



We are the Environment Agency. It's our job to look after your environment and make it a better place – for you, and for future generations.

Your environment is the air you breather the state of the

Your environment is the air you breathe, the water you drink and the ground you walk on. Working with business, Government and society as a whole, we are making your environment cleaner and healthier.

The Environment Agency. Out there, making your environment a better place.

Published by.
Environment Agency
Rio House
Waterside Drive,
Aztec West Almondsbury,
3ristol BS32 4UD
Tel: 0870 8506506
Email: enquiries@environment-agency

Email: enquiries@environment-agency.gov.uk www.environment-agency.gov.uk

© Environment Agency

All rights reserved. This document may be reproduced with prior permission of the Environment Agency. March 2009

GEHO0209BPIN-E-E

Contents

Contents	JAI081
	0,
Introduction	2
Installations covered	3
Key issues	4
Managing your activities 1.1 Energy efficiency	7
1.1 Energy efficiency	7
1.2 Avoidance, recovery and disposal of wastes	9
2. Operations	11
	11
3. Emissions and monitoring	13
3.1 Point source emissions to water	13
3.2 Point source emissions to air	15
3.3 Fugitive emissions	27
3.4 Monitoring	28
4. Annexes	32
Annex 1- Emission-benchmarks	32
Annex 2- Other relevant guidance and abbreviations	44
Annex 3- Regulatory information	45
3.1 Point source emissions to water	
O	

Introduction

In "Getting the basics right – how to comply with your environmental permit" (GTBR) we described the standards and measures that we expect businesses to take in order to control the risk of pollution from the most frequent situations in the waste management and process industries.

This sector guidance note (SGN) is one of a series of additional guidance for Part A(1) activities listed in Schedule 1 of the Environmental Permitting Regulations (the Regulations). We expect you to use the standards and measures in this note in addition to those in GTBR to meet the objectives in your permit.

Sometimes, particularly difficult issues arise such as problems with odour or noise. You may then need to consult the "horizontal" guidance that gives in depth information on particular topics. Annex 1 of GTBR lists these.

The IPPC Directive requires that the Best Available Techniques (BAT) are used. When making an application, explain how you will comply with each of the indicative BATs in this sector guidance note. Where indicative BAT is not included, where you propose to use an alternative measure or where there is a choice of options you should explain your choice on the basis of costs and benefits. Part 2 of Horizontal Guidance Note H1 Environmental Risk

Assessment (see GTPR Annex 1) gives a formal method of as essing options which you should use where major decisions are to be made.

We will consider the relevance and relative importance of the information to the installation concerned when making technical judgments about the installation and when setting conditions in the permit.

Modern permits describe the objectives (or outcomes) that we want you to achieve. They do not normally tell you how to achieve them. They give you a degree of flexibility.

Where a condition requires you to take appropriate measures to secure a particular objective, we will expect you to use, at least, the measures described which are appropriate for meeting the objective. You may have described the measures you propose in your application or in a relevant management plan but further measures will be necessary if the objectives are not met.

The measures set out in this note may not all be appropriate for a particular circumstance and you may implement equivalent measures that achieve the same objective. In cases where the measures are mandatory this is stated.

In response to the application form question on Operating Techniques, you should address each of the measures described as indicative BAT in this note as well as the key issues identified in GTBR.

Unless otherwise specified, the measures and benchmarks described in this note reflect those of the previous Sector Guidance Note. They will be reviewed in the light of future BREF note revisions. In the meantime we will take account of advances in BAT when considering any changes to your process.

Installations covered

This note applies to activities regulated under the following section of schedule 1 of the Regulations:

Section 1.1 Combustion Activities, Part A(1):

(a) Burning any fuel in an appliance with a rated thermal input of 50 mera vatts or more. The reference to thermal rated input shall be taken to refer to gross, not net, input.

Burning any of the following fuels in an appliance with a rated thermal input of 3 megawatts or more, but less than 50 megawatts, unless the activity is carried out as part of a Part A (2) or B activity -

- (i) waste oil;
- (ii) recovered oil;

(iii) any fuel manufactured from, or comprising any other waste.

Selected activities of Section 1.2 Part A(1):

- (e) Producing gas from oil or other carbonaccous material or from mixtures thereof, other than from sewage, unless the production is carried out as part of an activity which is a combustion activity (whether or not that combustion activity is described in Section 1.1).
- (j) Activities involving the pyrolysis, carbonisation, distillation, liquefaction, gasification, partial oxidation, or other heat treatment of coal (other than the drying of coal), lignite, oil, other carbonaceous material or mixtures thereof otherwise than with a view to making charcoal.

Operators of landfill gas engines should refer to LFTGN 6 Guidance on gas treatment technologies for Landfill Gas Engines¹

Appliances operated on the same site by the same operator should be aggregated. This note applies if the total exceeds 50MW. This includes those rated <3 MWth and standby plant.

3

¹ goto http://www.environment-agency.gov.uk/commondata/acrobat/lftgn06 treatment 93 6556.pdf

4

Directly associated activities

The installation will also include **directly associated activities**, which have a direct technical connection with the main activities, and which may have an effect on emissions and pollution. These may involve activities such as:

- the storage and handling of raw materials; the storage and despatch of finished products, waste and other materials
- the control and abatement systems, where not an integral part of the activity
- waste treatment or recycling

Tables 3.1 and 3.2 (Annex 3) provide information regarding which part of the regulations apply to your plant.

Certain biomass wastes are exempt from the requirements of WID. For further information refer to IPPC Co-Incineration Supplement, http://www.environment-agency.cov.uk/commondata/acrobat/coincinerator-924475 924475.pdf

involving the simultaneous use of two or more fuels, the LCPD sets the method by which emissions limit values should be calculated.

Plaster processes are included in this sector because of the aggregation rule. This part of the combustion sector differs

as it uses combustion techniques to modify mineral properties within its processes and does not generate power or steam.

Key issues

The key issues are:

Key emissions to air

- sulphur dioxide
- nitrogen oxides
- particulate matter, particularly from solid fuel fired plants
- trace substances, such as heavy metals and dioxins.

In any combustion activity, there is an inter-relation between the CO concentration, NOx concentration, carbon in ash and thermal efficiency. Combustion should be controlled to provide an optimum compromise between these factors.

Global warming

Global warming is addressed under climate change levy agreements and/or the Greenhouse Gas Emissions Trading Regulations. This environmental permit considers basic energy efficiency measures.

3r the manufacture of gyptum and cement manufacture. The Landfill Directive requires specialist pre-treatment of some

residues before they can be landfilled.

Burning alternative fuels

Assified as a classified as a classified as a classified as ander the Waste are (WID). The relativen within that Direction within that Direction and residues from air pollution control aipment require disposal. Where cossible, these should be recycled e.g. in

Releases to water

Cooling water discharge is usually the key release. There may also be some aqueous discharges from air pollution control equipment.

Managing your activities and wastes 1.1 Energy efficiency 1.2 Avoidance, recovery wastes

- avoid wastes

6

Energy efficiency

1. Managing your activities

1.1 Energy efficiency

For power generating cycles, the maximum theoretical (thermodynamic) efficiency is obtained by maximising the working temperature difference. This implies high combustion temperatures and minimising the temperature at which heat is rejected from the cycle.

Fuel should be combusted as completely as possible and the heat generated should be transferred efficiently to the working medium, usually water in the case of boilers.

In addition to efficient boiler heat transfer design, further efficiency improvements may be obtained by optimal heat recovery through combustion air preheating and feedwater heating. Practical limits are leached by the tendency for increased NOx formation with high air preheating and by the risk of condensation and corrosion by overcooled stack gases. You should account for these factors when designing plant.

Indicative BAT

You should where appropriate:

 Demonstrate that the proposed or current situation represents BAT where there are other considerations involved, e.g. recovering energy from waste affects the energy efficiency of the process.

Boilers and jurnaces

- 2. Carry out regular checks to minimize leakage of air into units operating below atmospheric pressure.
- 3. Ensure good design, operation and maintenance of burners.

team turbines

- Replace existing turbines with more efficient turbines.
- 5. Increase cycle efficiency by measures such as reheating steam between stages, improving the vacuum on condensers and using very high, including supercritical, pressures to increase the working temperature difference and cycle efficiency.
- 6. Take steam from between stages or from a backpressure exhaust for use in, for example, process or building heating.

Energy efficiency

Gas turbines

 In large installations, consider installing more than one smaller turbine to allow for more efficient load following efficient load following.

017A10812018

- 8. Consider measures to improve the efficiency of the turbine:
 - increasing the combustion temperature, but balanced against increase NOx levels and amounts of excess air required
 - using concentric shafts to connect different stages of compression and expansion this is common in aero derivative machines
 - intercooling between stages of air compression and reheating between stages of expansion. Such features are normally incorporated at the design stage and may not be relevant to retrofits
- 9. The exhaust from even the most efficient gas turbines contain large amounts of heat that should be recovered, and used for process or building heating (CHP), or steam may be fed to a steam turbine to provide additional power in a combined cycle (CCGT). Supplementary fuel may also be fired in the heat recovery boiler to meet the heat demands. The design of the veter needs to optimize the characteristics of the turbines and boiler to achieve the best overall performance.

Reciprocating engines

- 10. Maximise engine efficiency by measures such as turbo charging and air intercooling. However, this should be balanced against increased NOx emissions.
- 11. Recover exhaust gas heat for process or building heating or absorption chilling.
- These features are usually incorporated at the design stage.) 13. Where additional heat is required, supplementary fuel may be fired into the boiler, almough this can be complicated by the pulsating exhaust and size limitations.

16 Jan 10 Avoidance, recovery and disposal of wastes

1.2 Avoidance, recovery and disposal of wastes

Indicative BAT

You should where appropriate:

- 1. Store, handle and transport all waste streams to prevent the release of waste, dust, VOC, leachate or odour.
- 2. Store bottom ash and fly ash separately. This provides flexibility to re-use the different ash fractions.
- ash fractions.Where scale allows, store ash fractions and other dusty residues in closed silos fitted with high level alarms and dust abatement plan
- 4. Explore markets for waste streams, for example:
 - bottom ash for aggregate
 - PFA for cement manufacture and construction products
 - FGD gypsum and fused slags of construction products
- 5. Recycle materials back into the process whenever possible, e.g. re-using partially
- 6. Where recycling or re-use is not possible, then consider regeneration of other materials or return to the manufacturer e.g.:
 - ion exchange resins
 - reverse osmosis membranes
 - molecular sieves
- catalysts

Operations
2.1 Liquid fuels and 2.2 Gaseous facts

Atic

1 Liquid fuels
2.2 Gaseous fuels

Liquid fuels Gaseous fuels

2. Operations

2.1 Liquid fuels

Air: The in-furnace techniques for controlling releases of NOx and particulates are based on burner design, the method of atomization and the control of primary, secondary and tertiary air. A control loop system is required to govern the air and fuel supply and is significant in air pollution control. Such techniques may not be practical for small installations using compact, high thermal rating combustion chambers.

Solids removal from the fue I may be required. Efficient atomisation of oil fuels is necessary and fuel viscosity at the burners is a primary consideration. Oil (including

2.2 Gaseous fuels

These include natural gas and others, e.g. mine gas vaporised LPG, refinery gas and gases from the production of smokeless fuel; together with that from the casification of coal, oil or other carbonaceous matter in a separate plant, or the underground gasification of coal.

ithdrawn on 2 Miles 120, 85 bitumen) emulsions and many heavy fuel oils have high sulphur contents and may have high vanadium and nickel contents. The use of oil fuels containing sulphur will result in some sulphur trioxide releases, as well as sulphur dioxide. Where low sulphur oils (below 1% w/w sulphur) are used, this may preclude the need for flue gas desulphurisation (FGD). The Sulphur Content Of Liquid Fuels Regulations 2000 (SCOLF) allows for derogation from this limit where the Emission Limit Values set out in LCPD are met.

Air: If a plant designed for and normally operated on gaseous fuel not requiring the use of flue gas treatment, has to use standby fuel because of an interruption in the gas supply, then for short periods only such plant may be allowed to operate without flue gas treatment.

Indicative BAT

You should inform us, and the local authority, when standby fuel is used, and when you
return to gas firing.

Emissions and monitoring 3.1 Point source emissions to water 3.2 Point source emissions to air 3.3 Fugitive emissions 3.4 Monitoring

Point source emissions to water 3. Emissions and monitoring

3.1 Point source emissions to water

Generally, waste water can arise from:

- the process
- rain-water run-off where there is the potential for contamination
- storm water
- cooling water
- treatment of accidental releases of raw materials, products or waste materials
- fire-fighting.

At sites with large boilers and furnaces the following sources of waste streams to water are the most significant:

- Coal storage leachate from coal stacks is governed by the coal composition. In particular the presence of pyrite (iron sulpnide) and a low pH can cause pollution. Acidic conditions may increase mobility of trace metals, contributing to the polluting effect of the leachate.
- Cil storage leakage from oil storage a eas can cause pollution.

De-ionisation effluent - water deionisation plant regeneration effluent will usually be produced as an acid stream and an alkali stream, which are mixed together and pH adjusted for disposal. This effluent contains all the dissolved solids in the original water supply and any undissolved solids which may be present, particularly where river or estuary water is used.

Soluble sulphates are also likely to be present where sulphuric acid is used for repeneration of the ion exchange material. The presence of salts in the riease should be considered.

Boiler blowdown - during normal operations boilers are blown down to control the composition of the boiler water. This blowdown is a concentration of the small amounts of solids remaining in the boiler feed water from the water de-ionisation plant, plus any chemicals used for treating the water, e.g. phosphates, small amounts of alkalis, hydrazine, ammonia etc.

- Process water (e.g. wet scrubbing) the effluent will be acidic and be likely to contain metals, solids and organics.
- Cleaning liquids Wash waters and cleaning-out solutions arising from the maintenance of boiler plant may contain organic acids, alkalis, alkali phosphates, iron oxides in suspension, complex corrosion inhibitors, and hydrochloric or hydrofluoric acid.
- Waste water treatment many sites will have on-site wastewater treatment plants for treating domestic wastes and suitable other streams.
- Cooling tower purge Where evaporative cooling towers are used, biocides, most frequently chlorine or bromine based, lead to releases of

Point source emissions to water prescribed substances to both air - water. Timber --

prescribed substances to both air and water. Timber used in cooling towers is usually treated with CCA (copper sulphate, potassium dichromate, arsenic pentoxide), most of which remains well bound to the timber over

its operating life, but initial surface residues can lead to significant levels in the purge water. You should include the requirement for controlled washing at the treatment site in your specification for treated timber.

Indicative BAT

You should where appropriate:

Oil storage

- 1. Fit a high-level alarm to oil tanks.
- 2. Drain decanted water from oil storage tanks and storm water from bunded areas to a water treatment plant, or direct it to an appropriate disposal facility. You should deal with liquid effluents generated during periodic tank cleaning in a similar way.
- 3. Use oil removal facilities such as partition chambers or plate separators for water contaminated with oil.

Coal storage

4. Where there is a significant risk of pollution of water or groundwater, you should collect and treat leachate.

De-ionisation effluent

5. Neutralise water do-ionisation plant regeneration effluent before discharge.

Process water (e.g. wet scrubbing)

- 6. Chemically treat, neutralise and settle the effluent from wet scrubbing before discharge.
- 7. Quantify organics, including dioxins, furans and PAHs in the treated effluent. At low levels they 3 normally most effectively treated in a biological plant, usually by the sewerage undertaker.
- 8. Discharge volumes for sea water scrubbing make most treatment impracticable. Since contaminants are likely to be present in very low concentrations, focus your effort on minimising mass releases where practicable.

Ash handling

9. Handle ashes in a solid state and not as slurry. Hydraulic transport of ashes produces a wastewater stream that is likely to contain metal salts and organic compounds. If there is a market for the ash, you should not use a handling technique that will prevent its re-use.

Cleaning liquids

- 10. Neutralise or treat wash waters and cleaning-out solutions to produce an acceptable waste before discharge or disposal.
- 11. Boiler cleaning wastes require appropriate disposal.

Point source emissions to air All British Point Source emissions to air

- 12. Use an efficient oil/water separation/interceptor system. Further treatment may be required to remove dissolved hydrocarbons.
- 13. Direct discharge to controlled waters will only be allowed where discharges will meet discharge requirements under all conditions.

Waste water treatment

14. On-site wastewater treatment plant effluent must meet discharge standards.

Cooling tower purge

- 15. Optimise the dosing regime for biocides in evalurative cooling towers to minimise their use.
- 16. Chemically treating the main cooling water circuit may allow reduced use of biocides. Do not release accidental overdoses of biccides into the environment.

Thermal plumes

- 17. In terms of the overall energy efficiency of an installation, the use of once-through systems is an appropriate measure. It may be acceptable to use water from a river or an estuary for once-through cooling, provided that:
 - fish can still migrate through the extended heat plume in the receiving water
 - the cooling water intake minimises fish entrainment
 - heat load does not interfere with other users of the receiving surface water.

3.2 Point source emissions to air

For further information concerning the regulatory framework applicable to large coal and oil fired power stations in England and Wales, refer to Controls on Releases of Oxides of Sulphur and Nitrogen from Coal and Oil Fired Power Stations" at:

http://www.environmentagency.gov.uk/business/444304/44436 7/972464/?version=1&lang= e.

NOx control

Primary measures for NOx control

The most important oxides of nitrogen with respect to releases from combustion processes are nitric oxide (NO), nitrogen dioxide (NO₂) (i.e. NOx) and nitrous oxide (N₂O). Nitric oxide forms over 95% of the total NOx in emissions from most types of combustion plant. Higher proportions of NO₂ can be present in exhaust gases from gas turbines particularly modern gas turbines with dry low NOx burner

Point source emissions to air air sperations produce had sperations between the produce had sperations are produced by the produce had sperations are produced by the produce had been produced by the produced b

oxide than other types of combustion plant.

There are three recognised NOx formation mechanisms:

- "fuel NOx" by conversion of chemically bound nitrogen in the fuel
- "thermal NOx " by fixation of nitrogen in the combustion air
- "prompt NOx " by a mechanism in which molecular nitrogen is converted to NO via intermediate products in the early phase of the flame from with hydrocarbons participating reactions.

The first two mechanism are the only ones of major importance in most combustion plants.

Fuel NOx formation depends on the oxygen level in the vicinity of the flame. Reducing oxigen levels reduces fuel NOx. Fuel NOx is most important in coal fired and biomass fired combustion systems, as these fuels have a higher fuel N content.

Thormal NOx formation requires temperatures greater than 1000°C. Reducing peak temperatures reduces thermal NOx formation. The thermal NOx formation route is the most important source of NOx in emissions from oil and gas fired plant.

Baseline NOx emissions from unabated internal combustion engines vary with engine size and speed (RPM). Larger,

NOx than smaller high-speed engines. Similarly NOx emissions from unabated gas turbines are generally a function of engine load, combustor temperature and size.

Applying combustion modifications is limited by operational and fuel specific parameters, that influence the safe operation of plant, such as:

- ignition stability at the burner
- change in flame shape
- problems of corrosion and erosion
- incomplete burnout of the fuel
- increase in CO emissions
- change in flue gas temperature
- changes in the water-steam circuits.

As a result of the likely changes in one or more of these parameters, the suitability and choice of a primary measure and the resulting effects may not be directly transferable from one type of plant to another.

Large boilers and furnaces

Combustion modification techniques include:

NOx control by operational adjustments Reducing the level of excess air can reduce NOx emissions, but carbon monoxide and carbon in ash levels must also be controlled. Particulate emissions may also increase and thermal efficiency may be affected. All new plant should be

Point source emissions to ain who we haracteristics.

In large multiple burner boilers, the upper burners may be used to supply air only. This is not as effective as purpose designed over fire air (OFA) systems. This technique is most suitable for gas and oilfired boilers.

If you preheat your combustion air (up to 300°C), reducing the level of air preheat may reduce NOx emissions but at the expense of a reduction in thermal efficiency.

Low NOx burners

Low NOx burners are effective in educing NOx emissions, but may cause increased emissions of carbon monoxide to air, and a higher level of carbon in the ash. You should use them in all new plant. Retrofitting low NOx burners should be considered for a pant, but sometimes the modifications required make it impossible. Low NOx burners are highly effective for gas firing but there is a tendency to generate particulates when firing heavy fuel cil.

ver-fire air (OFA)

This technique involves introducing air above the primary combustion zone in a boiler. The technique is best used in conjunction with low NOx burners and may be applied to small and large boilers. 15-25% of the total combustion air can be supplied as OFA. The reduction of NOx emissions is variable, depending on boiler

application. The use of this technique may result in increases in carbon monoxide and unburnt carbon emissions. Carbon in ash levels must be controlled.

Flue gas recirculation (FGR)

Recipulation of flue gas dilutes the combustion air, lowering peak flame comperatures and reducing the thermal NOx produced. This technique is useful for gas and oil firing (distillate in particular), but is less effective for coal firing. Particulate emissions may increase with heavy fuel oil. This technique should be considered in the design of boilers. Space requirements may prevent its application to smaller designs. Retrofitting to some types of existing plant may be possible.

Reburn

This technique controls NOx emissions by injecting part of the fuel input at a level above the main combustion zone in the boiler. In this fuel rich region the NOx produced in the main combustion zone is reduced. Up to 20% of fuel may be introduced this way and the fuel need not be the same as the primary fuel. Thus gas or oil might be the reburn fuel for a coalfired boiler. This technique can be applied to new and existing boilers and in combination with low NOx burners. The higher the initial concentration of NOx, the more efficient reburn is at reducing NOx emission levels.

For smaller scale combustion plant, for example plant regulated only as a result of

Point source emissions to air who air who are the use of combined nay be a final and be a final

may be sufficient in the consideration of appropriate measures for the control of oxides of nitrogen emissions.

Pressurised fluidised beds

Pressurised fluidised bed combustion (PFBC) plants are now available up to 400 MW(th). Emissions of NOx are inherently lower than with conventional fuel systems. If required, ammonia can be directly injected (a form of SNCR) into the furnace to reduce NOx further, without using addon back-end catalysts (i.e. SCR). SQx levels are also inherently low due to the use of sorbent (limestone or dolonite), which also helps form the fluidised bed.

Recent developments in pressurized fluidized bed combustion (PFBC) technology include:

Freeboard firing Cinjecting a small quantity of oil into the freeboard at part load increases the plant's efficiency and also maintains the conditions for selective non-catalytic NOx reduction close to those valid at full load conditions.

Wash recirculation – improves SO₂ mission reductions for a given quantity of sorbent feed. Improvements in combustion efficiency can also be achieved. Thermal NOx is not produced. SO₂ emissions are controlled by feeding a sorbent (limestone or dolomite) into the fluidized bed. Elevated pressures in the PFBC improve SO₂ capture.

The use of combined cycle gas turbine (CCGT) plant is the appropriate measure for new gas turbine installations. Dry low NOx combustion is preferred to wet low NOx combustion for new plant. Low emission combustors can be retrofitted to some existing gas turbines, but it may be nore economical to replace others with ew higher-efficiency, lower-emissions models. Additional NOx formed by supplementary firing in the exhaust using a well-designed and adjusted burner can be much less than would be expected by firing in air. This is often used in CHP schemes to vary the ratio of heat to power produced. It has a high incremental thermal efficiency and low environmental impact. Very low NOx emissions are not achieved with auxiliary firing when the gas turbine is not running.

Reciprocating engines

Note that this technology is developing rapidly. You should ensure that you consider the latest developments when planning the installation of new plant.

The principal technique used to reduce NOx emissions is lean burn technology, where the fuel content of the charge is less than stoichiometric. This reduces local temperatures by dilution and ensures there is ample oxygen for good hydrocarbon conversion.

Low NOx emissions from both SI and CI engines can also be achieved by combustion modification. Applying the

Point source emissions to air lead to the echniques listed below is limited by engine design. plant of the last of

fuel parameters. The suitability and choice of a primary measure may not be directly transferable from one engine to another (or between SI – CI engines):

- reduction of charge temperature by the addition of water (CI Engines). This reduces NOx, but is impractical in most circumstances
- tuning for NOx (ignition timing). Reducing the timing angle reduces NOx emissions but it also reduces engine efficiency and increases levels of CO and VOCs
- the use of SCR for gas fuelled SI engines. SI engine exhausts are prone to rapid fluctuations in pressure, which can destroy the SCR catalyst
- fuel/air mixing improvements (CI engines.) This is generally only available on new engines, although multiple discharge injectors can be retrolated to existing engines
- reduction of air manifold temperature. Increased after-cooling may reduce the temperature of the air charged into the cylinders and reduce NOx levels. This technique should be applicable to both existing and new engines
- exhaust Gas Recirculation (EGR). Recycling exhaust gas into the air inlet feeds more inert mixture into the engine and reduces NOx emission.

All the above techniques except lean burn engines have the drawbacks of reduced efficiency and increased emissions of CO and VOCs Effective control of combustion conditions is essential if they are used.

Plasier processes

installations manufacturing plaster from gypsum, the combustion gas stream is intentionally diluted with additional air in order to provide, for example, a transport medium for dried and calcined material and a heat transfer medium, which interacts directly with individual particles. For effective process performance the oxygen content of exhaust gases often approaches ambient levels. For such installations, the excess air should be minimised.

Secondary abatement measures to control NOx

End-of-pipe flue gas technologies to reduce NOx emissions rely on the injection of ammonia, urea or other compounds to react with the NOx in the flue gas and reduce it to molecular nitrogen. They can be divided into:

- selective catalytic reduction (SCR)
- selective non catalytic reduction (SNCR).

Selective catalytic reduction (SCR)

SCR reduces NO and NO₂ to N₂ with the addition of ammonia or urea solution in the

Point source emissions to air we have been been a catalyst at 300-400°C corplant can be operated.

presence of a catalyst at 300-400°C. SCR plant can be operated at various locations in the flue gas stream. It may also be positioned after exhaust gas desulphurisation, although this arrangement would normally require an exhaust gas reheating stage, which may take up as much as 2% of the electric capacity. This measure typically removes between 80 and 90% of the NOx.

SCR is used on combustion plants from <50 MW(th) to 1000 MW(th). The use of SCR for gas turbines can achieve ultra low NOx emissions (<10 mg/m³ at 15% O Catalytic combustion is in commercial development for some types of operations.

The use of SCR requires the storage of liquid ammonia or ammonia solution.

Selective non-catalytic reduction (SNCR)

SNCR reduces NOx emissions by chemically reducing them to nitrogen and

water through the injection of NH₂-X compounds into the furnace. SNCR is operated without a catalyst at a temperature of 850 to 1100°C. The temperature window and residence time can strongly influence the reagent used (ammonia or urea). Ammonia tends to give use to lower nitrous oxide formation but area may be more effective over a signtly larger temperature window and is easier to handle. Abatement efficiencies of up to 80% have been claimed although efficiencies of 30-50% are more typical.

SNCR also requires a sufficient retention time for the injected reagents to react with NO. Reagent distribution/injection must be optimized and computational fluid dynamics modelling may be useful, and essential for all new plant.

Forms of SNCR have been successfully used at lower temperatures in both CFBC and PFBC systems.

Point source emissions to air Albardania Point Source emission emission to air Albardania Point Source emission emission

- Maswill

 - flue/exhaust gas recycling
 - reburn
 - selective catalytic reduction (SCR)
 - selective non catalytic reduction (SNCR)
- 2. Use low NOx burners for coal- and oll-fired plant.
- 3. Use OFA or equivalent for existing coal-fired plant above 100 MWth).
- 4. Use dry low NOx burners in new natural gas-fired gas turbines. For natural gas-fired gas turbines, use water/steam injection, or convert to dry low NOx burning.
- 5. Where air quality standards or other environmental standards must be met, you must use SCR or SNCR for smaller plant (<100 MW).
- 6. For new coal and oil-fired plant above 100MW, use SCR or primary measures to achieve equivalent NOx levels.
- 7. Only combustion optimisation and SCR are feasible on >500MW PF plant firing low volatic coal. In these cases you need SCR for new plant. You need a site specific assessment for existing plant.

SOx control

Fuel Selection

The source of sulphur in emissions to air from combustion processes is the fuel. During combustion the major oxide of sulphur produced is sulphur dioxide (SO₂). Sulphur trioxide is adsorbed and reacts

with ash particles in the flue gases. It is more likely to pose a problem with low ash fuels (e.g. fuel oils) and its presence is an important contributing factor in acid smutting. Measures used to control SOx will also result in lower emissions of hydrogen chloride and hydrogen fluoride.

Point source emissions to air leading to the selection needs to consider a wide range of quality parametrizement.

complex issue.

For smaller scale combustion plant, use of low sulphur fuels (ie less than 1.2 % S) may be sufficient in the consideration of BAT for control of oxides of sulphur emissions.

Coal sulphur content varies widely (range 0.1-3.5%). If coal is the selected fuel, flue gas desulphurisation (or a technique that delivers equivalent emissions) is required Where life or size of the plant precludes FGD as BAT, then low sulphur coals. should be used.

For fuel oils, the Sulphur Content of Liquid Fuels (SCOLF) Regulations set limitations on the sulphur content of liquid fuels. A similar approach should be adopted to that used for coal fired plant.

Natural gas that meets the standard for acceptance into the National Transmission System is considered to be a sulphur free fuel. Natural gas that does not meet this standard, and industrial gases (e.g. some refinery gas and gases from gasification plants) may contain sulphur compounds and may require desulphurisation.

The sulphur content of biofuels can range widely, dependent upon the fuel used. For example straw and barley crops can contain between 0.2 and 0.7% sulphur whilst concentrations in willow are expected to be at round 0.06%

Coal washing is well developed. Most techniques involve crushing coal to fine particle sizes and then separating pyrites and ash content from coal. Depending on the pyrites content, up to 50% of the original coal sulphur may be removed. However, extra energy is consumed, a waste stream is produced and solids waste will require disposal.

For industrial gas fired plant:

Gas cleaning reduces the sulphur content of raw gas streams. Sulphur compounds in the raw fuel gas may be recovered as elemental sulphur by additional treatment. Where gasification is used as an alternative to the direct combustion of coal, oil and oil emulsion fuels, it should be possible to remove about 99% of the sulphur present in the original feedstock.

In-furnace sulphur control

Emissions of sulphur dioxide from combustion plant can be reduced using infurnace injection of sorbents. Fluidised bed combustion (FBC) techniques are suited to this technique. In FBCs, between 10-15% of sulphur is retained in the ash as a result of reaction with calcium carbonate naturally present. Adding crushed sorbent material (e.g. limestone) to the bed may increase sulphur retention in the ash. Wider environmental considerations include the use of limestone and its disposal, and possible uses of the combustor bottom ash and the fly ash streams.

Point source emissions to air with pulverised final form

efficiency of SO₂ removal tends to be lower than with some other wet and dry FGD techniques.

Sulphur dioxide abatement

For large coal-fired plant, end-of-pipe flue gas desulphurisation (FGD), or equivalent techniques are considered appropriate measures. Removal efficiency is "the ratio of sulphur not emitted over a given period to the sulphur in the fuel over the same period"; or, the "Rate of desulphurisation" The given period will be when the FCD plant is 'Available' for use and excludes start up and shutdown periods.

Wet limestone scrubbing

Sulphur dioxide removal is currently about 90-92% at existing FGD plant in the UK (500-600 Mwe), depending on coal sulphur content. The by-product is gypsum. Hygrogen chloride in the flue gas is captured in the scrubber liquor and is purged in the wastewater stream. Sulphur dioxide removal may be increased up to 92-98% (BREF reference), but the buffers used to assist in controlling the chemistry of the process may increase problems with waste-water treatment from scrubbing

This technique represents an appropriate measure for large coal-fired combustion plants above 100 MWth scale and large CI engines using high sulphur residual/fuel

only operates for a limited period this may change the cost benefit equation and make wet lin estone scrubbing uneconomic.

Sea water scrubbing

This technique is considered an appropriate measure for large combustion plants situated on the coast. Up to 98% sulphur dioxide removal is possible. The reheat system may limit efficiency if a bypass is required. Localised effects of the additional sulphate, chloride, heavy metals and elevated temperatures in the discharge must be addressed.

Spray dry scrubbing

This has been used on coal and oil fired systems. 70-95% sulphur dioxide removal is possible. Hot flue gases dry the wet scrubbing medium, which results in a dry residue and an aqueous waste steam is not produced.

Dry sorbent scrubbing

A solid residue is produced which can be used. 80-95% sulphur dioxide removal is possible depending on the process and the quantity of sorbent used. The sorbent may be lime, limestone or other material. A fabric filter is needed to remove alkaline dusts. The filter coated with alkaline dust enhances the scrubbing effect.

Point source emissions to air air four should "' igkamb ou

You should where appropriate:

- 1. Use low sulphur fuels as a primary measure.
- 2. For large coal or oil-fired plant, use wet limestone scrubbing or seawater scrubbing for flue gas desulphurisation (FGD).
- 3. Consider dry sorbent injection for pulverised and liquid fuel furnaces which are too small to justify FGD.
- 4. For fluidised bed combustors, consider in-bed sulphur capture.
- 5. Consider IGCC for new large-scale solid and liquid fuel fired plant.

Particulate matter control

Solid and liquid fuel fired plants en it particulate matter. Gas fired plant using natural gas will not general equire particulate control. Some industrial gases contain particulates and should be filtered on production or, if that is not possible, before combustion

Abating particulate matter, especially the finer fractions, is a significant method of controling the release of heavy metals, dioxins and polynuclear aromatic hydrocarbons (PAHs).

Where particulate abatement is required, the options include:

- electrostatic precipitators (EPs)
- fabric filters
- ceramic filters
- wet scrubbers
- cyclones

For Compression Ignition (CI) engines, particulate removal efficiency is very dependent upon fuel quality. For many CI engine sites particulate abatement may not be required. New CI engines should be capable of achieving the given unabated release levels.

Electrostatic precipitators

Electrostatic precipitators located downstream of FGD can consistently achieve levels below 25 mg/m³. Where FGD is not fitted, levels of 50mg/m³ should be achieved. Electrostatic precipitators are used in both solid and liquid fuelled combustion plants and are available for small and large-scale plant.

Fabric Filters

Bag (fabric) filters when correctly operated and maintained provide reliable abatement to below 10mg/m³ and are likely to be the appropriate measure for many

Point source emissions to air applications. They cannot generally used at temperature

used at temperatures over 250°C.

In oil-fired plant, using a pre-coat of lime or ash on the bags adsorbs any oil carryover. Bags in oil fired plant need to be cleaned less often than with other fuels and fresh coating material must be injected to coincide with cleaning cycles. Using filters in this way must also be done in conjunction with maintaining good combustion control.

prevent thermal damage to bags and blinding by the condensation of moisture

Ceramic filters

Ceramic files are available for small combustion plant and are being developed for larger plant. They can normally achieve reliable levels below 10 mg/m³ and have to v maintenance requirements. Ceramic filters are able to withstand high temperatures around 800-900°C. Operating temperatures are usually limited by the housing rather than the filter media.

Indicative BAT

- 1. For coal and oil-fired plant above 100 MWth, electrostatic precipitators (EP) are required. At smaller scale plant, other methods may be acceptable to meet emission limits.
- 2. For 'opted in' 2 plant FGD is an appropriate measure, and therefore particulate abatement is FGD + EP.
- 3. For large (>150 MW) existing plant and where FGD is not required (not 'opted in') EP is an appropriate measure.
- 4. Where low sulphur fuel is used with EPs, use sulphur trioxide injection to improve particulate control.

25 **Environment Agency** How to comply with your environmental permit Combustion activities (EPR1.01)

ris doci

² 'Opted in' plant have to comply with the requirements of the Large Combustion Plants Directive.

Point source emissions to air leading to be because of the control of CO₂, CO and volces organic come.

organic compounds (VOC)

All measures to reduce fuel use will also reduce CO₂ emissions. Selecting raw materials with low organic matter content and fuel with a low ratio of carbon content to calorific value reduces CO₂ emissions.

Elevated CO and VOC emissions indicate poorly controlled combustion and may also indicate higher releases of other pollutants. Good combustion conditions are required to minimize releases.

Where necessary, the use of catalytic? oxidation in the exhaust gas stream will reduce carbon monoxide emissions to less than 100 mg/m³.

Other releases

Metals and their compounds

The largest proportion of metals and compounds released to air are in the particulate phase, except for mercury and boron, which are released in the vapour phase Controlling particulate levels (see above) and selecting residual fuel oils with a low ash content will control levels of mest metals. Vapour phase mercury and other heavy metals will be partially removed from flue gases during FGD when using a wet scrubbing system.

Halogens

Hydrogen chloride and hydrogen fluoride are produced during the combustion of

abating sulphur dioxide will also reduce these gases. UK coal contains 0.1 - 0.6% chloride and so emissions of hydrogen chloride will be significant from larger unabated combustion plant.

Polycyclic aromatic hydrocarbons (PAHs)

Total PAH (unspecified) emissions for large coal and oil fired combustion plant are of the order of 10-100 ng/m³ in the flue gas. Some PAH is associated with particles, so abating particulates (see above) will reduce emissions of PAHs. although the ash and dust will be contaminated.

Dioxins

Dioxins can potentially be formed by the combustion of any carbon containing fuel in the presence of trace quantities of chloride. Dioxins are usually present in both the particulate and vapour phases and accordingly measures to reduce particulate matter emissions will also significantly reduce emissions of these compounds. FGD systems can enhance dioxin removal as can some SCR systems employed for NOx reduction. These systems should not be relied upon as primary measures to abate dioxins, which should be at concentrations below or close to the limits of current measurement and analysis techniques.

Fugitive emissions

3.3 Fugitive emissions

On many installations fugitive or diffuse emissions may be significant.

Coal, limestone and other solid raw materials may create a dust problem during receipt and subsequent handling and processing operations. Particulate emissions to air should be minimised.

Liquid feedstocks should be stored in roofed tanks. Venting arrangements will depend on the nature of the feedstock, its vapour pressure and composition. The size and number of tanks will depend on whether the oil is the main or standby uel. Alternatively, at larger sites, oil nay be supplied from nearby refinery sites by

pipeline. Oil and oil e mulsion storage facilities require heating plant to reduce the viscosity of the fuels. Releases will include volatile organic compounds (VOC's) to air, oil spills from storage and transfer operations and sludge from tanks.

Gaseous fuels are supplied direct by poeline. Where gas is used as a fuel, local compression facilities may be required in order to inject sufficient fuel. Releases of gaseous fuels can arise from controlled venting during maintenance and purging of the fuel system.

Indicative BAT

- 1. Windbreaks should be created by natural terrain, banks of earth or planting of long grass and evergreen trees in open areas. This has aesthetic benefits and such vegetation is able to capture and absorb dust without suffering long-term harm. Hydro seeding should be used to rapidly establish vegetation on waste tips, slag heaps or other apparently infence ground.
- 2. Where materials are delivered by sea and dust releases could be significant, use self-discharge vessels or enclosed continuous unloaders.
 - Minimise dust generated by grab-type ship unloaders by ensuring adequate moisture content of the material as delivered, minimising drop heights and using water sprays or atomised mist at the mouth of the ship unloader hopper.
- 4. **Fugitive emissions** from fly ash should be prevented by dust suppression, or by enclosing its handling and storage.
- 5. Intercept rainwater run off from open areas, especially coal and raw materials stocking areas, and remove the suspended solids by settlement or other techniques. Where there are potentially vulnerable receptors, monitor the quality of the water discharged from the storage and blending areas.
- 6. Plant used to pre-treat and store raw materials should be totally enclosed, with extraction

Monitoring

on 2410812018 and arrestment plant as appropriate, to prevent emissions to atmosphere. For some gasification processes coal is milled to a very fine dust and needs to be handled with an inert gas blanket.

- 7. Gasifiers should be coal-charged using a double lock system, whereby the gases released from the reactor during charging are contained within the lock hopper. After closure of the charge valve they are routed either to reconverssion for re-injection into the crude gas stream or to a vent treatment system. Attendatively, a wet feed (slurry)
- system may be used with comparable features.

 8. You should demonstrate that the potential risks of contamination of land by deposition of dust, leachate or run-off are not significant and that you can comply with the requirements of the Groundwater Directive

3.4 Monitoring

Indicative BAT

1. Emissions to air

Many plants in this sector will be subject to the detailed monitoring requirements of Annex VIII of the Large Combustion Plant Directive (LCPD). For plants co-incinerating waste, the provisions of Articles 10, 11 and Annexes II and III of the Waste Incineration Directive (WID) should also apply. The monitoring requirements of the LCPD and WID are considered to represent appropriate measures for this sector. The LCPD requires that:

For existing plant using continuous emission monitors the emission limit values shall be regarded as having been complied with if the evaluation of the results indicates, for operating hours within a calendar year, that:

- none of the calendar monthly mean values exceeds the emission limit values; and in the case of:
 - sulphur dioxide and dust: 97% of all the 48 hourly mean values do not exceed 110% of the emission limit values
 - nitrogen oxides: 95% of all the 48 hourly mean values do not exceed 110% of the emission limit values.

For new plants, emission limit values shall be regarded, for operating hours within a calendar year, as complied with if:

no validated daily average value exceeds the relevant figures set out in part B of Annexes III to VII, and

Monitoring

on 2^{A10}812018 95% of all the validated hourly average values over the year do not exceed 200% of the relevant figures set out in part B of Annexes III to VII.

For both existing and new plants, start-up and shut-down periods snall be disregarded.

Installations that do not fall within the scope of WID or LCPD should meet the benchmark standards except where you can clearly demonstrate that this is not the appropriate measure.

We may require you to monitor and report releases more frequently than required by WID or LCPD where it is considered appropriate to do so.

You should process the readouts from continuous emission monitors using software that reports monitoring compliance, to enable direct comparison with the emission limit values specified in relevant European legislation and in this guidance.

In order to relate emission concentrations to mass releases, you will need to measure or otherwise determine the stack gas flow rate. In order to relate measurements to reference conditions, you will need to determine temperature and pressure. Determination of oxygen or water vapour content may also be required. All such measurements should be recorded.

2. Emissions to water and sewer

For combustion plants co-incinerating waste and operating air pollution control equipment with an aqueous discharge, you should comply with Article 8 and Annexes III and IV of the Waste incineration Directive (WID).

Environmental monitoring (beyond installation)

Specific environmental monitoring requirements which may be required are:

air:

Where there is a significant public concern/record of complaint:

- you should keep 24 hour time/date coded CCTV/video recordings of chimney stack emissions
- you should record wind speed and direction data to help with complaint investigation.

- effluent treatment plant discharges to controlled waters
- cooling water discharges

To land:

Monitoring surveys may be needed where sensitive soil systems or terrestrial ecosystems are at risk from deposition of air borne pollutants or direct impacts of any on-site operations.

Monitoring

- 017 2410812018 solid and liquid fuel fired plants - fuel feedstock should be sampled and analysed at a frequency and manner
- potential difference across EP plates

... J fuel the dpl.
should be sample
d at a frequency and
... opriate(b) he type of pla.
concerned

potential difference across EP

reagent injection or feed rates.

Annex 1 Emission benchmarks
Annex 2 Other relevant guidance
and abbreviations
Annex 3 Regulatory information

Annex 1-Emission benchmarks

4. Annexes

Annex 1- Emission benchmarks

indraint of Allos 120/8. The emissions quoted below are daily averages based upon continuous monitoring during the period of operation.

New plants

- No daily average value exceeds the benchmark value.
- 95% of all hourly averages over a defined period, e.g. a calendar year do not exceed 200% of the benchmark value.

Retrofit plant

- No monthly average value (where defined in table 3.1) exceeds the benchmark
- For SO₂ and dust, 97% of the 48 hour average values do not exceed 110% of the benchmark value.
- For NOx, 95% of the 48 hour average values do not exceed 110% of the benchmark value.

Where spot tests are employed:

- It may be necessary to set periodic limits taking account of sampling times practical for the spot test method involved.
- Where the continuous and spot test sampling periods are compatible, the mean of three consecutive tests taken during a calendar year shall not exceed the relevant continuous monitoring benchmark value by more than 10%.

Processes, which come under the provisions of the Large Combustion Plant Directive or the Waste Incineration Directive, will also need to address the specific requirements of these directives.

Annex 1-Emission benchmarks

Min oin 24108/2018 Release values shown are for consolidated but not agglomerated units and NOx releases values under LCPD are 'not opted out' >2000 hr 5 vr rolling average.

Reference conditions for releases to air

The reference conditions of substances in releases to air from point-sources are:

For combustion gases:

dry

temperature 273K (0°C)

pressure 101.3kPa (atmosphere)

oxygen content

- 3% y/v, dry for liquid and gaseous fuels burned at large boilers
- 15% //v, dry for liquid and gaseous fuels burned at CI engines and
- 6% v/v, dry for solid fuels burned at large boilers and furnaces.
- or non-combustion gases:

pressure 273K (0°C)

pressure 101.3kPa

no correction for water vapour or oxygen content.

To convert measured values to reference conditions, see Monitoring Guidance³.

33 **Environment Agency** Combustion activities (EPR1.01) How to comply with your environmental permit

³ Environment Agency Technical Guidance Notes M1 and M2 provide extensive guidance on the monitoring of stack emissions to air. The conversion referred to is given in TGN M2.

Annex 1-Emission benchmarks Table 1.1 - Summary of achievable releases to air (normal font- daily averages, underlined font- monthly averages).

Note 1 where FGD used

Note 2 Low sulphur coal will produce typically 100mg/m³ per 0.1% Cl in coal. Dust Sorbent Injection will reduce this by up to 90%

Note 3 Where straw or other higher Cl containing biomass material are used HCl concentrations may be up to 300mg/m³

Note 4 Upper end of range applies where higher moisture content biomass is BAT.

Fuels and	Size MWth	Process	Ref	Achievable concentrations, mgm ³ , dry at 0°C, 101.3kPa (at ref O ₂)					
processes			%						
		40	O ₂	PM	SO ₂	NOx	N ₂ O	СО	HCI
Coal	50-100	New Grates/PC	6	20	400	300	10	150	10 Note 1,2
		New CFBC and PFBC		20	400	300	30-120	150	30
		New BFBC		20	400	300	30-120	100	5
		Retrofit PC	6	100	1300	<u>600</u>	2-10	400	400 Note 2
		Retrofit CFBC and PFBC		100	1300	600	30-120	150	30
		Retrofit BFBC		100	1300	600	30-120	100	1 - 5
·	100-300	New PC	6	20	200	200	2-10	150	10 Note 1,2

Annexes	\$							0,	
								A series and a series of the s	
							10		
							`\ <i>\</i>		
							•		
						9			
		<u> </u>							
	nev 1	-Emission h	മ	near	naxx				
					116/01/				
				.0	S				
		T	1		,				
		New CFBC and PFBC		20	200	200	30-120	150	30
		New CFBC and PFBC New BFBC		20	200	200	30-120 30-120	150	30
		New CFBC and PFBC New BFBC Retrofit PC	a a	20	200 200 400	200 200 600	30-120 30-120 2-10	150 100 400	30 1-5 400 Note
		New CFBC and PFBC New BFBC Retrofit PC Retrofit CFBC and PFBC	0	20 20 100 100	200 200 400 400	200 200 600 600	30-120 30-120 2-10 30-120	150 100 400 100	30 1-5 400 Note
		New CFBC and PFBC New BFBC Retrofit PC Retrofit CFBC and PFBC Retrofit BFBC	le o	20 20 100 100 100	200 200 400 400 400	200 200 600 600 600	30-120 30-120 2-10 30-120 30-120	150 100 400 100	30 1-5 400 Note 30 1-5
		New CFBC and PFBC New BFBC Retrofit PC Retrofit CFBC and PFBC Retrofit BFBC New PC	1						
	>300	New PC	6	10	150	150	2-10	150	10
		New PC New CFBC and PFBO	1	10 10	150 200	150 200	2-10 30-120	150 150	10 30
		New PC New CFBC and PFBC New BFBC	6	10 10 10	150 200 200	150 200 150	2-10 30-120 30-120	150 150 100	10
		New PC New CFBC and PFEC New BFBC Retrofit PC with FGD & OFA	1	10 10 10 25	150 200 200 400	150 200 150 500	2-10 30-120 30-120 2-10	150 150 100 400	10 30 1-5 0 Note 1
		New PC New CFBC and PFBC New BFBC Retrofit PC with FGD & OFAwith OFA	6	10 10 10 25 50	150 200 200 400 2,000	150 200 150 500 500	2-10 30-120 30-120 2-10 2-10	150 150 100 400 400	10 30 1-5 0 Note 1 400 Note
		New PC New CFBC and PFBC New BFBC Retrofit PC with FGD & OFAwith OFA Retrofit CFBC and PFBC	6	10 10 10 25 50 100-50	150 200 200 400 2,000 100-200	150 200 150 500 500 500	2-10 30-120 30-120 2-10 2-10 30-120	150 150 100 400 400 100	10 30 1-5 0 Note 1 400 Note 30
		New PC New CFBC and PFBC New BFBC Retrofit PC with FGD & OFAwith OFA	6	10 10 10 25 50	150 200 200 400 2,000	150 200 150 500 500	2-10 30-120 30-120 2-10 2-10	150 150 100 400 400	10 30 1-5 0 Note 1 400 Note

								-90	
								1100	
								J.X.	
						(0)			
Λ nn ϵ	5V 4_[Emissia			chm				
	3X 11-1	=11112210) GII		dinio			
		800000000000000000000000000000000000000			S	5566666666666	5666666666		
F-1.1. 4 4	. 4 ! l				7.0.				
1 2 1 1 CO	ATINIIAA				~ 1.3				
able 1.1 col	Size MWth	Process	Ref	Achiel	able conce	ntrations, mo	n/m³. drv a	t 0°C. 101.3kPa	(at
Tuels and processes	Size MWth	Process	Ref	Achie	able conce	ntrations, mg	g/m³, dry a	t 0°C, 101.3kPa	(at
able 1.1 col Fuels and processes	Size MWth	Process	Ref % O ₂	Achie ref ()	SO ₂	ntrations, mg	g/m³, dry a N₂O	t 0°C, 101.3kPa	(at
Fuels and processes Biomass	Size MWth	Process New PF/Grates	Ref % O ₂	Achie ref () ₂) PM	SO ₂	NOx 300	g/m³, dry a	t 0°C, 101.3kPa CO 100-250 Note 4	(at HCI 25 Note 3
Tuels and processes Biomass	Size MWth	Process New PF/Grates New FBC	Ref % O2	Achie ref ().) PM	(a) le conce SO ₂ 300	NOx 300 250	g/m³, dry a N₂O	CO 100-250 Note 4 100-150 Note 4	(at HCI 25 Note 3
Tuels and processes Biomass	Size MWth 50-100	Process New PF/Grates New FBC Retrofit PF/Grates	O.	Achie ref () ₂) PM 20	(a) ble conce SO ₂ 300 300	NOx 300 250 300	g/m³, dry a	CO 100-250 Note 4 100-250 Note 4 100-250 Note 4	(at HCI 25 Note 3 25 Note 3
Fuels and processes Biomass	Size MWth 50-100	New FBC	O.			250	g/m³, dry a	100-150 Note 4	
Guels and processes Biomass	Size MWth 50-100	New FBC Retrofit PF/Grates	O.			250	g/m³, dry a N₂O	100-150 Note 4	
Fuels and processes Biomass	Size MWth 50-100	New FBC Retrofit PF/Grates Retrofit FBC	6	50	300	300	g/m³, dry a	100-150 Note 4 100-250 Note 4	25 Note 3
able 1.1 colling and processes Biomass	Size MWth 50-100	New FBC Retrofit PF/Grates Retrofit FBC Retrofit PF/Grates	6	50	300	300	g/m³, dry a	100-150 Note 4 100-250 Note 4 100-250 Note 4 100-150 Note 4	25 Note 3
Tuels and processes Biomass		Retrofit PF/Grates Retrofit FBC Retrofit PF/Grates Retrofit FBC New PF/Grates	6	50	300	300 300	g/m³, dry a	100-150 Note 4 100-250 Note 4 100-250 Note 4 100-150 Note 4 100-250 Note 4	25 Note 3 25 Note 3
Tuels and processes Biomass		New FBC Retrofit PF/Grates Retrofit FBC Retrofit FBC Retrofit FBC New PF/Grates New FBC	6 6 6	50 50 20	300 300 200	300 300 300 150	g/m³, dry a	100-150 Note 4 100-250 Note 4 100-250 Note 4 100-150 Note 4 100-250 Note 4 100-150 Note 4	25 Note 3 25 Note 3
	>300	Retrofit PF/Grates Retrofit FBC Retrofit FBC Retrofit FBC New PF/Grates New FBC Retrofit PF/Grates	6	50	300	300 300	g/m³, dry a	100-150 Note 4 100-250 Note 4 100-250 Note 4 100-150 Note 4 100-250 Note 4 100-150 Note 4 100-250 Note 4	25 Note 3 25 Note 3 25 Note 3
		Retrofit PF/Grates Retrofit FBC Retrofit FBC Retrofit FBC New PF/Grates New FBC Retrofit PF/Grates	6 6 6	50 50 20	300 300 200	300 300 300 150	g/m³, dry a	100-150 Note 4 100-250 Note 4 100-250 Note 4 100-150 Note 4 100-250 Note 4 100-150 Note 4	25 Note 3 25 Note 3 25 Note 3

Table 1.1 continued Fuels and processes Ref Achie (able concentrations, mg/m³, dry (at ref O₂) O₂ PM SO₂ NOx N₂d		08/2018
Table 1.1 continued Fuels and processes Size MWth Process Ref (at ref O ₂) Achie (able concentrations, mg/m³, dr. (at ref O ₂) Do2 PM: SO2 NOx N2 Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 des5 450 des5	57 ²	
Table 1.1 continued Fuels and processes Size MWth Process Ref (at ref O₂) Achie (able concentrations, mg/m³, dr (at ref O₂) Do2 PM SO₂ NOx N₂th Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 (685) 450 (685)	\$	
Table 1.1 continued Fuels and processes Size MWth process Process (at ref O₂) Ref (at ref O₂) Achie (able concentrations, mg/m³, dr (at ref O₂) Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 (685) 450 (685)		
Table 1.1 continued Fuels and processes Size MWth Process Ref (at ref O2) Achie (able concentrations, mg/m³, dr (at ref O2) Do2 PM SO2 NOx N2t Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 450 (685) 450 (685)		
Table 1.1 continued Fuels and processes Size MWth Process Ref (at ref O ₂) Achie (able concentrations, mg/m³, dr. (at ref O ₂) Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 450 (885) 450 (885)		
Fuels and processes Size MWth Process Ref (at ref 02) Achie (a) le concentrations, mg/m³, dry (at ref 02) PM: SO2 NOx N2 Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 685 450 685		
Fuels and processes Size MWth process Process Ref (at ref O ₂) Achie (able concentrations, mg/m³, dry (at ref O ₂) PM SO ₂ NOx N ₂ (at ref O ₂) Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 (685) 450 (685)		
Fuels and processes Size MWth process Process Ref (at ref O ₂) Achie (able concentrations, mg/m³, dry (at ref O ₂) PM SO ₂ NOx N ₂ (at ref O ₂) Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 (685) 450 (685)		
Liquid Fuels 50-100 New 3 15 350 (175 post 2008) 300 post 2008) Retrofit 1%S 1%S +DSI 3 50 1700 685 450 685	y at 0°C, 10	101.3kPa
Liquid Fuels 50-100 New 3 15 350 (175 300 post 2008) Retrofit 1%S 3 50 1700 450 685	о со	HCI
Retrofit 1%S 3 50 1700 450 1%S +DSI 685	150	1-5
	30-50	0 1-5
100- New 3 13 200 130	150	1-5
300	150	1-5
Retrofit 1%S 3 <u>50</u> 1700 <u>450</u>	30-50	0 1-5
1%3 +DSI 400		
>300 Yew 3 15 150 100 Retrofit opt out 3 50 1700 1400	150	1-5

Annex 1-							. 90.	
4 Annexes						(200	
						28	V	
						OAN		
						-0'N		
					.(/O,		
					100			
Anney 1	Emissio	n k	ക്രദ	hma	wise.			
AHIIGA 17			/U11U		(avo			
				100				
	Opt in		\	<u> 400</u>	450-400			
			and		(300- 500MW)			
	Engines CI	15	23 0	112	150	150		
		90	Ť	66 from 2008				
	Gas Turbines	15		112	<u>125</u>	100		
	· · ·			(66 from 2008)				
	0							

38 Combustion activities (EPR1.01) How to comply with your environmental permit

4 Annexes							. N.C	3120	
Anne	x 1	-Emission	bei	nchr	nąd	Grawn.	5 5.		
Table 1.1 con	tinued			' MS	5				
Table 1.1 con Fuels and processes	tinued Size	Process	Ref	Achieva ref O₂)	able conce	ntrations, mg	/m³, dry at 0	°C, 101.3kPa	ı (at
Table 1.1 con Fuels and processes	tinued Size	Process	Ref % Q	Achieva ref O ₂) PM	able conce	ntrations, mg	/m³, dry at 0 N₂O	°C, 101.3kPa	ı (at HCI
Table 1.1 con Fuels and processes Natural gas	tinued Size	Process New Gas Turbines (incl CCGT)	Ref % (7) 02) 7 15	Achieva ref O ₂) PM	SO ₂	ntrations, mg NOx 20-50	/m³, dry at 0 N ₂ O	°C, 101.3kPa	ı (at
Table 1.1 con Fuels and processes Natural gas	tinued Size	(incl CCGT) Existing GTs(inc CCGT)DLN	Ref % Q 15	PM 5	SO ₂ 10	ntrations, mg NOx 20-50 75 110	/m³, dry at 0	°C, 101.3kPa CO 100	ı (at
Table 1.1 con Fuels and processes Natural gas	tinued Size	(incl CCGT) Existing GTs(inc CCGT)DLN Water or Steam injection Engines Cl(dual Fuel)	>			75	/m³, dry at 0		HCI
Table 1.1 con Fuels and processes Natural gas	tinued	(incl CCGT) Existing GTs(inc CCGT)DLN Water or Steam injection	15	5	10	75 110	/m³, dry at 0	100	HCI

39 Combustion activities (EPR1.01) How to comply with your environmental permit

					NO'	
4 Annexes					00,	
					100	
				ŋ	X V	
				20'		
				Ο).		
				0		
Anno	v 1 Emission	hone	ah mar	dian		
Anne	x 1-Emission	bend	chma	KS		
Anne	x 1-Emission	bend	chmar	Ks .		
Anne	x 1-Emission	bend	chmar	KS		
Anne	x 1-Emission Engines SI Lean Burn	bend	chmar	250	400	
Anne	x 1-Emission Engines SI Lean Burn	bend	chmat Nas	250	400 1000	
Anne	X 1-Emission Engines SI Lean Burn pre 2005	bend	chmar	250 500	400 1000	
Anne	X 1-Emission Engines SI Lean Burn pre 2005 New Boilers	bend	chmate	250 500 150/100	400 1000 30-50	
Anne	X 1-Emission Engines SI Lean Burn pre 2005 New Boilers (<100MW / >100MW)	bend	chmat Syncs	250 500 150/100	400 1000 30-50	
Anne	Engines SI Lean Burn pre 2005 New Boilers (<100MW / >100MW) Existing Boilers	5 5 3 5	chmate 5 10	250 500 150/100 300/200	400 1000 30-50 50	

40 Combustion activities (EPR1.01)

							108/	6,	
Anne	x 1-	-Emission Process All combustion plant	ı be	nchm	and som				
Table 1.1 conti	nued Size	Process	Ref	Achievable co	ncentrations m	g/m³ dry	at 0°C 101	1 3kPa (at	
Table 1.1 conti Fuels and processes	nued Size	Process	Ref %	Achievable co	ncentrations, m	g/m³, dry	at 0°C, 10 ⁴	1.3kPa (at	
Table 1.1 conti Fuels and processes Industrial gas	nued Size	Process All combustion plant	Ref % 03 30r 15	Achievable co ver O ₂) PM Appropriate measure is gas cleaning to remove fuel S and fuel PM	ncentrations, m SO₂ Appropriate measure is gas cleaning to remove fuel S and fuel PM	g/m³, dry NOx	at 0°C, 10′ N₂O	1.3kPa (at	HCI
		Process All combustion plant	Ref % C 3 or 15	to remove fuel	to remove fuel	g/m³, dry NOx	at 0°C, 101 N₂O	1.3kPa (at	HCI
Smokeless Fue Combustor/Inci Battery/Boiler F	ls nerator/	Process All combustion plant	Ref % Q 3 or 15	to remove fuel	to remove fuel	g/m³, dry NOx 300	at 0°C, 10′	1.3kPa (at CO	HCI
Table 1.1 continuity Fuels and processes Industrial gas Smokeless Fuel Combustor/Incity Battery/Boiler For Gases Dryer Vents	ls nerator/	Process All combustion plant	Ref % Q 3 or 15	S and fuel PM	to remove fuel		at 0°C, 101		HCI

41 Combustion activities (EPR1.01)

4 Annexes						00	
					c	8/	
						9	
					\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
					0,,		
					NO		
				0			
Annov		iaa ka	مطمم	0			
7 AN WOLLAND DI AND W AND WAY CONDINON			2				
Alliex	-EMISS			ICK NA			
Aillex	I-EMISS	NOU DE		May 2			
Annex	I-EMISS	NOIT DE	ر از	JUNO S			
Oryer Vents	-EMISS		25 10	ichino M	300		
Dryer Vents Screen Abatement	r-Emiss		25 70		300		
Dryer Vents Screen Abatement Coal/oil liquor	r-Emiss	SIUII DE	25 25 25 25		300		
Dryer Vents Screen Abatement Coal/oil liquor	r-Emiss	SIUIT DE	25 NO 25 25		300		
Dryer Vents Screen Abatement Coal/oil liquor handling Carbon black		NOIT DE	25 25 25 25	3	300		
		TOTT DE	25 25 25 25	2300	500	150	
gases Dryer Flue		1011 DE	25 25 25 25 25	2300	300 500 500	150	
gases Dryer Flue Gases Process/Purge/	0	O I					
Annexes Annexes Dryer Vents Screen Abatement Coal/oil liquor handling Carbon black Boiler/CHP flue gases Dryer Flue Gases Process/Purge/ dust filters Other Processes		10	25	2300			

4 Annexes Anne	x 1	-Emissi	on k	oenchr	naws	or Akios	120	V	
Table 1.1 con	tinued	-Emissi		No	S MI				
Table 1.1 con Fuels and processes	tinued Size	Process	Ref	Achievable con	centrations, mg/m³	, dry at 0°C, 101.3	kPa (at		
Fuels and	inuea					, dry at 0°C, 101.3	kPa (at	со	нсі
Fuels and	inuea		Ref	Achievable con- ref O ₂) PM Appropriate measure is gas	SO ₂ Appropriate measure is	NOx Use of low NOx technology in	T	1	HCI
Fuels and processes	inuea	Process	Ref % O ₂	Achievable con-	SO ₂ Appropriate measure is desulphurisation to remove S	NOx Use of low NOx	T	СО	HCI
Fuels and processes	inuea	Process Liquid and gas	Ref % O ₂ 3% v/\ 6%	Achievable con- ref (2) PM Appropriate measure is gas cleaning to remove fuel S	SO ₂ Appropriate measure is desulphurisation	NOx Use of low NOx technology in heating burners	T	СО	HCI

General Note Releases values shown are for consolidated but not agglomerated units and NOx releases values under LCPD are 'not opted out' >2000 hr 5 yr olling average.

> 43 **Environment Agency** How to comply with your environmental permit Combustion activities (EPR1.01)

Annex 2- Other relevant guidance and abbreviations For a full list of available Technical Guidance and other relevant guidance and other guidance and other guidance and guidanc

In addition to the guidance in GTBR the following guidance's relevant to this sector:

- 1. The Categorisation of Volatile Organic Compounds, 1995 HMIP Report No. DOE/HMIP/RR/95/009
- 2. BREF on Waste Water and Waste Gar Treatment Industry http://www.jrc.es/pub/english.cgi/0/733169 or http://eippcb.jrc.es
- 3. LFTGN 6 Guidance on gas treatment technologies for Landfill Gas Engines http://www.environment-agency.gov.uk/commondata/acrobat/lftgn06_treatment_936556.pdf

Abbreviations

BAT Best Available Techniques

BOD Biochemical Oxygen Demand

BREF BAT Reference Document

CEM Continuous Emissions Monitoring

CHP Combined heat and power plant

COD Chemical Oxygen Demand

FLY Émission Limit Value

EMS Environmental Management System

EQS Environmental Quality Standard

ETP Effluent treatment plant

FOG Fat Oil Grease

ITEQ International Toxicity Equivalents

MCERTS Monitoring Certification Scheme

NIEHS Northern Ireland Environment and Heritage Service

SAC Special Areas of Conservation

SECp Specific Energy consumption

SEPA Scottish Environment Protection Agency

SPA Special Protection Area

TSS Suspended solids

TOC Total Organic Carbon

VOC Volatile organic compounds

44 **Environment Agency** How to comply with your environmental permit Combustion activities (EPR1.01)

4 Annexes						(120)	
						1100	
						V	
					'() ()		
					100		
Anna	v 2_Doc	telur	on/ ir	oformati	400		
	V O I V C C	Julau					
			ď	1185			
	egulatory inform			9/1/32			
Annex 3- Re	egulatory inform 3.2 provide informatio	nation on regarding	which part of t	he rogulations apply t	o your plant.		
Annex 3- Re Tables 3.1 and 3 Note that this table	egulatory inform 3.2 provide informatio e is not applicable in So	nation on regarding cotland. Pleas	which part of the	he rogulations apply to	o your plant. plant regulatory issues for	· Scotland	
Annex 3- Re Tables 3.1 and 3 Note that this table	egulatory inform 3.2 provide informatio e is not applicable in So	nation on regarding cotland. Pleas	which part of the	he rogulations apply t	o your plant. plant regulatory issues for	Scotland	
Annex 3- Re Tables 3.1 and 3 Note that this table Table 3.1 Combu	egulatory inform 3.2 provide informatio e is not applicable in So	nation on regarding cotland. Pleas us combustic	which part of the contact SEPA	he regulations apply to discuss combustion where gas is produce	o your plant. plant regulatory issues for ed) plus combustion	T	
Annex 3- Re Tables 3.1 and 3 Note that this table Table 3.1 Combu Material Processed Fuel not comprising Waste e.g.	egulatory inform 3.2 provide informatio e is not applicable in So	nation on regarding cotland. Pleas us combustic <0.4MWth	which part of the contact SEPA	he regulations apply to discuss combustion where gas is produce	o your plant. plant regulatory issues for ed) plus combustion	T	WID DOES NOT APPLY
Annex 3- Re Tables 3.1 and 3 Note that this table	egulatory inform 3.2 provide informatio e is not applicable in So	nation on regarding cotland. Pleas us combustic <0.4MWth See note 1 Not regulated	which part of the contact SEPA on or pyrolysis 0.4 - 31/1/4/1	he regulations apply to discuss combustion where gas is produce 3 - 20MWth	o your plant. plant regulatory issues for d) plus combustion 20 - 50MWth	>50MWth	

4 Annexes						24/08/12	
Anne	x 3-Re	gulat	ory ir		ight of		
				105			
waste	purpose is destruction of waste)	5.1A(2)(a) if < 1t/hr	5.1A(2)(a) if < 1t/hr	5.1A(2)(a) if < 1t/hr		< 1t/hr	
waste Non hazardous waste	destruction of				1.1A(1)(b)(unless part of a part A(2) activity)	< 1t/hr 1.1A(1)(a)	
Non hazardous	destruction of waste) Co-incineration (Primary purpose is generation of) if < 1t/hr	< 1t/hr	5.1A(2)(a) if < 1t/hr 1.1A(1)(b)(unless part of a part			

Note:1 WID Guidance(ref) excludes certain small waste burners and space heaters or other waste oil burners. In these cases 1.1 B (b) or (c) may apply dependent on individual or aggregated capacity

1.2A(1)(j) 1.2A(1)(i) WID DOES NOT **Material not** 1.2A(1)(j) 1.2A(1)(i) 1.2A(1)(j) **APPLY** comprising Waste e.g. coppiced wood **WID Exempt** 1.2A(1)(j) 1.1A(1)(b)(unless 1.1A(1)(b)(unless 1.1A(1)(a) waste e.g. part of a part A(2) part of a part A(2) or biomass or B activity) B activity) (Primary purpose is 5.1A(1)(c) if 5.1A(1)(c) if >= 5.1A(1)(c) if >= 1t/hr5.1A(1)(c) Non 5.1A(1)(c) if >= WID DOES destruction of waste) **APPLY** 1t/hr 5.1A(2)(a) 5.1A(2)(a) if < 1t/hrhazardous >= 1t/hr 1t/hr 5.1A(2)(a) if <if >= 1t/hr5.1A(2)(a) if 5.1A(2)(a) waste if < 1t/hr1t/hr if < 1t/hrIncineration < 1t/hr

4 Annexes					ŋ	410812)
Anne	ex 3-Regu	ulator	v info	rmatio			
			<i>y</i>				
				705			
nazardous vaste Co-	(Primary purpose is generation of energy)	5.1A(2)(b)	5.1A(2)(b)	1.1A(1)(b)(unless part of a part A(2)activity)	1.1A(1)(b)(unless part of a part A(2)activity)	1.1A(1)(a)	
nazardous waste Co- ncineration Non nazardous		5.1A(2)(b) 1.2A(1)(j)	5.1A(2)(b) 1.2A(1,1)	1.1A(1)(b)(unless	part of a part	1.1A(1)(a) 1.2A(1)(j)	WID DOES NOT APPLY
nazardous waste Co- ncineration Non nazardous waste	generation of energy) Material products- No incineration of any products locally or		315	1.1A(1)(b)(unless part of a part A(2)activity)	part of a part A(2)activity)		
Non hazardous waste Co- incineration Non hazardous waste Hazardous waste Hazardous waste	generation of energy) Material products- No incineration of any products locally or remotely Incineration (Primary purpose is destruction	1.2A(1)(j)	1.2A(1)0	1.1A(1)(b)(unless part of a part A(2)activity) 1.2A(1)(j)	part of a part A(2)activity) 1.2A(1)(j)	1.2A(1)(j)	APPLY WID DOES

email
enquiries@environment-agency.gov

or visit our websitend was agency.gov

ncir

incident botline 0800 80 70 60 (24hrs) floodline 0845 988 1188