	4.4E+01	9.0E+00	6.3E+00	2.3E-01	3.6E-01	7.1E-03	5.4E-03	2.1E-03	1.7E-06	7.6E-07	1.9E-12	2.8E-06	3.8E-04	9.5E-08
Jiminey Cricket Long Day Care	298562	6259310	4.7E+01	9.6E+00	6.7E+00	2.5E-01	3.6E-01	7.5E-03	5.7E-03	2.2E-03	1.7E-06	7.6E-07	1.9E-12	2.8E-06	3.8E-04	9.5E-08
White Bunny Child Care Centre	299792	6259530	4.4E+01	9.2E+00	6.4E+00	2.4E-01	3.6E-01	7.1E-03	5.4E-03	2.1E-03	1.5E-06	6.7E-07	1.7E-12	2.5E-06	3.3E-04	8.3E-08
LITTLESMARTIES	296419	6258212	2.1E+01	4.3E+00	3.0E+00	1.1E-01	3.6E-01	3.3E-03	2.5E-03	9.9E-04	5.9E-07	2.6E-07	6.6E-13	9.9E-07	1.3E-04	3.3E-08
Kidz Fun Factory	298128	6259445	4.1E+01	8.4E+00	5.9E+00	2.2E-01	3.6E-01	6.6E-03	5.0E-03	2.0E-03	1.4E-06	6.3E-07	1.6E-12	2.4E-06	3.1E-04	7.9E-08
Closest	297450	6256754	1.8E+01	3.6E+00	2.5E+00	9.3E-02	3.6E-01	2.8E-03	2.1E-03	8.4E-04	8.2E-07	3.7E-07	9.1E-13	1.4E-06	1.8E-04	4.6E-08
receptors to	297500	6256754	1.8E+01	3./E+00	2.6E+00	9.5E-02	3.6E-01	2.9E-03	2.2E-03	8.6E-04	8.5E-07	3.8E-07	9.4E-13	1.4E-06	1.9E-04	4./E-08

Table J-1: Predicted ground level concentrations at particular sensitive receptors – short term averaging periods