

Suomen Soodakattilayhdistys ry

HAJUKAASUJEN POLTTOSUOSITUKSEN PÄIVITYS

PROJEKTIKOKOUS 1/2014

AIKA 28.5.2014 klo 16.00 – 19.00

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LIITE 1 Hajukaasusuosituksen englanninkielinen käännös 28.5.2014

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24.6.2014

2 (4)

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2 HAJUKAASUJEN POLTTOSUOSITUKSEN PÄIVITYS

2.1 Projektin tavoite

Päivittää suositus viime vuosina toteutettujen sellutehdasprojektien kokemusten perusteella ja ehdottaa missä laajuudessa suositus käsittelee myös hajukaasukeräilyjärjestelmää.

2.2 Työsuunnitelma ja tehtävänjako

Jokainen lukee tahollaan suosituksen läpi ja merkitsee kommentit/päivitystä vaativat kohdat. Kokouksessa kommenttien läpikäynti, keskustelu ja tekstin päivittäminen.

Lisäksi työryhmän laitetoimittajajäsenet käyvät läpi suosituksen laimeiden/väkevien/hönkien kapasiteettirajoitukset (milloin hajukaasun polton voi aloittaa) ja selvittävät voidaanko nykyisiä rajoja perustellusti alentaa.

Työryhmän tehdashenkilöt kokoavat tahollaan listan kokemukseen perustuvista asioista jotka tulisi huomioida laimeiden/väkevien/hönkien keräilyn sekä polton osalta ylös/alasajossa ja seisokkitilanteissa.

3 PÄIVITYKSEN TILANNE

Suomenkielinen suositus julkaistu 29.10.2013. Kokouksessa käytiin läpi englanninkieliseen käännökseen saadut kommentit, LIITE 1.

Suositus vaati vielä läpikäynnin koska käännöksen tehnyt henkilö ei ole soodakattila-alan asiantuntija, käännöksessä on käytetty eri termejä ja lauseita kuin mitä normaalisti alalla käytetään.

3.1 Standardi 12952-8

Standardissa 12952-8 “Water-tube boilers and auxiliary installations. Part 8: Requirements for firing systems for liquid and gaseous fuels for the boiler” todetaan että:



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- A.3.3.1 Introduction of any toxic or odorous gas or gas mixture into the furnace shall not be allowed unless at least 50 % of the rated capacity of the boiler is established by means of black liquor firing.

Tämän asian muutos täytyy laittaa vireille suosituksen julkaisun jälkeen.

4 AIKATAULU

Työ julkaistaan ulkoasun läpikäynnin jälkeen (sihteeri).

5 MUUT ASIAT

LIITE 1

**Englanninkielinen käännös hajukaasusuosituksesta
28.5.2014**



Finnish Recovery Boiler Committee

**Recommended procedure for
incineration of non-condensable gases**

Original version, May 30, 2002

Revision A, December 16, 2005

Revision B, March 27, 2014

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PREFACE

This recommendation issued by the Finnish Recovery Boiler Committee is the second updated version of the original recommendation of May 30, 2002. The recommendation has been updated based on the experiences gained in pulp mill projects during the last few years. The updating work started in late 2010 and was finished in the spring of 2013.

Responsible for preparing this revision B has been the working group established by the Finnish Recovery Boiler Committee, with the following members: Marja Heinola and Risto Honkanen of Andritz Oy, Kari Haaga and Tuomo Hilli of Metso Power Oy, Ismo Tapalinen of UPM-Kymmene Kaukas Mill, Lauri Mattila of UPM-Kymmene Oyj Pietarsaari Mill, Raine Rantanen of UPM-Kymmene Kymi Mill, Esa Vakkilainen of Lappeenranta University of Technology and with Markus Nieminen Pöyry Finland Oy as secretary. Group wishes to thank Reijo Hukkanen of Stora Enso Oy for his numerous expert comments.

Responsible for preparing revision A was the Finnish Recovery Boiler Committee's Environmental Working Group, with Pekka Posti of Oy Metsä-Botnia Ab as chairman, with Sebastian Kankkonen of Jaakko Pöyry Oy as secretary, with Aimo Hakkarainen of Andritz-Ahlstrom Oy, Jouni Hiltunen of Stora Enso Fine Paper Oy, Markku Isoniemi of Kvaerner Pulp Oy, Kari Parviainen of Jaakko Pöyry Oy, Matti Tikka of Kymi Paper Oy, Esko Talka of the Finnish Pulp and Paper Research Institute KCL, and Juha Tolvanen of Alstom Power Finland Oy as members and Esa Vakkilainen of Jaakko Pöyry Oy as expert adviser.

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Finland Oy as members, and with Esa Vakkilainen of Jaakko Pöyry Oy as expert adviser.

The recommendation does not attempt to present a uniform system for handling non-condensable gases nor promote the use of uniform equipment or process designs.

Instead, the recommendation provides basic information for use in engineering, manufacture and operation. As the intention is to develop the recommendation further, we ask you send your possible comments on mistakes, improvement proposals and experiences to the Secretariat of the Finnish Recovery Boiler Committee. Our contact information is found on the Committee's web site: <http://www.soodakattilayhdistys.fi>.

The Finnish Recovery Boiler Committee assumes no responsibility for errors in this recommendation nor for problems that might arise through use of this document. The possible updated document is available for members on the Committee's web site or through the Secretariat.

The Finnish Recovery Boiler Committee

CONTENTS

1	GENERAL	1
2	DEFINITIONS AND ABBREVIATIONS	3
2.1	DILUTE NON-CONDENSIBLE GASES	3
2.2	DISSOLVING TANK AND MIXING TANK VENT GASES	4
2.3	CONCENTRATED NON-CONDENSIBLE GAS	4
2.4	DEFINITIONS.....	5
2.5	ABBREVIATIONS	<u>65</u>
2.6	EXPLOSION LIMIT FORMULAS.....	6
2.7	HYDROGEN SULPHIDE	7
2.8	METHYL MERCAPTAN	8
2.9	DIMETHYLE SULPHIDE/DIMETHYLE DISULPHIDE.....	8
2.10	METHANOL	8
2.11	TURPENTINE.....	<u>98</u>
3	IMPACTS ON RECOVERY BOILER EMISSIONS.....	11
3.1	IMPACTS ON RECOVERY BOILER SULPHUR EMISSIONS.....	11
3.2	IMPACTS ON RECOVERY BOILER NOX EMISSIONS.....	11
3.2.1	IMPACT OF DNCG	<u>124</u>
3.2.2	IMPACT OF DISSOLVING TANK VENT GASES	12
3.2.3	IMPACT OF CNCG	<u>134</u>
3.2.4	IMPACT OF METHANOL	<u>134</u>
3.3	IMPACTS ON RECOVERY BOILER TRS EMISSIONS.....	13
3.4	IMPACTS ON RECOVERY BOILER FOULING.....	<u>1413</u>
3.5	IMPACTS ON OTHER RECOVERY BOILER FUNCTIONS	<u>1413</u>
4	RISKS AND CORROSION RELATED TO NON-CONDENSIBLE GAS SYSTEMS	
	<u>1615</u>	
4.1	LEAKAGE OF NON-CONDENSIBLE GASES INTO MANNED AREAS.....	<u>1615</u>
4.1.1	ODOUR.....	<u>1746</u>
4.1.2	SAFETY EQUIPMENT	<u>1746</u>
4.2	CONDENSATE DRAINING PROBLEMS	<u>1716</u>
4.2.1	OVERFLOWING OF CONDENSATE TRAP	<u>1847</u>
4.2.2	DRYING OUT OF WATER SEALS.....	<u>1847</u>
4.2.3	PLUGGING OF WATER SEALS	<u>1847</u>
4.3	EXPLOSION RISKS IN NON-CONDENSIBLE GAS LINES	<u>1817</u>
4.3.1	SHUT-DOWN AND START-UP SITUATIONS.....	<u>1948</u>
4.3.2	FIRE HANDLING AND SMOKING BAN	<u>1948</u>
4.4	GAS EXPLOSION IN BOILER.....	<u>1918</u>
4.5	SMELT-WATER EXPLOSION IN BOILER.....	<u>1918</u>
4.6	CORROSION PROBLEMS	<u>2019</u>
4.7	SPARKING/STATIC ELECTRICITY	<u>2019</u>
5	DILUTE NON-CONDENSIBLE GAS SYSTEMS IN A RECOVERY BOILER	<u>2120</u>

5.1	COMPOSITION AND AMOUNT	<u>2120</u>
5.2	COOLING/HEATING.....	<u>2322</u>
5.3	STARTING/TRIPPING LOGIC FOR THE INCINERATION OF DNCG.....	<u>2322</u>
5.3.1	STARTING LOGIC FOR INTRODUCING DNCG.....	<u>2423</u>
5.3.2	TRIPPING LOGIC FOR INTRODUCING DNCG	<u>2524</u>
5.4	FAN.....	<u>2624</u>
5.5	DILUTION AIR INLET	<u>2625</u>
5.6	DUCTS	<u>2625</u>
5.6.1	FLOW	<u>2726</u>
5.6.2	POTENTIAL BALANCING	<u>2726</u>
5.7	CONDENSATE DRAINING	<u>2726</u>
5.7.1	SIZE OF CONDENSATE DRAIN LINES	<u>2826</u>
5.7.2	DUCT SLOPES AND LOCATIONS OF CONDENSATE DRAINS.....	<u>2826</u>
5.7.3	WATER SEALS	<u>2827</u>
5.8	SCRUBBERS/CONDENSERS.....	<u>2927</u>
5.9	VALVES/DAMPERS	<u>2927</u>
5.10	BYPASS.....	<u>2928</u>
5.11	DROPLET SEPARATOR	<u>3028</u>
5.12	FLAME ARRESTER	<u>3028</u>
5.13	UNDERPRESSURE PROTECTION DEVICE	<u>3028</u>
5.14	OVERPRESSURE PROTECTION DEVICE	<u>3028</u>
5.14.1	RUPTURE DISC	<u>3028</u>
5.15	CONCENTRATION MEASUREMENT	<u>3028</u>

6	DISSOLVING TANK VENT GAS HANDLING SYSTEMS AT THE RECOVERY BOILER.....	<u>3130</u>
6.1	COMPOSITION AND AMOUNT	<u>3130</u>
6.2	MIXING TANK VENT GASES.....	<u>3231</u>
6.3	COOLING/HEATING.....	<u>3231</u>
6.4	STARTING/TRIPPING LOGIC FOR THE INCINERATION OF VENT GASES.....	<u>3332</u>
6.4.1	STARTING LOGIC FOR INTRODUCING DISSOLVING TANK VENT GASES.....	<u>3332</u>
6.4.2	TRIPPING LOGIC FOR INTRODUCING DISSOLVING TANK VENT GASES	<u>3433</u>
6.5	FAN.....	<u>3433</u>
6.6	DUCTS	<u>3533</u>
6.6.1	FLOW	<u>3533</u>
6.6.2	POTENTIAL BALANCING	<u>3534</u>
6.7	CONDENSATE DRAINING	<u>3534</u>
6.7.1	SIZE OF CONDENSATE DRAINS.....	<u>3534</u>
6.7.2	DUCT SLOPES AND LOCATIONS OF CONDENSATE DRAINS.....	<u>3634</u>
6.7.3	WATER SEALS	<u>3634</u>
6.8	SCRUBBER	<u>3735</u>
6.9	VALVES/DUMPERS	<u>3735</u>
6.10	BYPASS.....	<u>3735</u>
6.11	DROPLET SEPARATOR	<u>3836</u>
6.12	FLAME ARRESTER	<u>3836</u>
6.13	OVERPRESSURE PROTECTION.....	<u>3836</u>
6.13.1	RUPTURE DISC	<u>3836</u>
6.14	CONCENTRATION MEASUREMENT	<u>3836</u>

7	INCINERATION OF CNCG IN THE RECOVERY BOILER	<u>3937</u>
7.1	COMPOSITION AND AMOUNT	<u>4038</u>
7.2	STARTING/TRIPPING LOGIC FOR INCINERATION OF CNCGs	<u>4139</u>
7.2.1	STARTING LOGIC FOR THE CNCG INCINERATION.....	<u>4240</u>
7.2.2	CAPACITY LIMIT	<u>4241</u>
7.2.3	TRIPPING LOGIC FOR THE CNCG INCINERATION.....	<u>4241</u>
7.2.4	BURNER AND USE OF SUPPORT FLAME.....	<u>4342</u>
7.3	PIPELINE SYSTEM	<u>4644</u>
7.3.1	STEAM PURGING OF PIPELINES	<u>4644</u>
7.3.2	PIPELINE PRESSURE MEASUREMENTS	<u>4644</u>
7.3.3	POTENTIAL BALANCING	<u>4745</u>
7.4	CONDENSATE DRAINING	<u>4745</u>
7.4.1	SIZE OF CONDENSATE DRAINING LINES.....	<u>4745</u>
7.4.2	PIPE SLOPES AND LOCATIONS OF CONDENSATE DRAINS.....	<u>4745</u>
7.4.3	WATER SEALS.....	<u>4846</u>
7.5	VALVES	<u>4846</u>
7.6	TRANSFER OF CNCGs.....	<u>4947</u>
7.6.1	PRESSURE-RETAINING STEAM.....	<u>5047</u>
7.6.2	HEATING/COOLING	<u>5047</u>
7.6.3	DROPLET SEPARATOR.....	<u>5048</u>
7.6.4	FLAME ARRESTER	<u>5048</u>
7.6.5	UNDERPRESSURE PROTECTION	<u>5048</u>
7.6.6	OVERPRESSURE PROTECTION.....	<u>5048</u>
7.6.7	RUPTURE DISCS.....	<u>5148</u>
7.7	CONCENTRATION MEASUREMENT	<u>5149</u>
7.8	ALTERNATIVE LOCATION	<u>5149</u>
7.8.1	VALVES.....	<u>5249</u>
7.8.2	STEAM PURGING	<u>5249</u>
7.8.3	LOCATION OF FLARE	<u>5250</u>
8	METHANOL/TURPENTINE INCINERATION IN THE RECOVERY BOILER	
	<u>5351</u>	
8.1	STARTING/TRIPPING LOGIC OF THE METHANOL/TURPENTINE INCINERATION.....	<u>5351</u>
8.1.1	STARTING LOGIC FOR METHANOL AND TURPENTINE INCINERATION.....	<u>5452</u>
8.1.2	TRIPPING LOGIC FOR METHANOL AND TURPENTINE INCINERATION	<u>5552</u>
8.2	ELECTRICAL AND AUTOMATION SYSTEMS OF THE INCINERATOR.....	<u>5553</u>
8.3	VALVES	<u>5553</u>
9	ACTIONS IN SPECIAL SITUATIONS	<u>5755</u>
9.1	ACTIONS DURING DISTURBANCES	<u>5755</u>
9.2	ACTIONS DURING SHUT-DOWNS.....	<u>5755</u>
9.3	ACTIONS DURING START-UPS.....	<u>5755</u>
9.4	ACTIONS DURING OUTAGE.....	<u>5856</u>
9.4.1	INSTRUCTIONS FOR PREPARING NCG SYSTEM FOR SHUT-DOWN OR MAINTENANCE.....	<u>5856</u>
9.4.2	INSTRUCTIONS FOR MAINTENANCE WORK DURING OUTAGES.....	<u>5957</u>
10	SPECIAL INSTRUCTIONS FOR THE DESIGN OF NCG LINES	<u>6058</u>

10.1	CLASSIFICATION.....	<u>6058</u>
10.2	SIGNS.....	<u>6159</u>
10.3	CONDENSATE DRAINING	<u>6159</u>
10.4	DUCT/PIPELINE MATERIAL	<u>6159</u>
10.5	DUCT INSULATION	<u>6159</u>
10.6	PIPE/DUCT SLOPES.....	<u>6260</u>
10.7	VALVES/DAMPERS	<u>6260</u>
10.8	FLANGES, CONNECTIONS	<u>6260</u>
10.9	FANS.....	<u>6260</u>
10.10	EJECTORS/LIQUID RING COMPRESSOR.....	<u>6361</u>
10.11	DUCT SIZING.....	<u>6361</u>
10.12	LOCATION OF FLAME ARRESTER	<u>6361</u>
10.13	LOCATION OF RUPTURE DISCS	<u>6462</u>
10.14	LOCATION OF BYPASS	<u>6462</u>
10.15	PLACE CLASSIFICATIONS	<u>6462</u>
10.16	MANUFACTURING PERMITS	<u>6563</u>
11	OTHER INSTRUCTIONS AND REGULATIONS AFFECTING DESIGN AND OPERATION.....	<u>6664</u>
11.1	FINNISH LAW	<u>6664</u>
11.1.1	HAZARD AND OPERABILITY STUDY (HAZOP)	<u>6664</u>
11.1.2	CLASSIFICATION OF PIPELINES	<u>6664</u>
11.1.3	CLASSIFICATION OF POTENTIALLY EXPLOSIVE ATMOSPHERES (ATEX)	<u>6765</u>
11.2	EUROPEAN STANDARDS	<u>6765</u>
11.3	EXISTING INSTRUCTIONS.....	<u>6866</u>
11.3.1	SODAHUSKOMMITTÉN	<u>6866</u>
11.3.2	BLRBAC	<u>6866</u>
12	DEVIATIONS FROM PRACTICES DESCRIBED IN OTHER INSTRUCTIONS	
	<u>7068</u>	
12.1	DEVIATIONS COMPARED TO BLRBAC'S INSTRUCTIONS.....	<u>7068</u>
13	EXAMPLES OF ACCIDENTS	<u>7169</u>
13.1	EXAMPLE 1:.....	<u>7169</u>
13.2	EXAMPLE 2:.....	<u>7270</u>
13.3	EXAMPLE 3:.....	<u>7472</u>
13.4	EXAMPLE 4:.....	<u>7572</u>
13.5	EXAMPLE 5:.....	<u>7573</u>
13.6	EXAMPLE 6:.....	<u>7775</u>
13.7	EXAMPLE 7:.....	<u>7876</u>
13.8	EXAMPLE 8:.....	<u>7977</u>
13.9	EXAMPLE 9:.....	<u>7977</u>
14	CLOSING WORDS	<u>8179</u>
15	REFERENCES	<u>8280</u>



APPENDICES

| Appendix I

Appendix II

Appendix III

Example of calculation procedure for furnace heat loading

Alfa pinene; Turpentine balance curve

Methanol balance curve



1 ~~GG~~GENERAL

Incineration of non-condensable gases (NCG) in recovery boilers is a common procedure. The Finnish Recovery Boiler Committee does not recommend take a stand on where NCGs should be incinerated. Incineration of NCGs in the recovery boiler means an added risk to boiler operation and the handling of NCGs may cause safety problems. However, since NCGs are nevertheless being incinerated, the members of the committee have called for a clear instruction for the handling of non-condensable pulp mill gases and their incineration in recovery boilers. The Finnish Recovery Boiler Committee wants to improve safety and the performance of the overall process.

This recommendation refers to non-condensable gas systems of recovery boilers and related equipment. The recommendation does not express a view as to where and how NCGs should be collected or on the equipment related to these functions. A holistic focused approach should be used when designing NCG collection systems. Also, the recommendation does not attempt to present a uniform system for handling NCGs, nor promote the use of uniform equipment or process designs. Instead, the recommendation provides basic information for use in engineering, manufacture and operation.

The recovery boiler as an operating environment requires highly reliable and safe systems. It makes good sense to invest in quality equipment and in correct usage. Even very short operating shut-downs caused by disturbances often cost more than the investment required to prevent them. The incineration of NCGs makes the recovery boiler operation more complicated, and the handling of inflammable toxic gases increases the risk of accidents. Complying with the instructions in this recommendation will help minimise hazards and dangerous situations.

This recommendation does not express a view on the contents of operator training required for operating NCG collection and incineration systems. Appropriate training and orientation programmes are vital before new equipment is taken into use and before new operators are allowed to take control over NCG incineration systems.



The recommendation does not ~~give guidelines to take a stand on~~ the mechanical service and maintenance functions of equipment related to NCG systems, but it does express a view on inspection intervals and methods.

In drawing up this recommendation, special care has been taken to confirm careful preparation, while taking into account the opinions of pulp mill operators, equipment suppliers and insurance companies concerning incineration of NCGs. This recommendation is based on the assumption that the equipment of a NCG system primarily meets the requirements imposed by the current national laws, regulations and supervisory authorities, including work safety legislation and laws and regulations concerning inflammable liquids and gases. The equipment of NCG systems is also assumed to be professionally designed, first-class, functional and reliable in use and manufactured with top-quality materials and workmanship.

The responsibility for the performance and process design solutions of NCG systems remains with the equipment ~~suppliers~~ designers. The responsibility for correct and careful operation of NCG systems rests with the operators of the plant. The Finnish Recovery Boiler Committee's recommendation is that the supplier and ultimate user of the plant together perform a hazard and operability study (e.g. HAZOP) of the installation and review the information required by risk assessment. In its report series, the Finnish Recovery Boiler Committee has published report No. 11/2001 entitled "Recovery boiler risk assessment" (in Finnish).

In spite of what is said in this instruction, precedence shall always be given to current laws and regulations and to instructions and guidelines issued by the authorities.



2 ~~D~~EFINITIONS AND ABBREVIATIONS

Non-condensable gases ~~contain gases released during pulp manufacturing, volatile compounds entering with wood, water vapour and air. The gases and vent gases~~ released during cooking, black liquor handling and causticising, that is hydrogen sulphide, methyl mercaptan, dimethyl sulphide and dimethyl disulphide, methanol and turpentine, are inflammable, explosive and strongly odorous compounds giving non-condensable gases their characteristic smell. The volatile compounds ~~from entering with the~~ wood ~~raw material~~, in other words turpentine and methanol, are not odorous in a pure state, but in pulp mills they contain impurities in the form of odorous components.

In this recommendation, odorous gases are generally divided into the following categories:

- diluted non-condensable gases (DNCG)
- dissolving and mixing tank vent gases
- concentrated non-condensable gases (CNCG)
- liquid methanol
- liquid turpentine

2.1 Dilute non-condensable gases

Dilute non-condensable gases are collected from non-pressurised tanks and equipment in the fibre line, evaporation plant, tall oil cooking plant and causticising plant. Gases from pressurised black liquor tanks should be led to the CNCGs system. Leading of gases from non-pressurised black liquor tanks to the collection system for CNCGs or DNCGs should be considered on a case-by-case basis.

Dilute non-condensable gases contain the same components as CNCGs. DNCGs include so much leakage air that their concentrations are below the explosion limit. Processes used to handle DNCGs should ensure that these gases remain dilute in all operating circumstances.

The English abbreviation HVLC refers to High Volume Low Concentration gases. In the Recommended Good Practice published by the BLRBAC, the American term DNCG (Dilute Non-Condensable Gas) is used.



2.2 Dissolving tank and mixing tank vent gases

The ash and black liquor mixing tank can be connected to the DNCGs or dissolving tank vent gases, for example via a ~~dump~~drain-tank. The collection system for mixing tank vent gases must be designed to keep the gases at a dilute concentration under any conditions.

Dissolving tank vent gases contain inorganic dust, ammonia and malodorous sulphur compounds. Vent gases have not traditionally been classified as non-condensable gases. However, vent gases ~~conducted~~introduced into the recovery boiler furnace must be handled in accordance with the instructions for handling DNCGs, ~~e.g. because of their high moisture content.~~

In the Recommended Good Practice published by the BLRBAC, the American term DTVG (Dissolving Tank Vent Gases) is used.

2.3 Concentrated non-condensable gas

Concentrated non-condensable gases originate mainly from the evaporation plant, stripping, methanol plant, ~~firing~~combustion liquor tank and ~~heavy~~concentrated waste liquor tank. They may also come from the cooking plant.

Since CNCGs contain inflammable gaseous compounds their handling involves a risk of explosion. This risk is eliminated by preventing air leaks into the system and by eliminating sources of ignition energy.

Reduced sulphur compounds (TRS, Total Reduced Sulphur) and especially turpentine are explosive in a wide concentration range.

The English abbreviations CNCG are used for Concentrated Non-Condensable Gas and LVHC for Low Volume High Concentration Gas.

SOG (Stripper Off-Gases) refers to concentrated non-condensable ~~concentrated~~ gases originating from the stripper. This gaseous mixture contains methanol (30% - 50% by weight) as well as TRS compounds, different terpenes and other high molecular weight organic compounds. ~~In those gases where for example methanol is in the steam phase. The stripper off-gas is led directly to incineration or to a~~

liquid methanol plant. If stripper off-gases are subjected to methanol liquefaction, the resulting non-condensable gases are led to the CNCG collection system. When ~~there are no~~ SOG is led to methanol liquefaction, a more stable and reliable NCG incineration process will be achieved, also in disturbance situations in the evaporation plant and stripper.

Figure 2-1 is an example of a CNCG burner, showing for example CNCG inlet line, liquid methanol lance, flame ~~scanners~~detectors, flame arrester and support fuel lance.



Figure 2-1. Concentrated non-condensable gas burner

2.4 Definitions

Lower Explosion Limit, LEL	Above this limit the concentration of inflammable gases may be high enough to cause an explosion (air as such does not explode).
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Upper Explosion Limit, UEL Below this limit the concentration of inflammable gases may be low enough to cause an explosion (fuel as such does not explode).

Explosive Region Region between LEL and UEL (when above LEL there is enough fuel to cause explosion, when below UEL there is enough oxygen to cause explosion)

2.5 Abbreviations

Abbreviation	Finnish definition	English definition
BLRBAC	Amerikan soodakattilayhdistys	Black Liquor Recovery Boiler Advisory Committee
CH ₃ SH (MM)	metyylimerkaptani	methyl mercaptan
(CH ₃) ₂ S (DMS)	dimetyylisulfidi	dimethyl sulphide
(CH ₃) ₂ S ₂ (DMDS)	dimetyylidisulfidi	dimethyldi sulphide
CH ₃ OH (MeOH)	metanoli	methanol
C ₁₀ H ₁₆	tärpähti	turpentine
C ₂ H ₅ OH	etanoli	ethanol
CO ₂	hiilidioksidi	carbon dioxide
CO	hiilimonoksidi	carbon monoxide
H ₂ S	rikkivety	hydrogen sulphide
H ₂ O	vesi	water
NH ₃	ammoniakki	ammonia
NO	typpioksidi	nitric oxide
O ₂	happi	oxygen
SO ₂	rikkidioksidi	sulfur dioxide
CNCG	väkevät hajukaasut	concentrated non-condensable gas
DNCG	laimeat hajukaasut	dilute non-condensable gas
HAZOP	poikkeamatarkastelu	hazard and operability study
HEL	ylempi räjähdysraja	higher explosion limit
HVLC	laimeat hajukaasut	high volume low concentration
LEL	alempi räjähdysraja	lower explosion limit
LVHC	väkevät hajukaasut	low volume high concentration
NCG	hajukaasu	non-condensable gas
SHK	Ruotsin soodakattilayhdistys	Sodahuskommittén
SOG	stripperin kaasut	stripper off-gas
TRS	pelkistyneet rikkiyhdisteet	total reduced sulfur compounds
UEL	ylempi räjähdysraja	upper explosion limit
VOC	haihtuvat orgaaniset yhdisteet	volatile organic compounds
(TLV)	haitalliseksi tunnettu pitoisuus	threshold limit value

2.6 Explosion limit formulas

The explosion limit is typically calculated based on the components in the gas mixture.

The Ministry of Social Affairs and Health Decree on Chemical Classification Principles and Labelling (807/2001), Annex 1, Item 7.7.1, provides a formula for determining inflammability. It is based on [Standard ISO 10156](#) and its Tables 1 and 2, which provide calculation constants for different inflammable and inert gases. Table 2-12 resumes the characteristics of the different components of non-condensable gases in air.

Table 2-1. Characteristics of different components of non-condensable gases in air. Burgess andja Young 1992.

		Hydrogen sulphide	Methyl- mercaptan	Dimethyl sulphide	Dimethyl disulphide	Turpentine (α pinene)	Methanol
Formula		H ₂ S	CH ₃ SH	(CH ₃) ₂ S	(CH ₃) ₂ S ₂	C ₁₀ H ₁₆	CH ₃ OH
Molecular mass		34	48	64	94	132	32
Explosion limit	LEL	4.3	3.9	2.2	1.1	0.8	5.5
	UEL	45.0	21.8	19.7	16.1	6.0	36.5
Flame velocity	m/s			0.55		0.62	0.5
Auto-ignition temperature	°C	260	197 (340)	206	300	255	385 (465)
Flash point	°C	-82 Gas	-18 Gas	- 49	24	34	11
Boiling point	°C	- 60	6	38	110	150	65
Heat value HHV	MJ/kg	15	22	31	23	41	22
Volume weight of gas compared to air		1.19	0.87	2.14	3.24	4.7	1.11
Odour threshold	ppm	0.008	0.0004-0.003	0.001-0.01	0.003-0.011	100	100
Solubility in water		Yes	No	No	Poor	No	Yes

When defining ignition limits, that is to say explosion limits, it is advisable to use a method which is guaranteed to give safe limits. Explosion limits can be calculated according to different principles. For gas mixtures the calculation of explosion limits in Section 5.2. of [Standard ISO 10156/2010](#) can be used.

Another method is to examine the problem thermodynamically (Hokynar 1999). There is not much difference in the definition of the lower explosion limit, but the definitions of the higher explosion limit differ because of the dissimilar impact of inert gas in the two methods.

2.7 Hydrogen sulphide

Hydrogen sulphide is a colourless, toxic, very easily flammable gas heavier than air, with the characteristic foul odour of rotten eggs. However, when reaching values of 100 ppm, an exposed person generally can no longer observe the odour of



hydrogen sulphide ~~generally is no longer malodorous~~. A leak of hydrogen sulphide in outdoor locations may cause ignition risks and in indoor locations explosion risks. Hydrogen sulphide corrodes several metals and plastics.

Hydrogen sulphide acts as a reducing agent and it may react violently with oxidising substances. The substance corrodes metals and generates metal sulphides. In water solutions hydrogen sulphide reacts as an acid. Hydrogen sulphide is generated when solutions containing sulphide, for example sodium sulphide, interact with acids, and it can be present in the whole mill area. It may spread over long distances along ducts and sewers. For this reason, gas alarms are installed in pulp mill areas where hydrogen sulphide may be present. When a detector notices a strong enough concentration of dangerous gas, the alarm lights installed in the system and close to measurement points are activated.

2.8 Methyl mercaptan

Methyl mercaptan is a colourless, very easily flammable, volatile and toxic gas, heavier than air. The detrimental effects of methyl mercaptan on health are similar to those of hydrogen sulphide.

2.9 Dimethyle sulphide/dimethyle disulphide

Under normal conditions, dimethyle sulphide and dimethyle disulphide are colourless liquids with a characteristic foul smell. Both compounds are easily flammable and explosive in contact with air.

2.10 Methanol

Pure methanol (CH_3OH) is a colourless, almost odourless and toxic gas/liquid, but in a sulphate pulp mill methanol is contaminated by numerous malodorous compounds. Methanol is soluble in water in all volumetric ratios and can therefore easily be separated from a gas mixture in a scrubber.

Methanol vaporizes from black liquor during evaporation and ends up to foul condensate. Foul condensate is purified in a stripping column and methanol is condensed from stripper off gas (SOG) in a methanol column. Liquefied methanol is pumped via a storage tank to incineration.



2.11 Turpentine

Turpentine ($C_{10}H_{16}$) is produced from volatile organic compounds (terpenes) in wood during chemical pulp production. Pure turpentine is a colourless and mildly odorous gas/liquid, but in sulphate pulp mills turpentine is contaminated by numerous malodorous compounds. A typical turpentine composition is shown in Table 2-1.

Table 2-21. Typical turpentine composition

Compound	Content, %
Alpha-pinene	80 - 90
Beta-pinene	5 - 10
Delta-3-carene	5 - 10
Other $C_{10}H_{16}$ compounds	< 1
Methyl mercaptan	0 - 5
Dimethyl sulphide	0 - 12
Dimethyl disulphide	0 - 1

It is important to remove turpentine from NCG as effectively as possible, because it is explosive in a very wide concentration range. It shall be noted that turpentine has the lowest LEL, about 0.8 %, and even the flame propagation velocity of turpentine is low, the detonation velocity after explosion is high (100-130 m/s), while it is only about 1 m/s in turpentine free gas.

When collecting gases from terpene (turpentine vapour) containing sources, high concentrations can be avoided by condensation of turpentine compounds. These are separated in a scrubber by condensing with cold water. The scrubber is installed after the turpentine condenser in the NCG line. Separated turpentine is stored in turpentine tank.

In spite of condensation, treated gas contains terpenes proportional to their partial pressure. This means that sources of DNCGs possibly with a high content of terpenes must be diluted after collection or condensation. If the collection point maximum temperature is exceeded, the collected stream must be vented. Partial pressures for some turpentine compounds are shown in Figure 2-1.

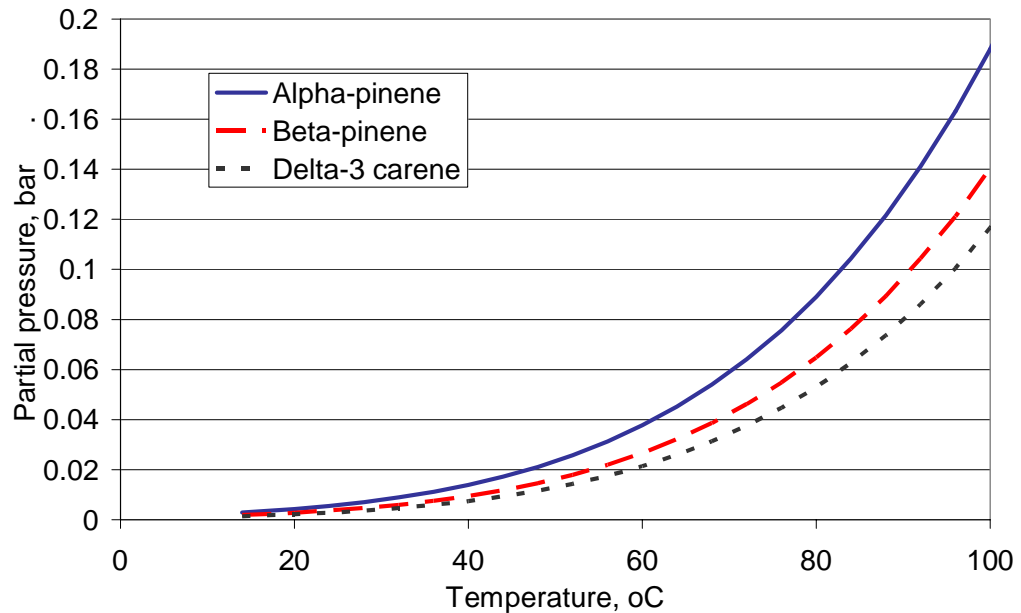


Figure 2-24. Partial pressures for some turpentine compounds, Drew et al., 1971.

Sources of terpenes in NCG collection can be: softwood chip bins, ~~fouled~~ contaminated condensate tanks, collection and weak liquor tanks, especially if they receive condensate containing ~~turpentine~~ terpene, even if only occasionally.

Also in shut-down and start-up situations, evaporated turpentine may vaporise with odorous gases on the surfaces of evaporation tanks or ducts.

In addition to NCG, turpentine can be separated from the bottom condensate in methanol distillation by means of a turpentine decanter. Turpentine separates to water surface, because it's lighter than water and poorly soluble. The raw turpentine is removed from decanter with overflow.



3 **IMPACTS ON RECOVERY BOILER EMISSIONS**

Incineration of NCGs in the recovery boiler may affect recovery boiler emissions. Incineration of NCGs may reduce the carbonate content and pH value of the recovery boiler precipitator ash. The additional sulphur in NCGs is first seen as replacement of carbonate with sulphate in the electrostatic precipitator ash. When the carbonate content becomes low, the remaining sulphur will form SO₂.

3.1 **Impacts on recovery boiler sulphur emissions**

The extra sulphur contained in NCGs always raises the sulphate content of recovery boiler ash and may therefore reduce its pH. Therefore, if there is not a sufficient carbonate buffer, ~~it may increase~~ the boiler's SO₂ emissions may increase. ~~A mill's~~ with high sulphidity, low black liquor dry solids content, low heat value of liquor ~~the solid matter and heat value of combustion liquor~~ and low boiler load affect the boiler's SO₂ emissions.

Non-condensable gases should not be incinerated in a recovery boiler that already has sulphur emissions. When this is the case, the NCGs should be incinerated for example in a separate incinerator with sulphur recovery. Such incinerators are for example ~~NCGspecial~~ boilers and lime kilns with scrubbers.

3.2 **Impacts on recovery boiler NO_x emissions**

Especially CNCGs contain nitrogen compounds. These nitrogen compounds consist mostly of ammonia, which is separated from black liquor in the evaporation plant.

In normal cooking, organic nitrogen compounds in the wood dissolve more or less completely in the alkaline cooking liquor and are thus transferred as part of the black liquor to the evaporation plant and further to the recovery boiler. Significant fraction of the organic nitrogen in the black liquor is transferred into the recovery boiler smelt as sodium cyanate, NaOCN. The cyanate nitrogen in the smelt passes into the green liquor through the dissolving tank along with the smelt's other salts. In the green liquor, alkaline hydrolysis gradually transforms cyanate nitrogen into ammonia.



The ammonia formed in the green liquor and in the white liquor partially evaporates into dilute malodorous gases. However, most of the ammonia continues as part of the white liquor back to the cooking. From the cooking, the ammonia passes into the black liquor together with the organic nitrogen compounds of the wood. The ammonia is finally quantitatively separated from the liquor in the evaporation plant, passing into concentrated malodorous gases and methanol.

~~Part of these nitrogen compounds have been formed in the causticising area from nitrogen entering the dissolving tank with recovery boiler smelt which is then recirculated through the cooking process to the recovery boiler.~~

3.2.1 Impact of DNCG

Dilute non-condensable gases typically account for 5...15% of the total amount of air fed into the recovery boiler. The nitrogen contained in DNCGs is primarily ammonia. The ammonia content in dry gases can be assumed to be about 10 ppm. About 30% of the ammonia could be assumed to be converted into NO. If the boiler's NO level is about 100 ppm, the increase in the boiler's nitrogen oxide emissions would be:

$$NO_x = \frac{(100 + 0.3 * 0.1 * 10)}{100} - 1 \leq 1\%$$

No measurable impact has been detected in practice, if DNCGs are directed to the correct temperature range in the furnace.

3.2.2 Impact of dissolving tank vent gases

The air contained in dissolving tank vent gases typically accounts for 5% of the total amount of air fed into the recovery boiler. The nitrogen contained in dissolving tank vent gases originates from the evaporation plant's secondary condensates or from the conversion into ammonia of nitrogen entering the dissolving tank with the smelt. The ammonia content in dry gases can be assumed to be about 100 ppm. About 30% of the ammonia could be assumed to be converted into NO. If the boiler's NO level is about 100 ppm, the increase in the boiler's nitrogen oxide emissions would be:



$$NO_x = \frac{(100 + 0.3 * 0.05 * 100)}{100} - 1 \approx 1\%$$

No measurable impact has been detected in practice, if DNCGs are directed to the correct temperature range in the furnace.

However, in some boilers slight NO_x emission increase has been reported when DNCG gases are led into the tertiary air level.

3.2.3 Impact of CNCG

CNCGs typically account for 0.2...3% of the total amount of air fed into the recovery boiler. The nitrogen contained in CNCGs is primarily ammonia. The ammonia content in dry gases can be assumed to be about 2000 ppm. About 30% of the ammonia could be assumed to be converted into NO. If the boiler's NO level is about 100 ppm, the increase in the boiler's nitrogen oxide emissions could be:

$$NO_x = \frac{(100 + 0.3 * 0.01 * 2000)}{100} - 1 \approx 6\%$$

3.2.4 Impact of methanol

Approximately 10 kg/ADt of methanol is generated, depending on the wood raw material used. The ammonia content in liquid methanol may be around 2%. The ammonia content in methanol may be assumed to be approximately 0.2 kg/Adt. It can therefore be assumed that around 30% of the ammonia is converted into NO. If the boiler's NO content is 200 mg/Nm³, 3.5 Nm³/kg of combustion gas is generated and the amount of liquor dry solids is 1600 kgds/Adt. Then the increase in the boiler's nitrogen oxide emissions may be as follows:

$$NO_x = \frac{(200 * 3,5 * 1600 / 1000000 + 0,3 * 0,2)}{200 * 3,5 * 1600 / 1000000} - 1 \approx 5\%$$

3.3 Impacts on recovery boiler TRS emissions

In modern boilers with efficient mixing of fuel and air and a NCG system that functions according to design specifications, the boiler's TRS emissions are not significantly increased, provided that the furnace temperature is at least 900 °C.



When CNCG ~~are~~ firing is started with low boiler load it is possible to get high SO₂ and TRS emissions.

If DNCGs ~~is~~~~are~~ fed to the boiler's above liquor gunstopmost air register, a small amount of NCGs may pass without reacting along cold regions of the boiler wall.

The measurable impact in total flow has in practice been much below one ppm.

3.4 Impacts on recovery boiler fouling

See item 3.1, Impacts on recovery boiler sulphur emissions.

If the combustion conditions for NCGs in the recovery boiler are not optimal and SO₂ emissions are generated, there is a risk that boiler bank, economizers, and electrostatic precipitator may get fouled and also corrosion may occur especially if low ash pH prevails for long period.

~~the boiler's and the electric filter's susceptibility to fouling and corrosion will increase, especially in the economiser area, due to acid ash salts (low pH).~~

3.5 Impacts on other recovery boiler functions

In modern boilers with efficient mixing of fuel, ~~and~~ air and sufficient base load operating according to design specifications, no impacts on other recovery boiler operation parameters functions have been observed. For example, changes in reduction rate have not been reported.

Incineration of NCGs gases generally reduces the mill's total sulphur emissions. This, however, affects the mill's chemical balances and tends to increase sulphidity. Therefore the sulphur balance at a mill has to be examined on a case-by-cbase basis.

Water vapours ~~is~~~~are~~ introduced into the boiler with dissolving tank vent gases and DNCGs, which may lower the boiler's capacity somewhat, depending on water vapour content. Figure 3-1 presents the maximum amount of water vapour contained in ~~dry~~ air at a specific temperature. Warm air can hold more vapour than cold air. The incineration of CNCGs, methanol and turpentine increases boiler



steam generation slightly vaporisation. In that case the design of the water and steam vapour systems has to be checked before starting incineration, especially in existing recovery boilers.

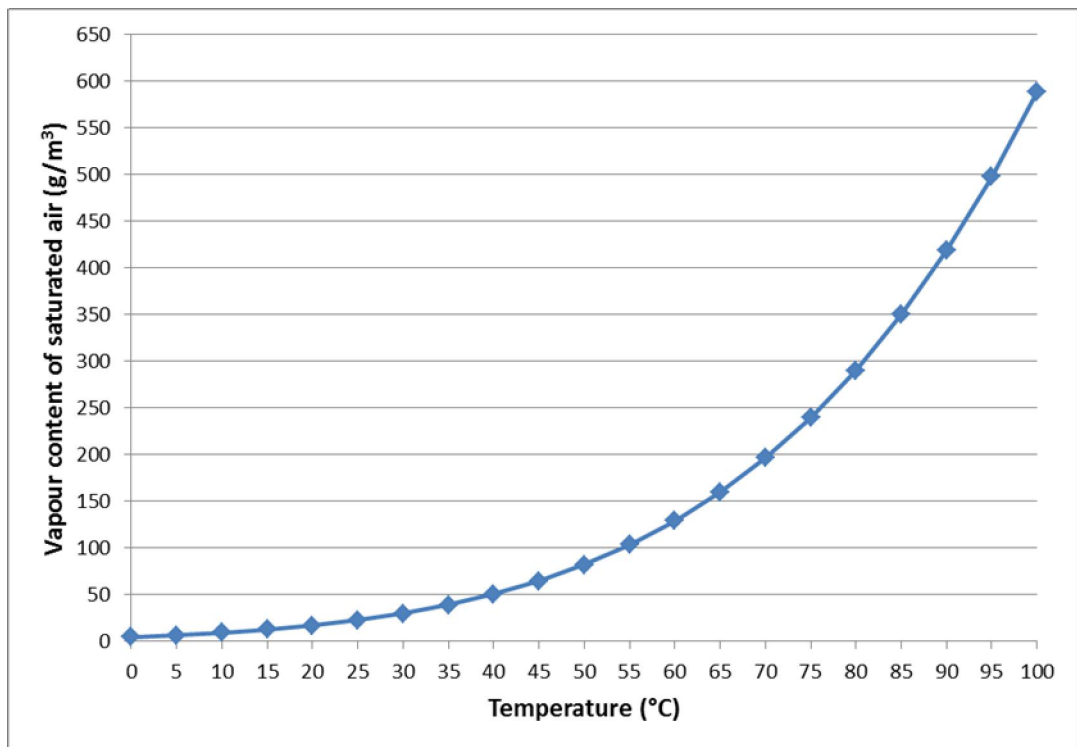


Figure 3-1. Saturated water vapour content in air as a function of temperature.



4 **RRISKS AND CORROSION RELATED TO NON-CONDENSIBLE GAS SYSTEMS**

Typical risks related to non-condensable gas incineration systems include explosion risk (gas concentration and ignition by sparks, for example as a result of overpressure in the furnace), danger of toxicat~~ion~~of being poisoned, condensate draining problems (water entering the furnace) and corrosion/erosion.

The Finnish Recovery Boiler Committee's recommendation is that a hazard and operability study (e.g. HAZOP) shall always be made when NCG system is modified~~in connection with a modification of the system~~ together with the supplier of the NCG system.

In addition, NCG incineration may cause process-related problems, such as sulphur emissions from the boiler, plugging of economiser and other boiler heating surfaces and combustion disturbances in the furnace.

4.1 Leakage of non-condensable gases into manned areas

Non-condensable gases may leak from joints, valves, drains or failure areas~~points of damage~~. The concentrations of reduced ~~organic~~ sulphur compounds and methanol in NCGs are so high that there is a danger to get toxicated~~of being poisoned~~. If a leak occurs into the boiler ~~house~~room or some other manned areas~~space~~, the entire ~~house~~room or space has to be evacuated, duly ventilated and concentrations measured to confirm that it is again safe to work in.

Non-condensable gas can be~~is~~ released into manned enclosures typically~~generally~~ from condensates during~~in connection with~~ maintenance work. Possible leakage sources are water seals in the NCG system, as well as the safety devices~~appliances~~ in the system. Special care is required when~~in~~ operating methanol and turpentine burners. Even a very small leak may cause a major odour problem.

Because concentrated NCG lines are kept at overpressure after the ejector in the recovery boiler building, there may develop a danger of gas poisoning in the recovery boiler building when there is a leak in the NCG line, and in some ineffectively ventilated areas there may also occur an explosion risk.



In accordance with the instruction entitled "Maximum concentrations of impurities in the working place" issued by the Finnish Ministry of Social Affairs and Health, the maximum allowed indoor hydrogen sulphide concentration is 5 ppm (7 mg/m³)/8 h and 10 ppm (14 mg/m³)/15 min. These so-called TLV values (threshold limit value) are the lowest concentrations of airborne chemicals in the workplace that could cause harm to workers.

Mills have typically both fixed and portable hydrogen sulphide and VOC meters. The fixed meters activate blinking warning lights and buzzers if the hydrogen sulphide or VOC content in the air is too high. When placing the meters, for instance air flowcurrents and working areaheights should be taken into consideration. Regular functional tests should be made of both fixed and portable meters and there should be instructions for the meters in the quality systems. The tests carried out should be documented.

Maintenance or repair work on NCG equipment requires special permits.

4.1.1 Odour

Sulphur compounds numb the sense of smell~~organ of smell~~ so it is not advisable to trust the nose alone. This may result in a lethal situation, unless appropriate protective action is taken. Employees working with equipment containing NCGs must familiarise themselves with the operating safety instructions concerning DNCGs and CNCGs.

4.1.2 Safety equipment

A pressurized air mask is the only safety equipment that protects from the effects of NCGs. Gas masks and personal breathing equipment are meant only for leaving a danger zone.

4.2 Condensate draining problems

If water vapour is allowed to condense in a NCG pipeline, there is a risk of a smelt water explosion, if this water enters the recovery boiler.



Special attention must be paid to water drains to confirm that condensate can be removed from all lines in a comprehensive manner.

4.2.1 Overflowing of condensate trap

A common problem with NCG systems is the occasionally large volume of condensate. Condensate traps close to the recovery boiler shall be provided with high-level alarm and, if necessary, with interlocking to lead NCG away from incineration.

4.2.2 Drying out of water seals

Drying of water seals is accelerated when NCGs are heated above condensation temperature. To prevent water seals from drying out, they should always be provided with a sufficient fresh water supply and equipped with flow alarms.

Measures should also be taken to prevent water seals from drying during shut-downs.

4.2.3 Plugging of water seals

Non-condensable gases carry with them small amounts of liquor foam/soap, which may accumulate at the lowest points of the line, for example at condensate drain points. If the drain pipe diameter is not large enough, complete plugging of the pipe may occur.

4.3 Explosion risks in non-condensable gas lines

Transfer and incineration of CNCGs shall be designed so that there is no explosion risk under any circumstances. An explosion risk arises if air leaks into CNCGs. The concentration of CNCGs is maintained above the upper explosion limit by ensuring system sealing.

Transfer and incineration systems for DNCGs are designed to keep the concentration of gases below the lower explosion limit. The concentration of DNCGs is kept below the explosion limit by ensuring sufficient dilution by air. In some cases an explosion risk may arise when water vapour in DNCGs is eliminated by condensation and gas concentration approaches the lower explosion limit.



Ignition requires ignition energy such as static electricity, heat or an electrical spark. Spark generation is best avoided by making sure that groundingsearthings, structures and equipment are selected and installed properly.

4.3.1 Shut-down and start-up situations

Special attention needs to be paid to the concentrations of DNCGs during shut-down and start-up situations. Explosions in recovery boilers caused by NCGs have several times been related to DNCG systems, where the gas concentration has risen too much duringin shut-downs and disturbance situations. A typical problem is the accumulation of explosive gases in the pipeline during a shut-down and their volatilization evaporation when NCG collection is started. in connection with a start-up.

~~4.3.2 Fire handling and smoking ban (tarvitaanko tätä)??~~

~~Smoking and the use of open fire are banned in areas where NCGs are handled. For this reason, NCG lines and permanent fire handling according to the hot work permit, as well as smoking areas, must be separated from each other. Critical areas must be provided with appropriate warning signs.~~

4.4 Gas explosion in boiler

Non-condensable gases are a mixture of inflammable components, oxygen and inert gas (nitrogen, water vapour). As is always the case when handling inflammable gases, there is a risk of explosion in the furnace. A gas explosion occurs if fuel is fed into the furnace without ensuring immediate combustion, after which the mixture absorbs so much energy that it is ignited.

4.5 Smelt-water explosion in boiler

Water condensate getting into the furnace from the NCG line may cause a smelt-water explosion. To prevent this from happening, DNCGs and dissolving tank vent gases must be sufficiently heated so that the major part of the water droplets turns into vapour, and the gas lines are equipped with a sufficient amount of condensate drainsremoval-outlets and slopes. When designing drain systems, attention should be paid for example to the size, amount and location of the drains.



A smelt-water explosion means extremely rapid vaporisation of water that gets into contact with the cooking chemical. The pressure wave caused by this vaporisation may damage the furnace and thus cause a leak of non-condensable gases/water/steam into the boiler ~~house~~room.

4.6 Corrosion problems

The burning of NCGs in recovery boilers has not been found ~~to increase to cause~~ corrosion.

4.7 Sparking/static electricity

For an explosion to occur there must be some kind of initial energy, such as a spark. To confirm good safety, NCG lines are potentially balanced from the collection point forward. The objective of potential balancing is to prevent the appearance of dangerous voltage differences between conducting parts that may be touched simultaneously. For instance in flanges, hatches and connections where a galvanic isolation may be formed for example due to isolation materials, potential balancing must be performed. Potential balancing shall be made according to the given instructions and verified with a field log.

5 ~~DILUTE~~ **DILUTE NON-CONDENSIBLE GAS SYSTEMS IN A RECOVERY BOILER**

Dilute non-condensable gases are not fuel. They primarily consist of air and water vapour. According to the Finnish Chemicals Act, DNCG are generally classified as harmful, so equipment purchases, engineering, installation and operation must be handled accordingly.

To confirm the operating safety of the recovery boiler, necessary precautions must be taken to prevent water from entering the boiler when DNCGs are incinerated. ~~in connection with the incineration of DNCGs.~~ DNCGs are cooled down (part of the water contained in the gases is condensed), transported by a fan and heated up to confirm evaporation of any water droplets before incineration. NCG lines are equipped with condensate drains to prevent water from accumulating in the gas line and from there entering the boiler.

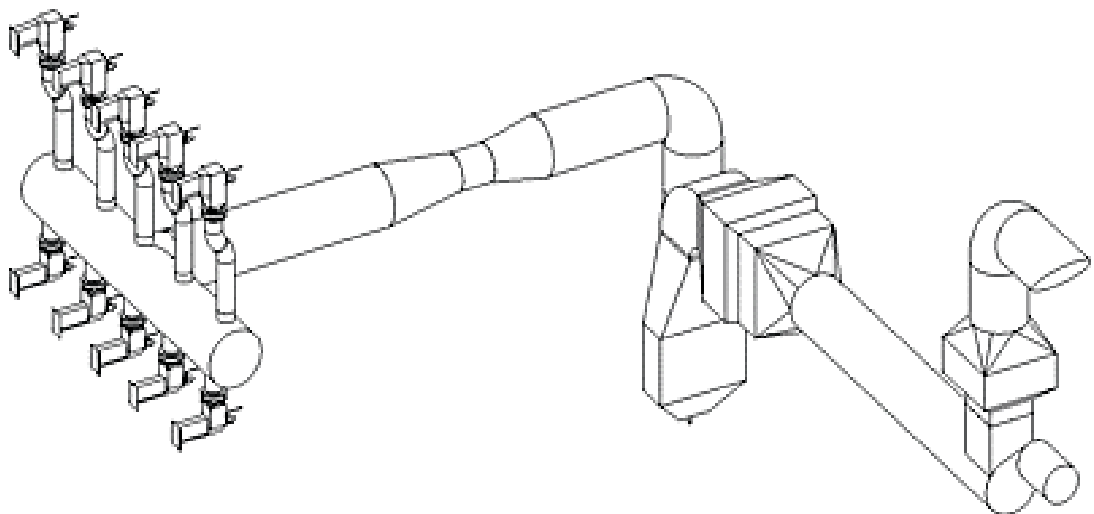


Figure 5-1. Example of a duct system for dilute non-condensable gases

5.1 Composition and amount

Dilute non-condensable gases consist of, among other things, tank and equipment vent gases from the pulp mill's fibre line, vent gases from the evaporation plant's tank farm, the causticising plant's vent gases, etc. DNCGs amount to 300 – 400 m³n/ADt, and their sulphur content is 0.1 – 0.5 kgS/ADt. DNCGs are typically treated in a scrubber-type vent gas cooler, which removes moisture and so reduces the amount of gases to be handled. Typical amounts of DNCGs collected from various pulp mill departments are shown in Table 5-1.

Table 5-1. Amount of DNCGs collected from different departments at a temperature of 40 °C

Department	kg S/ADt	m ³ n/ADt
Vent gases from continuous cooking	0.1 – 0.5	100 – 400
Vent gases from Superbatch cooking (evacuation air, vents from non-pressurised tanks)	0.1 – 0.5	150 – 300
Washing plant vent gases	0.05 – 0.1	100 – 200
Tall oil cooking plant vent gases	0.05 – 0.2	2 – 3
Tank vent gases, evaporation plant (atmospheric pressure tanks)	0.1 – 0.4	20 – 30
Causticising plant-lime kiln area	0.01 – 0.1	5 – 10
TOTAL	0.1 – 0.5	300 – 400

The concentration of DNCGs should continuously be kept below the explosion limit and any increase in concentration should be prevented. When the concentration and moisture content are kept low, DNCGs can be handled at the boiler plant in the same way as other combustion air. Examples of DNCG analyses are given in Table 5-2. The contents of the DNCG components vary extensively depending on the mill and measuring points.

Table 5-2. Examples of dilute non-condensable gas analyses at the point of origin.

Component	Mill A ppm	Mill B ppm	Mill C ppm	Mill D ppm	Mill E ppm	Mill F ppm
H ₂ S	*	*	1	0	725	1
MM, CH ₃ SH	200	40	70	2	14	2
DMS, (CH ₃) ₂ S	1 000	60	160	643	1 530	784
DMDS, (CH ₃) ₂ S ₂	90	*	270	1	10	365
Turpentine, C ₁₀ H ₁₆	1 500	15	*	69	470	12 400
MeOH, CH ₃ OH	900	150	*	834	6 943	2 046
Ethanol, C ₂ H ₅ OH	80	*	*	*	*	*
Oxygen, O ₂	*	*	*	*	*	*
Ammonia, NH ₃	200	*	*	*	*	*
Water, H ₂ O	70 000	50 000	*	*	*	*
Carbon dioxide, CO ₂	*	300	*	*	*	*

*) Not analysed or below detection limit

When incineration of DNCGs is taken into use for the first time at the mill or the process is modified, concentrations in the gas lines at different collection points must be measured to verify the contents of DNCGs in different collecting points. Special attention needs to be paid ~~forte~~ preventing the gas concentration in chip silo vent gases from rising too high in disturbance situations. In the event of a disturbance in the chip silo, vent gases from the silo must not be conducted to the



collection system for DNCGs. Table 5-3 shows the recommended maximum concentrations for DNCGs before incineration. DNCGs are typically much below 10% of LEL.

Table 5-3. Maximum recommended concentrations of DNCGs before incineration

Compound	Limit	Note
Air content, vol-%	> 90	
Water content, vol-%	< 7	40 °C (after condensing)
Water content, vol-%	< 12	50 °C (after condensing)
Sulphur content TRS, ppm	<200	
Turpentine content, ppm	< 1000	
Methanol content, ppm	< 1000	

In addition, it needs to be determined to where turpentine-containing condensates can be diverted safely in the event of a disturbance in the cooking plant. There are examples of cases where a careless connection has caused a significant increase in the turpentine content of the condensate during a cooking plant disturbance.

Dilute non-condensable gases are typically fed into the recovery boiler as a part of the boiler's combustion air. Because of the smelt-water explosion risk caused by the condensate [entering the furnace surge](#), it is not recommended to feed DNCGs into primary air.

5.2 Cooling/heating

Condensate from DNCGs must not be allowed to enter the recovery boiler furnace. The amount of water vapour entering the furnace with DNCGs must be minimised. The temperature of DNCGs should preferably be 40-50 °C after the condenser.

Dilute non-condensable gases are recommended to be heated to 30 °C above the condensate temperature to minimise the entrance of water drops into the furnace.

5.3 ~~Starting/tripping logic~~ **Logic for governing the incineration of DNCG**

The purpose of this section is to describe the process conditions to be fulfilled by the ~~control logic~~ [interlocks](#) ~~controlling~~ [governing](#) the incineration of DNCGs. The focus is on the most essential conditions that a safe automation system must fulfil. Because of differences in equipment design, process connections and technical



implementation, the interlockseontrol-logie of any recovery boiler will also include other conditionsvariables than those presented here.

This recommendation does not cover interlockings related to the start and operation of the collection of DNCGs, though these are often partly implemented in the same system as interlockings related to incineration. These interlockings have to be defined case by case.

The Finnish Recovery Boiler Committee recommends that the DNCG system has been in use for a sufficiently long time (for example over one hour). Several compounds can be condensed in the ducts. It is necessary to vaporise and vent out of the inflammable gas components from the system. If possible, the collection of DNCGs should be operateding during shut-downs. The objective is to prevent the concentration of DNCGs to explosive area in the system.

It must be possible to supervise the incineration of DNCGs in the recovery boiler as well as to control starting and stopping DNCG incineration from the same main control room where the other activities related to the operation of the recovery boiler are controlled.

5.3.1 Starting logicConditions for introducing DNCG

Dilute non-condensable gases can be introduced into the boiler furnace when e.g. the conditions below are fulfilled, Table 5-4. **Because of differences in commissioning year, equipment design, process conditions and technical implementation, the interlockseontrol-logie of any DNCG incinerationrecovery boiler will also include other conditionsvariables than those presented here.**

Table 5-4. To admit DNCGs to the recovery boiler, the following conditions should be fulfilled

Condition	Purpose of condition
1 Boiler steam load/feed water flow over 15% of boiler nominal load. This limit shall be defined on a case-by-case basis	Confirm sufficiently stable furnace conditions to oxidise <u>TRS compoundsgases</u> <u>NCGs</u> (temperature, retention time)
2 DNCG gas fan is on	Verify that there is sufficient flow in the duct to prevent flame propagation backward in the pipeline (burn-back)
3 Sufficient pressure in the duct system ahead of shut-off valve	Verify that there is sufficient flow in the duct to prevent flame propagation backward in



		the pipeline (burn-back)
4	Sufficient flow in the duct system ahead of shut-off valve	Verify that there is sufficient flow in the duct to prevent flame backward progression in the pipeline (backfire)
5	Temperature after cooling (ahead of heater) below temperature limit	Reduce gas volume and the amount of water vapour and gas volume
6	Temperature before introduction into boiler/after heater above temperature limit	Prevent water droplets from entering the boiler
7	Ventilation outlet to the roof (bypass) has been in operation long enough	Prevent concentration of DNCGs in the duct system
8	Trap Condensate pocket level in NCG system (if exists) below upper limit	Prevent condensate from entering the furnace. Indicate fault in the condensate system preventing condensate drainingremoval

5.3.2 Tripping logic for Interrupting introducing feeding of DNCG

The ~~introductionfeed~~ of DNCGs to the boiler furnace has to be ~~stopped interrupted~~ if any of the conditions below is fulfilled, Table 5-5. **Because of differences in commissioning year, equipment design, process conditions and technical implementation, the interlockingcontrol logic of any DNCG incinerationrecovery boiler will also include other conditionsvariables than those presented here.**

Table 5-5. Any of the following conditions will disallow introduction of DNCGs to the recovery boiler:

Condition	Purpose of condition
1 Boiler steam load/feed water flow is below 15% of boiler nominal load. This limit shall be defined on a case-by-bas case <u>case</u> basis	Confirm sufficiently stable furnace conditions to oxidise <u>TRS compounds</u> gases <u>DNCG</u> (temperature, retention time)
2 Emergency shut-down activated <u>tripped</u>	Prevent introducing of DNCGs when the boiler is not in <u>accepted condition use</u>
3 DNCG fan stopped	<u>Prevent flame propagation backward in the pipeline (burn-back)</u>
4 Temperature after cooling (ahead of heater) above temperature limit	Gas moisture content too high
5 Temperature before introduction into boiler/after heating below temperature limit	Amount of water droplets too high
6 Pressure in non-condensate gas duct below the lower limit	<u>Prevent flame propagation backward in the pipeline (burn-back)</u>
7 Trap Condensate pocket level in DNCG system above upper limit	Prevent condensate from entering the furnace. Indicate fault in the condensate system preventing condensate drainingremoval



5.4 Fan

The sealing characteristics of the fan must be sufficient and the fan must be equipped with a condensate ~~draining~~removal system.

5.5 ~~Dilution~~Make-up air inlet

~~Dilute non-condensable gas-DNCG~~ systems may incorporate a ~~make-up~~dilution air inlet (~~excess make-up~~mix air inlet) to allow air to be mixed into the dilute gases. If the safety of NCGs is dependent on the operation of the ~~make-up~~dilution air inlet, special care must be taken to confirm that the ~~make-up~~dilution air supply cannot be switched off for any reason.

For the above mentioned reasons, ~~air addition~~dilution must be sufficiently supervised and measured, for example by flow measurements, and necessary interlockings must be installed for insufficient air flow~~dilution~~.

~~If the Dilution air inlet must be designed so that NCG backflow to working areas is prevented. is installed in the fan ventilator suction inlet, possible pressure discharges from the inlet should be taken into account in the alignment of the inlet. The dilution air inlet, which protects the start-up of the fan, must open as a function of interlocking immediately when the fan stops and slowly close when the fan is started up, to confirm there is enough dilution to protect the fan.~~

5.6 Ducts

Dilute non-condensable gases must be prevented from leaking into the boiler ~~house~~room and the length of the duct inside the boiler ~~house~~room must be minimised.



Figure 5-2. Installation of a mill's dilute non-condensable gas incineration system.

5.6.1 Flow

It is recommended to measure the flow of DNCGs going to incineration. The flow measurement device is recommended to locate~~should be placed~~ as close as possible to the boiler's gas inlet.

5.6.2 Potential balancing

No potential balancing is required for DNCGs.

5.7 Condensate draining removal

Dilute non-condensable gas lines have to be equipped with condensate drains, removal outlets. Condensates shall primarily be conducted as a separate system to the fouled~~contaminated~~ condensate treatment system.

In designing condensate draining, special attention shall be paid for example to drain size, number and location. The last condensate drain is recommended to locate as close to the recovery boiler as possible so that the length of the duct from this drain to the furnace is minimised. This drain should have condensate pocket with limit switch.



5.7.1 Size of condensate ~~drain~~removal lines

Because of the risk of plugging, the recommended minimum diameter for condensate ~~drains~~ ~~removal-outlets~~ ~~and condensate lines starting from these~~ is 50 mm. The condensate drain lines closest to boiler should be at least 100 mm. If there is a possibility of additional water (e.g. washing water) feed into the system, then condensate drain lines shall be sized accordingly.

5.7.2 Duct slopes and locations of condensate ~~drains~~removal-outlets

Ducts and pipes must slope towards the condensate ~~drains~~outlets. Condensate ~~draining~~removal against flow is very difficult. The recommended slope for ducts, elsewhere than in the recovery boiler plant, in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:25.

Because of the risk of water getting into the furnace, the preferred slope for ducts in recovery boiler plants in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:1 or more. A condensate ~~drain~~removal-point must be installed before every point where the NCG line angles upward. If the recommended slopes cannot be achieved in some duct sections, the condensate ~~drains~~outlets in this part of the system must be designed with special care.

~~In designing condensate ~~draining~~removal, special attention shall be paid for example to ~~drain~~inlet size, number and location. The last condensate ~~drain~~outlet should be placed as close to the recovery boiler as possible to minimise the length of the duct from the outlet to the furnace.~~

5.7.3 Water seals

Condensate outlet lines shall be equipped with water seals and a pumping tank or with a common seal tank (condensate collection tank). In this way, DNCGs are prevented from leaking via the condensate ~~drain~~outlet.

To prevent water seals from drying out, they shall be equipped with level indicators or provided with a continuous water flow controlled by flow indicators.



To confirm that the pressure in the DNCG duct will not rise so high as to drain the water seal, the difference between the water levels in the water seal must be allowed to vary by at least 10 kPa, or by an amount corresponding to the duct's design pressure. 10 kPa corresponds to a water level variation of 1 m in the water seal.

If condensates are collected into a pumping [tankbasin](#), the level difference in the [tankbasin](#) must be allowed to vary by at least 10 kPa, or by an amount corresponding to the duct's design pressure.

5.8 Scrubbers/condensers

The dilute non-condensable gas system shall incorporate a sufficient number of scrubbers or condensers to remove the water vapour and VOCs (for example turpentine) contained in the gases and reduce the collected gas volume.

5.9 Valves/dampers

The valves/dampers for the DNCG incineration system shall be selected with care. Special attention shall be paid to their sealing characteristics. Normally, effectively sealing throttle valves/dampers are used for large ~~duct~~[pipe](#) sizes.

The DNCG line should have automatic shut-off valves with “fail-close” actuator which will close DNCG introduction to the recovery boiler automatically if operating energy is turned off. At the same time valve of the alternative location or bypass duct must open (fail-open). Valves must be provided with limit switches indicating position to the automation system.

~~Shut-off valves must be equipped with limit information indicating shut-off to the automation system.~~

Recommended material for the fan is EN 1.4301 (SS2333) or equivalent.

5.10 Bypass

The bypass duct for DNCGs must be led to as high a point as possible. Normally, the bypass duct is led to the roof of the recovery boiler plant or to the ~~chimney~~ stack.



5.11 Droplet separator

A droplet separator should preferably be used at the incineration end of the DNCG duct before the gases are heated up, and always after DNCGs have been cooled or treated in a scrubber.

5.12 Flame arrester

There is no need to use a flame arrester in DNCG ~~system~~ ducts.

5.13 Underpressure protection device

Dilute non-condensable gas lines can be equipped with underpressure protection, if the pipeline's design and fan suction so demand.

5.14 Overpressure protection device

There is no need to use overpressure protection in DNCG lines.

5.14.1 Rupture disc

There is no need to use a rupture disc in DNCG lines. If rupture discs are used in the systems as pressure protection, their condition shall be controlled by an alarm in the automation system. The outlet of the rupture disc shall be aligned in a safe direction.

5.15 Concentration measurement

There is no need for continuous concentration measurement in DNCG lines.

The Finnish Recovery Boiler Committee recommends measuring the concentrations of DNCGs at sources ~~the point of generation~~ when any equipment changes are made at that point.

In addition, the Finnish Recovery Boiler Committee recommends measuring the average concentration of sulphur and oxygen in DNCGs to incineration at least once a year.



6 DISSOLVING TANK VENT GAS HANDLING SYSTEMS AT THE RECOVERY BOILER

Dissolving tank vent gases are dilute non-condensable gases primarily consisting of air and water vapour. To recover odorous sulphur compounds contained in dissolving tank vent gases and to reduce dust emissions, these gases are being incinerated in the recovery boiler. The Finnish Recovery Boiler Committee recommends incineration of dissolving tank vent gases in the recovery boiler ~~plant~~.

For instance, vent gases from the mixing tank, vent gases from the ash handling process and the mill's DNCGs can be led to the dissolving tank vent gas system.

To confirm safe operation of the recovery boiler, water must be prevented from entering the boiler ~~when in connection with the incineration of~~ dissolving tank vent gases are incinerated.

Dissolving tank vent gases are scrubbed, cooled, blown with a fan and heated up in order to vaporize ~~to promote vaporisation of any~~ water droplets before incineration. Vent gas lines are equipped with condensate ~~drains~~ outlets to prevent water from accumulating in the vent gas line and thus from entering the boiler.

6.1 Composition and amount

Dissolving tank vent gases primarily originate from leakage air, steam due to boiling of the green liquor in dissolving tank caused by the heat in the smelt, and smelt-atomising steam. The most problematic aspect with a view to the treatment are the small green liquor droplets in dissolving tank vent gases, which, when drying, tend to ~~foul/plug~~ contaminate the vent gas system.

Typical dissolving tank vent gas volumes and compositions are shown in Table 6-1



Table 6-1. Dissolving tank vent gas volumes and compositions, Rantanen 1987

Vent gas volume from dissolving tank	m ³ n/kgds	0.4 – 0.8
Vent gas temperature from dissolving tank	°C	85 - 95
Vent gas moisture content from dissolving tank	vol-%	40 - 80
Vent gas total particulates from dissolving tank	mg/m ³ n (dry)	1000 – 5000
Vent gas TRS from dissolving tank	mg/m ³ n (dry)	150 - 700
Vent gas total sulphur after scrubber	kg S/ADt	0.01 - 0.1
Vent gas total particulates after scrubber	mg/m ³ n	100 - 200
Vent gas TRS after scrubber	mg/m ³ n	1 - 10

Dissolving tank vent gases can be fed into the recovery boiler for example mixed with air or with other DNCGs. Another possibility is to introduce the dissolving tank vent gases to boiler through separate nozzles.

6.2 ~~Dissolving~~Mixing tank vent gases

~~If When leading~~ mixing tank vent gases ~~are led~~ to the dissolving tank vent gas system, attention must be paid to their fouling effect. A proper scrubber can be added to the mixing tank vent gas system.

~~If~~ mixing tank vent gases ~~are led~~ to the furnace as a system of its own, the vent gas incineration ~~recommendations and interlockings~~ must be taken into account.

6.3 Cooling/heating

Dissolving tank vent gas condensate must not be allowed to enter the furnace. The amount of water vapour entering the furnace together with dissolving tank vent gases must be minimised. ~~The dissolving tank vent gas system shall be equipped with a gas cooler heat exchanger. Gas coolereondenser~~ is used to remove water vapour, thus reducing the amount of moisture going to the furnace. It must be possible to wash the cooler while the recovery boiler is in operation.

The temperature of dissolving tank vent gases should after the ~~coolereondenser~~ preferably be 40-50 °C. ~~The cooling device shall be installed in connection with the scrubber ahead of the fan.~~



Dissolving ~~inger~~-tank vent gases should preferably be heated to a temperature of 30 °C above condensing temperature, to prevent water droplets from getting into the furnace.

6.4 **Starting/tripping ~~L~~logic ~~for~~governing ~~the~~ incineration of vent gases**

The purpose of this section is to describe the process conditions to be fulfilled by the ~~interlocking~~control logic ~~controlling~~governing the incineration of dissolving tank vent gases. The focus is on the most vital conditions that a safe automation system (SIS) must fulfil. Because of differences in equipment design, process connections and technical implementation, the ~~interlocking~~control logic of any recovery boiler will also include other ~~conditions~~variables than those presented here.

It must be possible to supervise the incineration of dissolving tank vent gases in the recovery boiler, as well as to control starting and stopping dissolving tank vent gas incineration from the same main control room where other activities related to the operation of the recovery boiler are controlled.

6.4.1 **Starting logic ~~Conditions~~ for introducingfeeding dissolving tank vent gases**

Dissolving tank vent gases can be ~~introduced~~fed into the recovery boiler furnace when the conditions below are fulfilled, Table 6-2. **Because of differences in commissioning year, equipment design, process conditions and technical implementation, the ~~interlocking~~control logic of any vent gas incinerationrecovery boiler will also include other ~~conditions~~variables than those presented here.**

Table 6-2. To admit dissolving tank vent gases to the Recovery Boiler, the following conditions should be fulfilled:

Condition	Purpose of condition
1 Boiler steam load/feed water flow is over 15% of boiler nominal load. This limit shall be defined on a case-by-case basis	Confirm sufficiently stable furnace conditions to oxidise <u>TRS compounds</u> gases <u>NGGs</u> (temperature, retention time)
2 Dissolving tank vent gas fan is switched -on	Verify that there is sufficient flow in the duct to prevent flame propagation backward in the pipeline (burn-back)
3 Sufficient pressure in duct system ahead of shut-off valve	Verify that there is sufficient flow in the duct to prevent flame propagation backward in



- | | | |
|---|---|--|
| | | the pipeline (burn-back) |
| 4 | Temperature after cooling (ahead of heater) below temperature limit | Reduce gas volume and amount of water vapour and gas volume |
| 5 | Temperature before introductionfeed to boiler/after heater above temperature limit | Prevent water droplets from entering the boiler |
| 6 | CondensateTrap pocket level in NCG system (if exists) below upper limit | Prevent condensate from entering the furnace. Indicate fault in the condensate system preventing condensate drainingremoval |
| 7 | Duct flow before shut-off valve sufficient (if separate nozzles) | Verify that there is sufficient flow in the duct to prevent flame backward progression in the pipeline (backfire) |

6.4.2 ~~Tripping logic for introducing~~Interrupting feeding of dissolving tank vent gases

The ~~introductionfeed~~ of dissolving tank vent gases into the ~~recovery~~ boiler furnace has to be ~~stoppedinterrupted~~ if any of the conditions below is fulfilled, Table 6-3. Because of differences in commissioning year, equipment design, process conditions and technical implementation, the ~~interlockingsecontrol~~ logic of any ~~vent gas incinerationrecovery boiler~~ will also include other ~~conditionsvariables~~ than those presented here.

Table 6-3. Any of the following conditions will divert dissolving tank vent gases from the Recovery Boiler:

Condition	Purpose of condition
1 Boiler steam load/feed water flow is below 15% of boiler nominal load. This limit shall be defined on a case-by-bas case basis	Confirm sufficiently stable furnace conditions to oxidise TRS compounds gases NCGs (temperature, retention time)
2 Emergency shut-down tripped activated	Prevent introductionfeeding of NCGs when the boiler is not in accepted condition -use
3 Dissolver tank vent gas fan stopped	Prevent flame flame propagation backward in the pipeline (burn-back)
4 Temperature after cooling (ahead of heater) rises above temperature limit	Reduce gas volume and amount of water droplets and gas volume
5 Temperature before introducingfeeding into boiler/after heater falls below temperature limit	Prevent water drops from entering the boiler
6 Trap Condensate pocket level above upper limit	Prevent condensate from entering the furnace. Indicate fault in the condensate system preventing condensate drainingremoval

6.5 Fan

The fan structure must be sufficiently tight and the fan must be equipped with a condensate ~~drainingremoval~~ system.



6.6 Ducts

Dissolving tank vent gases must be prevented from escaping into the boiler ~~house~~room and the length of the duct inside the boiler ~~house~~room shall be minimised.

Because of the risk of corrosion, the ducts and nozzles for dissolving tank vent gas incineration shall be made from stainless steel EN1.4301 (SS2333) or better.

6.6.1 Flow

The flow of dissolving tank vent gases can be measured. The flow measuring point shall be close to the point where dissolving tank vent gases are ~~introduced~~~~fed~~ into the boiler.

6.6.2 Potential balancing

No potential balancing is required for dissolving tank vent gas ducts.

6.7 Condensate ~~draining~~removal

Dissolving tank vent gas lines must be provided with condensate ~~drain~~outlets. Condensates shall primarily be conducted to the dissolving tank or to the treatment system for ~~fouled~~contaminated condensates.

In designing condensate ~~drains~~removal, special attention shall be paid for example to the size, number and location of the ~~drains~~inlets. The last condensate drain is recommended to locate as close to the recovery boiler as possible so that the length of the duct from this drain to the furnace is minimised. Drain lines closest to the boiler should have condensate pockets with limit switches. The last condensate drainoutlet should placed as close to the recovery boiler as possible to minimise the length of the duct from the outlet to the furnace.

6.7.1 Size of condensate ~~drains~~removal lines

Because of the risk of plugging, the recommended minimum diameter for condensate ~~drain~~outlets and condensate lines starting from these is 50 mm. The condensate drain line closest to boiler should be at least 100 mm. -If there is a



possibility of additional water (e.g. washing water) feed into the system, then condensate drain lines shall be sized accordingly.

6.7.2 Duct slopes and locations of condensate ~~drains~~~~removal~~~~outlets~~

Ducts and pipes must slope towards the condensate ~~drains~~~~outlets~~. Condensate ~~draining~~~~removal~~ against flow is very difficult. The recommended slope for ducts, elsewhere than in the recovery boiler plant, in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:25.

Because of the risk of water getting into the furnace, the preferred slope for ducts in recovery boiler plants in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:1 or more. A condensate ~~drain~~~~removal~~~~point~~ must be installed before every point where the ~~vent gas~~~~NCG~~ line angles upward. If the recommended slopes cannot be achieved in some duct sections, the condensate outlets in this part of the system must be designed with special care.

6.7.3 Water seals

Condensate ~~drain~~~~removal~~ lines must be provided with water seals ~~or~~ a common seal tank (condensate collection tank). In this way, NCGs are prevented from leaking via the condensate ~~drains~~~~outlets~~.

To prevent water seals from drying out, they shall be equipped with level indicators or with a continuous water flow controlled by flow indicators.

To confirm that the pressure in the dissolving tank vent gas duct will not be sufficient to drain the water seal, the difference between the water levels in the water must be allowed to vary by at least 10 kPa, or by an amount corresponding to the duct's design pressure. 10 kPa corresponds to a water level variation of 1 m in the water seal.

~~If condensates are collected into a pumping tankbasin, the level difference in the tankbasin must be allowed to vary by at least 10 kPa, or by an amount corresponding to the duct's design pressure.~~



6.8 Scrubber

The dissolving tank vent gas system shall incorporate an effective cooling scrubber to remove dust and water vapour. It must be considered possible to wash the scrubber and the main equipment ~~heat exchangers cooling the scrubber circulation~~ while the recovery boiler system is in operation.

~~6.9 Cooling~~

~~The dissolving tank vent gas system shall be equipped with a cooler connected to the scrubber or separate. The cooler is used to remove water vapour and thus reducing the amount of gas going to the furnace. It must be possible to wash the cooler while the system is in operation.~~

~~6.10~~ 6.9 Valves/dampers

The valves/dampers for dissolving tank vent gas incineration shall be selected with care. Special attention shall be paid to the valve sealing characteristics. Normally, effectively sealing ~~throttle~~ valves/dampers are used.

The dissolving tank vent gas line should have automatic shut-off valve with “fail-close” actuator which will close vent gas introduction to the recovery boiler automatically if operating energy is turned off. At the same time valve of the bypass duct must open (fail-open). Valves must be provided with limit switches indicating position to the automation system.

Recommended material for the fan is EN 1.4301 (SS2333) or equivalent.

~~6.11~~ 6.10 Bypass

The bypass for dissolving tank vent gases must be led to as high a point as possible. Normally, the bypass duct is led to the roof of the recovery boiler plant or to the chimney stack.

~~The automatic shut-off valve must open by itself if operating energy is turned off.~~



6.126.11 Droplet separator

A droplet separator after gas cooling is recommended in the dissolving tank vent gas system. Droplet separator removes possible water droplets from gas before incineration.

6.136.12 Flame arrester

There is no need for a flame arrester in dissolving tank vent gas ducts.

6.146.13 Overpressure protection

The dissolving tank vent gas system must be able to withstand the overpressure generated by the fan. In addition, there must be a separate duct from the dissolving tank to the roof of the recovery boiler plant which is opened by overpressure in the dissolving tank.

6.14.16.13.1 Rupture disc

There is no need for a rupture disc~~s~~ in dissolving tank vent gas ducts.

6.156.14 Concentration measurement

There is no need for continuous concentration measurement of dissolving tank vent gases.

6.15 Washing of droplet separator and vent gas heater

Dissolving tank vent gases include a lot of fine dust which plug droplet separators, heater and ducting. When water washing of these equipment is carried out then extra care must be paid on proper condensate/water removal. If condensate pockets limit switches are indicating level the washing must be stopped. During extensive washing, vent gases shall be by-passed.

7 INCINERATION OF CNCG IN THE RECOVERY BOILER

Concentrated non-condensable gases are a low-calorific fuel and the system for incinerating such gases must comply with the same recommendations and regulations concerning safety and ~~control~~logics as for any other explosive gases.

Sources for CNCGs are for example non-condensable gases from the vacuum pit of the evaporation plant, gases from the stripper and methanol distilling plant, vent gases from the foul condensate tank, gases from cooking plant condensers, gases from turpentine system and gases from pressurised firing liquor tank~~liquor concentration~~. Every effort must be made to prevent leakage of air into CNCGs under any circumstances.

~~The ATEX Directive must be observed in the design of CNCG systems.~~



Figure 7-1. Bottom~~Lower~~ part of the recovery boiler furnace with a separate ~~incinerator for~~ CNCG burner opening in the centre.

7.1 Composition and amount

The total amount of sulphur in CNCGs in a modern chemical pulp mill is in the range of 2–4 kgS/ADt. This amount is increased considerably if black liquor is evaporated to a high dry solids content (over 80%). For process reasons, the amount of CNCGs may vary widely at different [sources points](#) in time. Typical departmental amounts of CNCGs collected from different [sourcespoints](#) in a chemical pulp mill are shown in Table 7-1.

Table 7-1. Typical amounts of CNCGs collected in different departments

Department	kg S/ADt	m ³ n/ADt
Batch cook blowing	0.4 – 0.8	5 - 15
Batch cook gassing	0.1 – 0.2	1.0 – 3.0
Continuous cooking	0.1 – 0.4	1.0 – 4.5
Stripper	0.5 – 1.0	15 –25
Evaporation plant	0.4 – 0.8	1 – 10
Methanol processing	0.5 – 1.0	1.0 – 2.0
Black liquor heat treatment	2 – 3	1.5 – 3.0
Concentrator	2 – 5	1.5 – 6.0

CNCGs contain significant amounts of sulphur compounds and ammonia. Examples of analyses of CNCGs are shown in Table 7-2. Depending on their components, CNCGs are classified as extremely toxic, toxic or inflammable. Heat values of CNCGs components are presented in Table 7-3.

Table 7-2. Examples of analyses of CNCGs

Component	Mill A ppm	Mill B ppm	Mill C ppm	Mill D % vol
H ₂ S	*	50 000	81 300	<u>1.5</u>
CH ₃ SH	80 900	110 000	188 300	<u>2.0</u>
(CH ₃) ₂ S	22 000	50 000	116 000	<u>2.0</u>
(CH ₃) ₂ S ₂	800	30 000	3 000	<u>1.5</u>
C ₁₀ H ₁₆	1 900	*	*	<u>0.1</u>
CH ₃ OH	*	*	*	<u>0.2</u>
O ₂	*	*	*	<u>9.0</u>
NH ₃	*	*	*	*
H ₂ O	20 000	330 000	*	<u>6.0</u>
CO ₂	*	*	*	*

*) not analysed or below detection limit



Table 7-3. Examples of the heat values of the components of CNCGs, BLRBAC

Component	kJ/kg(dry)
H ₂ S	15 280
CH ₃ SH	26 100
(CH ₃) ₂ S	30 890
(CH ₃) ₂ S ₂	23 630
Turpentine	41 560
Methanol (gas)	22 720

Concentrated non-condensable gases are usually not cooled or heated up in the same way as DNCGs. An ejector or a liquid ring compressor is used to transfer CNCGs. After these devices in the [CNCG](#) system are situated the droplet separator, [valve group](#), flame arrester and rupture disc before incineration.

CNCG incineration and collection systems must be designed to confirm that leaks from the gas lines are impossible, that water cannot enter the furnace with the gases and that the risk of fire/explosion is minimised.

7.2 Starting/tripping~~Control~~ logic for governing incineration of CNCGs

It must be possible to supervise the incineration of CNCGs in the recovery boiler and to control starting and stopping of the incineration of CNCGs from the same main control room where other activities related to the operation of the recovery boiler are controlled.

The purpose of this section is to describe the process conditions to be fulfilled by the ~~interlocking~~control logic ~~controlling governing~~ the incineration of CNCGs. The focus is on the most vital conditions that a safe automation system must fulfil. Because of differences in equipment design, process conditions and technical implementation, the ~~interlocking~~control logic of any [CNCG burning](#)~~recovery boiler~~ will also include other conditions~~variables~~ than those presented here.

This recommendation does not cover interlockings in the collection points of CNCGs, though these are often partly implemented in the same system as interlockings related to incinerations.



7.2.1 **Starting logic**~~Conditions~~ for starting the **CNCG** incineration~~nr of CNCGs~~

The incineration of CNCGs can be started when e.g. the conditions below are fulfilled, Table 7-4. **Because of differences in commissioning year, equipment design, process conditions and technical implementation, the ~~interlocking~~control logic of any **CNCG burning**~~recovery boiler~~ will also include other ~~conditions~~variables than those presented here.**

Table 7-4. To ~~introduce~~~~admit~~ CNCGs to the Recovery Boiler, the following conditions should be fulfilled:

Condition	Purpose of condition
1 Emergency shutdown not activated	Prevent CNCG feed when boiler is not in accepted condition
2 Boiler furnace heat loading is above 0.7 MW/m ² . This limit shall be defined on a case-by-basecase basis	Confirm sufficiently stable furnace conditions to oxidise TRSCNCGs compounds gases (temperature, retention time)
3 Burner support flame is on burning	Confirm sufficient incineration conditions in the burner
4 Flame scanner detector shows that the re is flame is burning in support fuel	Confirm burning of flame on
5 Burner combustion air inlet pressure and flow above minimum	Confirm sufficient air flow to burner
6 Burner opening is not plugged (operator should check)	Confirm a safe CNCG incineration

7.2.2 Capacity limit

The incineration of CNCGs requires sufficient temperature and retention time in the furnace to allow the gases to be combusted into carbon dioxide, water vapour and sulphur dioxide. A high enough temperature is achieved by raising the boiler load sufficiently [and/or using support fuel](#). The minimum load as steam flow is determined by the equipment supplier. Because nominal load is often a somewhat obscure concept, it is recommended that this capacity should correspond to a furnace [heat](#) loading of at least 0.7 MW/m²_{bottom}. An example of how to calculate the furnace [heat](#) load is given in Appendix I.

7.2.3 **Tripping logic for the**~~Interrupting~~ CNCG incineration

The incineration of CNCGs is interrupted for example if any of the conditions below is fulfilled, Table 7-5. **Because of differences in commissioning year, equipment design, process conditions and technical implementation, the**



~~interlocking control~~ logic of any ~~CNCG burning recovery boiler~~ will also include other ~~conditions variables~~ than those presented here.

Table 7-5. Any of the following conditions will divert CNCGs from the Recovery Boiler:

Condition	Purpose of condition
1 Emergency shut-down activated tripped	Prevent CNCG feed when boiler is not in accepted condition use
2 Boiler furnace heat loading is below 0.7 MW/m². This limit shall be defined on a case-by-case basis	Confirm sufficiently stable furnace conditions to oxidise TRS compounds gases (temperature, retention time)
3 High Level controllers tripped in CNCG condensate draining removal system	Indicate a malfunction in the condensate removal system preventing condensate draining removal. Prevent water from entering the boiler
4 Pressure of CNCG before burner incinerator is drops below low limit	Prevent flame flame propagation backward progression in the pipeline (burn backfire)

It is recommended that the incineration of CNCGs is interrupted if for example any of the following alarms is activated, Table 7-6:

Table 7-6. CNCGs alarms

Alarm	Purpose of alarm
1 Boiler CO emission exceptionally high (over 1000 ppm for more than 1 min)	There is a noticeable disturbance in incineration black liquor burning
2 Boiler house room hydrogen sulfide sulphur concentration alarm (H ₂ S) is activated tripped	Check if there is a leak in the CNCG system
3 Combustion black liquor dry solids drops below the alarm low limit causing alarm	Confirm a stable and safe burning Confirm incineration
4 Flame arrester pressure difference over upper limit	Flame arrester plugged/becoming plugged
5 CNCG flow to burner incinerator drops below alarm low limit (the actual flow is monitored secured by pressure)	Verify CNCG introduction feed to incineration burner

7.2.4 Burner and use of support flame

Concentrated non-condensable gases can only be incinerated in an appropriately designed separate burner. The burner for CNCGs shall be equipped with a separate air register and a separate lance for support/pilot fuel, which may be for example oil, gas, methanol, turpentine or tar oil. The use of DNCGs as air in the burner of NCGs is not recommended because of their moisture and impurities.



The electrical and automation systems of the CNCGs incinerator must meet the requirements for electrical and automation systems related to gas incineration. If oil is used as auxiliary fuel, the corresponding requirements for oil must be met.

The incineration of CNCGs requires a sufficiently high incineration zone temperature. In general, a temperature of 900 °C is required. The temperature requirement is met if CNCGs are led below the liquor guns when the boiler operation is stable and boiler load sufficient. If CNCGs are led above the liquor guns, attention must be paid to leading them to a place in the boiler where this temperature is exceeded. Another possibility is to use a support flame with sufficient capacity.

With small loads the purpose of the support flame is to ensure stable ignition ~~promote ignition~~ and incineration of CNCGs. ~~The burner support flame must not fall off or become extinct.~~ The ~~furnace~~ loading of the support flame shall be at least 10% of the maximum ~~furnace~~ heat input ~~loading~~ of the burner. The support flame shall be supervised by a flame ~~scanner~~ controller.

The ~~CNCG~~ concentrated non-condensable gas support fuel lance shall be equipped with local control ~~pane~~ le ~~abinets~~ from where support ~~fuel burning~~ incineration is always started. The incineration of CNCG can then be started from the control room. Both of the above-mentioned operations can be stopped from the control room or from local control panel.

Figure 7-3 shows an example of a CNCG burner photographed ~~both~~ from the outside and from the inside of furnace. The CNCG burner, methanol lance, flame scanners, flame arrester and support fuel lance are shown.



Figure 7-23. Inside and outside of a CNCG burner.

When the boiler operates at a furnace heat loading of over 0.7 MW/m^2 , two attempts can be made to ignite the support flame from the control room. **However, this limit shall be defined on a case-by-case basis.** The first ignition attempt shall always be made by using local control box at the burner.

When the support flame scanner doesn't detect flame goes out, CNCGs may be led to the recovery boiler for a while (for example 3 minutes) before shutting the quick-acting valves. During this time the start-up sequence for the alternative burning location can be started.

No support flame is needed when the following conditions apply, Table 7-7. **The condition may vary depending on commissioning year, equipment design, process conditions, and technical implementation.**

Table 7-7. Conditions for CNCG-incineration without a support flame

Condition	Purpose of condition
1 Boiler furnace <u>heat</u> loading is above 1.5 MW/m^2 . This limit shall be defined on a case-by-<u>base</u> case basis	Confirm sufficiently stable furnace conditions (temperature, retention time)



7.3 Pipeline system

The length of the pipeline system for CNCGs inside the boiler ~~house~~room should be kept as short as possible. The pipelines should be installed with a view to avoiding critical locations, such as the dissolving tank and its environment, weak corners, emergency exits and stairways.

The ~~stipulations of~~ pressure vessel and chemical handling legislation shall be taken into account in the design and installation of the pipeline system.

Pipelines should preferably be made of acid proof steel EN1.4436 (SS2343) or corresponding. Certain compounds in NCGs, such as turpentine and methanol, are strong solvents, so the use of glass fibre and plastic parts is prohibited.

7.3.1 Steam purging of pipelines

When incineration is stopped, the pipelines for CNCGs should be flushed from the shut-off valves to the boilers by purging them with an inert medium. In practice, the pipelines are always purged with steam. ~~Inert media are for example steam and nitrogen.~~ Purging with air is not recommended, because an explosive mixture is created at the interface between the purging air and the CNCGs to be purged. ~~In practice, the pipelines are always purged with steam.~~

Steam purging implies a risk of water condensate in the lines reaching the furnace, thus causing a dangerous situation. When designing the pipeline, attention should be paid to leading water possibly condensing in the pipeline to the condensate collection system.

7.3.2 Pipeline pressure measurements

The pipeline for CNCGs shall be equipped with pressure gauges after the ejector and immediately ahead of the incinerator for CNCGs. An alarm is activated if overpressure or minimum pressure is detected in the pipeline.

Plugging of the pipeline can be supervised based both on flow rate and pressure measurements.



7.3.3 Potential balancing

Potential balancing is carried out for CNCG lines. For instance in flanges, hatches and connections where a galvanic isolation may be formed, for example due to the isolation materials used, potential balancing must be performed. Potential balancing shall be made according to the given instructions and verified with a field log.

7.4 Condensate ~~draining~~removal

Pipelines for CNCGs shall be provided with condensate drains. The condensates contain large amounts of malodorous compounds. The condensates shall be collected into a pumping tank and pumped ~~from the there~~via the tank to ~~stripping of~~ ~~fouled~~contaminated condensate ~~tanks~~ in the evaporation plant.

7.4.1 Size of condensate ~~draining~~removal lines

Because of the risk of condensate overflow and plugging, the recommended minimum pipeline diameter for condensate ~~drain~~removal connections and pipelines starting from these is 50 mm. Connections and condensate pipes installed outdoors shall be equipped with electric tracing ~~if there~~because of ~~is a~~the ~~risk~~danger of freezing.

7.4.2 Pipe slopes and locations of condensate ~~drains~~removal-connections

Pipelines must slope towards the condensate ~~drains~~removal-connections. The recommended slope for pipes, elsewhere than in the recovery boiler plant, when condensate travels in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:25. Pipeline diameters shall be large enough to prevent flooding of condensates in the system.

Because of the risk of water entering the furnace, the aimed slope of pipes in the recovery boiler plant in the direction of the gas flow is 1:100 and counter to the direction of the gas flow 1:1 or more. In pipeline sections where the recommended slopes cannot be achieved, special care shall be taken in designing the condensate drainage system for the pipe in question.



When designing condensate ~~draining~~~~removal~~ systems, attention should be paid e.g. to the size, amount and location of ~~drain~~~~connections~~. The last condensate drain shall be placed so close to the recovery boiler that the length of the duct from this drain to the furnace is minimised. Special attention shall be paid to the ~~condensate draining removal of condensate from CNCGs in of~~ pressure retaining steam lines.

7.4.3 Water seals

To prevent ~~CNCGs~~ from leaking via the condensate drain~~sage~~~~connections~~, the condensate ~~drainage~~ lines must be equipped with water seals and a pumping tank, or a common condensate seal tank.

To confirm that the pressure generated by the ejector in the pipelines for CNCGs will not rise so high that it causes the water seal to be emptied, the rated values of the ejector (overpressure and underpressure), as well as the process requirements (for example boiler system bottling) must be taken into account in the dimensioning of the water seal.

The water seal dimensions should be large enough so that the suction of the ejector does not drag droplets with it.

The water level in the water seal must be monitored. One solution is to lead a constant flow to the water seal, which is verified by flow measurements.

7.5 Valves

The valves of the CNCG incineration system shall be selected with care. Special attention shall be paid to their sealing characteristics.

There should be two quick-~~acting~~~~closing~~ valves in the inlet line for CNCGs to the ~~burner~~~~furnace~~. Between them there should be a ventilation pipe with corresponding valve. The CNCG line shut-off valves should be equipped with “fail-close” actuator which will close CNCG introduction automatically if operating energy is turned off. The valve of the ventilation pipeline between the shut-off valves must open at the same time (fail-open). Valves must be provided with limit switches indicating position to the automation system.



It must be possible to close the pipelines for CNCGs to the boiler house, also from outside the boiler house, with so-called emergency shut-off valves. The need for closing the pipelines from the outside arises from the fact that in a danger situation the CNCG flow inside the boiler house must be stopped. ~~pipelines cannot necessarily be closed from inside the boiler house. In addition, there is a need to prevent CNCGs from flowing into the boiler house in a situation where for example a valve in the boiler house is leaking.~~

The valves selected must comply with the corresponding regulations for inflammable gases, for example fire endurance. Recommended material for the valves is acid proof steel EN 1.4436 (SS2343) or equivalent.

~~7.5.1 Shut-off valves~~

~~The pipelines for CNCGs shall be equipped with tightly sealing shut-off valves. Shut-off valves shall be provided with limit information.~~

~~It must be possible to close the pipelines for CNCGs to the boiler room, also from outside the boiler room, with so-called emergency shut-off valves. The need for closing the pipelines from the outside arises from the fact that in a danger situation the CNCG pipelines cannot necessarily be closed from inside the boiler room. In addition, there is a need to prevent CNCGs from flowing into the boiler room in a situation where for example a valve in the boiler room is leaking.~~

~~Shut-off valves in the CNCGs incineration system must close without external operating energy. The valves of the ventilation pipeline between the shut-off valves must open at the same time.~~

7.6 Transfer of CNCGs

Fan is not allowed to use~~Blower must not be used~~ for the transfer of CNCGs. Instead, the use of an ejector or liquid ring compressor is recommended. The temperature of the steam ejector and of the steam must not exceed 200 °C (corresponds to a steam pressure of approximately 12 bar(y)), because of the auto-ignition temperature of the gas components in CNCG gas.



If necessary, the CNCGs ejector/liquid ring compressor can be placed inside the recovery boiler building~~plant~~.

7.6.1 Pressure-control~~retaining~~ steam

Low pressure steam is used for controlling CNCG pressure before burner~~as pressure retaining steam~~. The functioning of the ~~drainage of the~~ steam lines drains must be monitored at all times to prevent water from entering the furnace.

7.6.2 Heating/cooling

Generally, the CNCGs are not heated or cooled ~~in connection with incineration~~.

7.6.3 Droplet separator

A droplet separator should be used after ejector/liquid ring compressor to remove possible water droplets from the gas.

~~7.6.3~~7.6.4 Flame arrester

A flame arrester must be used for CNCGs. The purpose of the flame arrester is to ~~prevent~~protect ~~against flame propagation or burn back~~ ~~stop the fire front from advancing~~ along the pipeline from the furnace towards the incoming CNCGs.

If there is a system for cleaning the flame arrester installed, the cleaning pipelines must not be permanently connected. The pressure difference of the flame arrester must be measured and the flame arrester equipped with an alarm.

~~7.6.4~~7.6.5 Underpressure protection

Underpressure protection by using air as make-up gas is not allowed for CNCGs. It is recommended that pipelines and equipment be dimensioned so that no underpressure protection is needed.

~~7.6.5~~7.6.6 Overpressure protection

Overpressure protection opening direct into the boiler house is not allowed for CNCGs in the boiler building. The flow of CNCGs caused by breaking~~tripping~~ of the overpressure protection shall be conducted to a point where it does not represent a hazard and where its negative impacts are minimised. Attention should



be paid to draining of pipes with overpressure leading to ~~outside the exterior~~ to prevent freezing. ~~Breaking~~Tripping of the overpressure protection must activate an alarm.

The most common type of overpressure protection is a rupture disc which activates an alarm.

~~7.6.6~~7.6.7 Rupture discs

~~Piping section shall be equipped with rupture discs if a process upset condition or equipment malfunction could lead to pressure build up over the design pressure. If there is a possibility of the pressure between two shut-off valves rising above the design pressure of the pipes concerned, this section of the pipe shall be provided with rupture discs.~~ This kind of pressure is for example the pressure generated by medium-pressure steam blowing. ~~Breaking~~Tripping of the rupture disc must activate an alarm.

The recommended material for rupture discs is acid-proof steel or graphite.

Rupture discs shall be installed in places where they will not represent a hazard to people working on the operating floors.

7.7 Concentration measurement

There is no need for continuous measurement of the concentrations of CNCGs.

The Finnish Recovery Boiler Committee recommends that the concentrations of CNCGs be measured at the point of origin, if equipment changes have been made at this point.

In addition, the Finnish Recovery Boiler Committee recommends that the content of sulphur and possibly also of oxygen to incineration be measured at least once a year.

7.8 Alternative location

~~A condition for incinerating CNCGs in the recovery boiler is that there is an alternative~~ There shall be an alternative location/system where the gases can be



diverted automatically ~~if in situations when~~ the incineration of CNCGs in the recovery boiler is interrupted. ~~There are multiple process equipment which can be considered as alternative location, for example power boiler, auxiliary boiler, flare, lime kiln and stack.~~ The intention of diverting to an alternative ~~location~~outlet is to ~~ensure low odour emissions. prevent CNCGs from escaping into the environment.~~

~~The automatic valve used to divert CNCGs from recovery boiler to the alternative location must be fail-open i.e. open automatically, if the supply of operating energy is turned off. Valve must be equipped with limit switches indicating position to the automation system.~~

~~7.8.1~~ **Valves**

~~The valves for CNCGs in the recovery boiler plant used to divert gases to the alternative outlet/incinerator must open automatically, if the supply of external operating energy is interrupted.~~

~~7.8.2~~7.8.1 **Steam purging**

It must be possible to steam-purge the pipeline of the ~~alternative location~~backup /incineration system for CNCGs.

~~7.8.3~~7.8.2 **Location of flare**

If the alternative outlet/incinerator for CNCGs is a flare, it should preferably be located on the roof of the recovery boiler plant.



8 METHANOL/TURPENTINE INCINERATION IN THE RECOVERY BOILER

In pulp mills, liquefied methanol/turpentine ~~can be burnt is, in addition to in the recovery boiler, also incinerated~~ in the lime kiln, in a separate incinerator boiler for NCGs, in the power boiler and in a flare. If there is no backup incineration system, methanol/turpentine storage capacity corresponding to for example 24 hours' production must be available. ~~Liquefied methanol and turpentine may be fired separately, or in a mixture. blended combination.~~ If turpentine is mixed to methanol characteristics of these streams must be taken into consideration for example turpentine heat value is ~40 MJ/kg and methanol ~20 MJ/kg and turpentine's tendency to collect on top of the storage tanks due to poor solubility. Typically turpentine is led to the suction side of methanol pump.

The methanol and turpentine storages must comply with applicable regulations for storing inflammable liquid chemicals. Tanks ~~It~~ must not be placed in the boiler house ~~room~~.

8.1 Starting/tripping ~~Control~~ logic of the governing methanol/turpentine incineration

The purpose of this section is to describe the process conditions to be fulfilled by the interlocking ~~control~~ logic controlling governing the operation of the methanol and/or turpentine burner ~~incinerator~~, in addition to those of a normal gas/oil burner. The focus is on the most essential conditions that a safe automation system must fulfil as a minimum. Because of differences in equipment design, process connections and technical implementation, the interlocking ~~control~~ logic of any recovery boiler will also include other conditions ~~variables~~ than those presented here.

Methanol/turpentine can be used in CNCG burners ~~incinerators~~ as support ~~backup~~ fuel and as fuel in start-up burners. When using methanol/turpentine as fuel in the start-up burner, the temperature in the furnace must be sufficiently high. Methanol firing ~~The incineration of methanol~~ can be started earlier than the incineration of CNCGs. In this case the interlocking conditions must be defined on a case-by-case basis.



To ensure safe and stable burning the methanol/turpentine quality (e.g. water content) must be monitored for example measuring methanol/turpentine density. Normal density of methanol is 0.80 g/cm³ and turpentine 0.85 g/m³. If certain limit is exceeded, the stream must be diverted from the burner.

It must be possible to supervise the ~~burning~~incineration of methanol and turpentine in the recovery boiler as well as to control starting and stopping methanol/turpentine ~~burning~~incineration from the same main control room where other activities related to the operation of the recovery boiler are controlled.

8.1.1 Starting logic~~Conditions~~ for ~~start of~~ methanol and turpentine ~~burning~~incineration

Methanol and turpentine ~~burning~~incineration can be started in the recovery boiler when the following conditions are fulfilled, Table 8-2. Because of differences in commissioning year, equipment design, process conditions and technical implementation, the interlocking logics of any methanol/turpentine burning recovery boiler will also include other conditions than those presented here.

Table 8-~~12~~. To admit methanol/turpentine to the recovery boiler, the following conditions should be fulfilled:

Condition	Purpose of condition
1 Fire signal in boiler	Prevent odour nuisances
2 Methanol/turpentine lance in place and connected	Confirm the right position of the methanol/turpentine lance
3 Methanol/turpentine pressure above minimum pressure (delay)	Confirm <u>proper atomization of methanol sufficient supply of methanol/turpentine to the burner</u>
4 Burner air pressure above low air pressure limit	Confirm sufficient supply of incineration air to the burner as well as sufficient atomising
5 Pressure of atomising air sufficiently high	Confirm sufficient atomisation of methanol and prevent methanol from entering compressed air pipelines
6 Burner d umper in ignition position	Confirm sufficient incineration air for ignition
7 Flame scanner detector shows flame (ignition delay)	Confirm that the fuel flame is burning
8 It has been checked that the burner opening orifice is not plugged blocked	Confirm a safety of methanol/turpentine burning incineration
9 Methanol/turpentine density below upper limit	Confirm that burner heat value does not drop too much <u>(too much water)</u>



8.1.2 Tripping logic for~~Events that will interrupt~~ methanol and turpentine incineration

Methanol and turpentine incineration is ~~stopped~~~~interrupted~~ if any of the following conditions is fulfilled, Table 8-3. Because of differences in commissioning year, equipment design, process conditions and technical implementation, the interlocking logics of any methanol/turpentine burning recovery boiler will also include other conditions than those presented here.

Table 8-~~23~~. Any of the following conditions will divert methanol/turpentine from the recovery boiler:

Condition	Purpose of condition
1 Fire signal is lost	Prevent odour nuisances
2 Methanol/turpentine lance not in place	Confirm the right position of the methanol/turpentine lance
3 Methanol/turpentine pressure below minimum pressure (delay)	Confirm sufficient supply of methanol/turpentine to the burner
4 Burner air pressure drops below low air pressure limit	Confirm sufficient supply of incineration air to the burner as well as sufficient atomising
5 Pressure of atomising air too low	Confirm sufficient atomisation of methanol and prevent methanol from entering compressed air pipelines
6 Methanol/turpentine density rises above the upper limit	Confirm that burner heat value does not drop too much
7 Flame scanner detector indicates <u>no flame</u> that the flame has gone out	Confirm that the <u>fuel flame</u> is burning

8.2 Electrical and automation systems of the burner~~incinerator~~

The electrical and automation systems of the methanol/turpentine burner must fulfil the conditions for corresponding systems related to oil/gas burning.

The methanol and turpentine lance are purged with steam or air. ~~In that case fire in boiler information must be available.~~ There must be boiler fire signal condition on before purging can be started (start-up burners on).

8.3 Valves

The valves for methanol and turpentine incineration systems shall be selected with care. The lines should have quick-acting shut-off valves with “fail-close” actuator which will close methanol/turpentine flow automatically if operating energy is turned off. Shut-off valves must be equipped with limit switches indicating shut-off to the automation system.



~~It must be possible to close the lines also from outside the boiler house. The need for closing the pipelines from the outside arises from the fact that in a danger situation the methanol/turpentine pipelines cannot necessarily be closed from inside the boiler house. In addition, there is a need to prevent methanol/turpentine from flowing into the boiler house in a situation where for example a valve in the boiler house is leaking.~~

The valves selected must comply with the corresponding regulations for inflammable gases, for example fire endurance. Acid-proof stainless steel EN 1.4336 (SS2343), or corresponding, is recommended for the valve construction material. The valve sealing material shall be resistant to dissolution (methanol and turpentine are solvents). Special attention shall be paid to the selection of sealing material.

8.3.1 Quick-closing valve

~~It must be possible to close the lines also from outside the boiler room. The need for closing the pipelines from the outside arises from the fact that in a danger situation the methanol/ pipelines cannot necessarily be closed from inside the boiler room. In addition, there is a need to prevent methanol/turpentine from flowing into the boiler room in a situation where for example a valve in the boiler room is leaking.~~

~~The quick-closing valves of methanol/turpentine incineration system close without external operating energy.~~



9 ACTIONS IN SPECIAL SITUATIONS

This section deals with the actions to be taken in situations where the recovery boiler is not operating at stable load.

9.1 Actions ~~during in connection with~~ disturbances

The Finnish Recovery Boiler Committee recommends that the incineration of CNCGs, ~~and~~ DNCGs, vent gases, methanol and turpentine in the recovery boiler shall be stopped and they should be the gases diverted to the backup incineration system in connection with any major disturbance situation at the recovery boiler.

If the boiler main interlock is initiated activated tripped, the incineration of methanol, turpentine, dissolving and mixing tank vent gases, CNCGs and DNCGs in the recovery boiler shall be stopped.

If there is a risk of explosion or fire the import of methanol, turpentine, dissolving and mixing tank vent gases, CNCG and DNCG in the recovery boiler building shall be stopped. Therefore it is not recommended to continue flow of CNCG through boiler house to back-up incinerators.

9.2 Actions ~~during in connection with~~ shut-downs

~~In connection with~~ During a shut-down, the incineration of NCGs in the recovery boiler is recommended to be stopped when approaching interlocking limits; see tables 5.5, 6.3 and 7.5. and gases are diverted to alternative location. When the incineration of CNCGs is stopped, the pipeline – from shut-off valves to the boiler – shall be purged with steam and when the incineration of DNCGs is stopped, the ducts are recommended to be flushed with air. If possible, the collection of DNCGs should be operated during shut-downs.

9.3 Actions ~~during in connection with~~ start-ups

Non-condensable gas handling systems should preferably be kept in operation also during shut-downs. If there has been a stop break in the incineration of NCGs in the recovery boiler, the following items must be checked before the NCG system is started up again:



- that any maintenance or repair work done during the shut-down is completed and that NCG equipment is no longer being serviced/maintained
- that the equipment and pipelines subject to maintenance or repair have been appropriately installed and cleaned of any debris
- that the valves connected to equipment and pipelines subject to maintenance or repair are in correct conditions and position for start-up
- that the equipment and pipelines subject to maintenance or repair are ready for use and tightly sealed. The ~~tightness~~sealings must be checked by test runs.

Before start-up, all DNCG lines that were out of use during the shut-down have to be ventilated and CNCG lines purged with steam and, if necessary, with nitrogen. The purpose of ventilation is to confirm that any NCGs that have leaked into or condensed inside ducts and other equipment will not cause harm or risk situations.

Before start-up, all condensate tanks must be checked for overflows or drying-out.

Before start-up, all water seal ~~leakage~~ fresh water flows must be checked. ~~and activated. to prevent water seals from drying out.~~

After start-up, when equipment, piping and connection flanges are warmed up to process temperature, it is good to check once again the tightening of bolts. Check also that groundings are connected.

9.4 Actions during ~~outages~~shut-downs

During ~~outages~~shut-downs, NCGs must be prevented from leaking into the boiler building. The purpose of separating NCG lines is to prevent gases from leaking into and accumulating within ducts and pipelines inside the boiler building during ~~outages~~shut-downs.

9.4.1 Instructions for preparing NCG system for shut-down or maintenance

Non-condensable gas lines, in the same way as other lines in the recovery are~~department~~, must be made safe before any work is started. CNCG lines require steam purging and, if necessary, purging with nitrogen. When the incineration of DNCGs is stopped, the gas ducts are recommended to be flushed with air. If shut-



down or maintenance works are to be done in NCG systems, the relevant parts and equipment of the system must first be separated from the rest of the system and the separated equipment then effectively ventilated at the very beginning of the shut-down. Ventilation streams shall always be discharged into the atmosphere. NCG lines require also LEL and sulphur-containing gas measurement before starting work.

Flushing of CNCG lines is recommended to be done 24 hours before concentration measurement and the start of service/maintenance work on CNCG lines. Flushing of DNCG ducts is recommended to be done eight hours before concentration measurement and the start of service/maintenance work on DNCG ducts.

Before starting work, a risk assessment shall be made in which especially all pipelines coming to the working area are mapped. A written work permit practice before starting work is recommended for example working inside NCG devices always needs written work permits. ~~for tanks.~~

9.4.2 Instructions for maintenance work during ~~outages~~**shut-downs**

Welding generates sparks inside the NCG system and therefore involves an explosion risk. Maintenance work involving the NCG system always requires a special permit. In connection with a outage, the maintenance staff shall be told which lines are NCG lines and how repair and maintenance work shall be carried out on these lines.

~~9.5 — Actions in connection with disturbances~~

~~If the boiler protection is tripped, the incineration of methanol, turpentine, dissolving and mixing tank vent gases and concentrated and DNCGs in the recovery boiler shall be stopped and the gas lines ventilated/purged with steam.~~

~~If the emergency shut-down has been activated, the incineration of methanol, turpentine, dissolving and mixing tank vent gases and concentrated and DNCGs in the recovery boiler and import to the recovery boiler building shall be stopped. In that event, the backup burner on the roof of the recovery boiler plant cannot be used.~~



10 SPECIAL INSTRUCTIONS FOR THE DESIGN OF NCG LINES

These instructions have been prepared for designers of non-condensable gas lines. Because NCG lines are subjected to explosive, dissolving and corrosive compounds, design practices and ~~construction~~ materials shall be selected to meet specific design criteria for NCG pipelines/ducts, in addition to the criteria for normal ducts.

Non-condensable gas ducts shall be resistant to mechanical, chemical and thermal impacts caused by their contents and ambient conditions (snow, ice, vibrations, hot and cold surfaces). Pipeline connection methods shall be adapted to the material used.

10.1 Classification

Non-condensable gas lines are subject to the stipulations of relevant chemical legislation. These lines are divided into different classes, depending on their toxicity.

The division of NCGs into DNCGs, vents and CNCGs is not sufficient to characterise the hazardousness of the contents. Gas mixtures must be classified in accordance with regulation 807 (of the 26 of September, 2001) of the Finnish Ministry of Social Affairs and Health concerning classification criteria and labelling of chemicals. On this basis, each NCG line must be classified in accordance with the specific contents of the line in question.

If no other specifications are given, CNCG~~concentrated non-condensable gas~~, methanol and turpentine lines can be designed preliminarily in accordance with the classification "toxic and inflammable".

If no other specifications are given, a DNCG~~dilute non-condensable gas~~ line can be designed preliminarily in accordance with the classification "hazardous".

The classification must be confirmed, however.



10.2 Signs

Non-condensable gas incineration equipment and pipelines shall be provided with appropriate markings. Markings shall always include the texts “Concentrated non-condensable gases”, “Turpentine”, “Methanol” and “Dilute non-condensable gases”, and whenever necessary, “Inflammable”, “Toxic” and “Must not be welded or cut without supervision”, and “Equipment must not be dismantled without permission in writing”. In addition, all pipelines/ducts shall be provided with markings showing the direction of flow.

Smoking and the use of open fire are prohibited in areas where NCGs are handled. Appropriate warning signs must be set up.

10.3 Condensate ~~drain~~removal

In the design of condensate ~~drain~~removal systems, special attention shall be paid for example to the size, amount and location of ~~drain~~outlets. See Chapters: 5.7, 6.7 and 7.4.

10.4 Duct/pipeline material

Recommended material for DNCG ducts is corrosion-resisting steel EN 1.4301 (SS2333) or equivalent.

Pipe PN \geq 10 shall be used for CNCGs. Recommended material for pipelines is acid-proof steel 1.4436 (SS2343) or equivalent.

Recommended material for pipelines for incineration of methanol/turpentine is EN 1.4436 (SS2343) or equivalent.

Some NCG compounds, such as turpentine and methanol, are strong solvents, so the use of plastic or glass fibre parts is prohibited.

10.5 Duct ~~pipeline~~ insulation

Non-condensable gases contain large amounts of water and other condensing substances. Condensation and the handling of condensates shall be minimised. NCG lines shall be insulated in places where their surface temperature is high.



~~Water-d~~Drains shall be insulated and outdoor ~~water~~-drains shall, if necessary, be provided with ~~heat electrical~~-tracing.

10.6 Pipe/duct slopes

Almost without exception, NCGs contain significant amounts of water vapour. Even if ducts are carefully insulated and/or provided with ~~heatelectrical~~ tracing, there is always a possibility of condensation. The slope instructions and locations of condensate ~~drainingremoval~~ connections are described in more detail in the following chapters: 5.7, 6.7 and 7.4.

10.7 Valves/dampers

Valves/dampers for NCG incineration systems shall be selected based on the characteristics of the gas.

EN 1.4301 (SS2333) or corresponding is the recommended ~~construction~~ material for DNCG valves/dampers.

Stainless steel EN 1.4436 (SS2343) or corresponding is the recommended construction material for valves for CNCGs.

Stainless steel EN 1.4436 (SS2343) or corresponding is the recommended construction material for valves of methanol/turpentine incineration systems. The valve sealing material must be resistant to dissolution (methanol and turpentine are solvents).

10.8 Flanges, connections

Special attention shall be paid to effective sealing of flanges and connections under any conditions. When replacing seals, special care shall be taken to select the right kind of sealing material. ~~NCGs contain dissolving and corroding compounds.~~

Connections shall retain their good sealing properties also in use (oxidation, reactions with sulphur compounds contained in NCGs, freezing, bending, etc.).

10.9 Fans

Fans are recommended for transferring DNCGs and vent gases.



10.10 Ejectors/liquid ring compressor

A fan must not be used for transferring CNCGs. Instead, a steam ejector or a liquid ring compressor shall be used. The maximum allowed steam temperature of the steam ejector is 200 °C (corresponding to a steam pressure of approximately 12 bar~~(yp)~~) because of the auto-ignition temperature of the gas components. In connection with a disturbance or start-up situation, the CNCGs may contain so much oxygen that the gas mixture is ignitable.

10.11 Duct/~~pipe~~-sizing

When incinerating CNCGs, operating gas velocities should exceed the flame propagation velocity under all circumstances.

The operating pressure of CNCG ~~pipelines~~ is -10...+10 kPa and that of DNCG ~~ducts~~ -5...+10 kPa.

~~DNCG Dilute non-condensable gas ducts~~ must withstand underpressure up to 10 kPa. ~~CNCG Concentrated non-condensable gas pipelines~~ must withstand full underpressure (100 kPa).

CNCGs are typically classified as extremely toxic, ~~toxic~~ or inflammable. The pipelines for such substances shall be sized in accordance with the operating pressure, with a minimum design pressure of 4 bar (g). A pressure of 1.3 times the design pressure is enough for sizing valves and other pipeline auxiliary equipment. The pressure of the purging steam shall be taken into account in selecting the design pressure.

DNCGs are typically classified as harmful. The design pressure of a pipeline for harmful NCGs shall be at least 1.3 times the operating pressure, with a minimum of +10 kPa (g).

10.12 Location of flame arrester

The flame arrester should be installed as close to the actual ~~burner/incinerator~~ as possible. If the bottom of the flame arrester is below the bottom level of the ~~pipeline~~, the flame arrester must be provided with a condensate drain.



10.13 Location of rupture discs

Rupture discs shall be installed in places where they will not represent a hazard to people working on normal operating floors. Their condition shall be monitored in the automation system by activating an alarm.

Rupture discs should be installed as a direct continuation of the duct section in question, without bends. The duct continues by means of a t-branch.

10.14 Location of bypass

A bypass duct/pipe outlet shall be installed as high as possible. The bypass is recommended to be taken as a separate pipeline to the ~~chimney~~ stack or the roof of the recovery boiler plant.

~~The bypass stack should be situated as close to the boiler as possible, in order to minimise the length of the pipeline between the bypass stack and the recovery boiler. For practical reasons, the bypass stack is typically situated close to the recovery boiler because the ejector or fan is often in the recovery boiler building or its vicinity, and the stack is always on the pressure side, after the ejector or fan.~~

The automatic valves for bypass duct must have fail-open actuators i.e. open automatically, if the supply of operating energy is turned off. This will confirm that gases will be led always to safe place even in case there is no instrument air available. Valves must be provided with limit switches indicating position to the automation system.

10.15 Hazardous area classification ~~Place classifications~~

The classification of different places where NCG equipment is located is based on the instructions and stipulations in the legislation on chemicals.

For CNCGs and turpentine-containing ~~fouled~~ contaminated condensates, the Hazardous area classification ~~place classification~~ often has to be based on the classification of places involving a risk of explosion. Turpentine is lighter than water and poorly soluble, so in case leakage it separates to water surface and evaporates to the air.



~~Place classes~~Hazardous area classification and impact areas are also defined for possible leakage points. Equipment that will be situated in the impact area of a specific place class will be designed from components in conformity with the classification. Zone 1 means that explosive gas will occur sporadically and Zone 2 that gas is not expected to occur in the area in question during normal operation.

If a place is considered risk-prone, it can be equipped with automatic high efficiency ventilation in case that the H₂S concentration rises too much.

The boiler ~~house room~~ and the possible area in the immediate vicinity of the CNCG, methanol and turpentine burner shall be equipped with a measuring device that will activate an alarm if the hydrogen sulphide (H₂S) or VOC~~sulphur~~ concentration becomes too high, ~~if a NCG line passes through a typically manned area.~~

10.16 Manufacturing permits

A manufacturer making pipelines whose operating pressure exceeds 0.5 bar (g) and which conduct inflammable gases (usually CNCGs) must have a manufacturing permit as stipulated in the pressure vessel legislation, and also an officially appointed manufacturing supervisor.

If dilute non-condensable gases, because of their contents, are classified as harmful substances, the pipeline manufacturer does not need a manufacturing permit or manufacturing supervisor as stipulated in the pressure vessel legislation.

Installation works related to methanol and turpentine lines shall be carried out in accordance with relevant regulations (gas installations, oil installations, and pressure vessel manufacture). In practice, it is advisable to handle the installation in the same way as installation of oil and gas pipelines for recovery boiler start-up burners.

In all works related to manufacture and installation, it is recommended to use companies with relevant documentation proving their competence.

Installation works for the burner and support fuel pipelines can only be carried out by a company~~firm~~ holding the relevant installation permits.



11 OTHER INSTRUCTIONS AND REGULATIONS AFFECTING DESIGN AND OPERATION

11.1 Finnish law

The most important Finnish laws and regulations concerning NCGs are:

- Chemicals Act (744/1989)
- Decree on the Industrial Handling and Storing of Dangerous Chemicals (59/1999)
- Decree of the Ministry of Social Affairs and Health on Chemicals referred to in Annex VI to the CLP regulation (5/2010)

11.1.1 Hazard and operability study (HAZOP)

The Chemicals Act defines hazardous gases and requires a hazard and operability study (HAZOP study) to be carried out for NCGs.

11.1.2 Classification of pipelines

The Chemicals Act and the Decree on the Industrial Handling and Storing of Dangerous Chemicals specify technical equipment requirements and stipulate, in practice, for example how pipes are to be classified, how regular inspections are to be carried out and that CNGs are to be transferred with an ejector and not with a fan.

At present, there are two parallel legislations in force concerning the classification, labelling and packaging of chemicals. The classification and labelling regulations according to the so-called Substance Directive (67/548/EEC) and Mixture Directive (1999/45/EC) are still in force and they have been enforced in the Chemicals Act (744/1989), Chemicals Decree (675/1993), as well as in the Ministry of Social Affairs and Health Decree on Chemical Classification Principles and Labelling (807/2001) (the so-called outgoing system).

EU's new Regulation on Classification, Labelling and Packaging of Substances and Mixtures, the so-called CLP Regulation (EU) no. 1272/2008 entered into force on January 20, 2009. This regulation is as such in force in all EU member states. The outgoing legislation is in force within certain transition periods.



The Ministry of Social Affairs and Health Decree on Chemical Classification Principles and Labelling (September 26, 2001/807) provides a formula for determining inflammability, using standard ISO 10156, Determination of fire potential and oxidizing ability for the selection of cylinder valve outlets, and its Tables 1 and 2. DNCGs are classified as harmful gases but not as explosive gases.

Systems for separate incineration of CNCGs are, in practice, constructed in accordance with the requirements for natural gas equipment. There are detailed instructions for natural gas, which specify accurate requirements for example for valves and flushing systems.

11.1.3 Classification of potentially explosive atmospheres (ATEX)

The EU ATEX Directive (94/9/EU) refers to potentially explosive atmospheres. In addition, instructions concerning potentially explosive atmosphere classification are given in standard EN 60079-10: 1996, which has been translated into Finnish and published as SFS Manual no. 59 (4th edition 1998). The classification of potentially explosive atmospheres is applied to CNCGs. The zones and their impact areas are defined for possible leakage points. Equipment to be situated in the impact area of the zone class in question is designed with components in conformity with zone class. Zone 1 refers to a place in which an explosive atmosphere is likely to occur in normal operation occasionally and Zone 2 to a place in which an explosive atmosphere is not likely to occur in normal operation.

11.2 European standards

The European standard on pressure vessels (EN 12952-8, Annex A.3.3.) imposes certain restrictions on the incineration of NCGs in a recovery boiler. CNCGs can be incinerated in a recovery boiler, provided that the steam generation is at least 50% of the boiler's nominal load with black liquor.

In addition, the standard requires precautions to be taken in order to prevent aqueous solutions from penetrating into the boiler furnace together with gases or fuel, and to prevent CNCGs from escaping into the boiler ~~house~~room's indoor atmosphere. The above-mentioned clause does not apply to DNCGs.



All NCG lines must be equipped with a automatic shut-off valves.

11.3 Existing instructions

Sweden's Sodahuskommittén and the American BLRBAC have issued instructions for the handling and incineration on non-condensable gases. BLRBAC's instructions are very detailed and include accurate technical descriptions of design solutions.

11.3.1 Sodahuskommittén

Our Swedish sister organisation, Sodahuskommittén, published in December 2008 a decision in principle that their recommendation from then on will be published only in Swedish and they are intended for the sole use of the members of the committee.

11.3.2 BLRBAC

BLRBAC has issued instructions for minimising risk situations, entitled "Recommended good practice for the thermal oxidation of waste streams in a black liquor recovery boiler", updated in February October 20120.

The document lists inflammable NCGs and defines their explosion limits. It explains where NCGs are collected and how they can be destroyed, giving instructions on how to incinerate methanol and turpentine, among others. A whole chapter is dedicated to the handling of chip silo vents. The document defines the conditions at which black liquor solids contents NCGs can be incinerated safely.

In a problem situation, and especially in practice when the Emergency Shutdown Procedure (ESP) is activated, NCGs shall be instantly diverted from the recovery boiler in the same way as other fuels.

BLRBAC's document also warns of the toxicity of NCGs and the risk of corrosion that they involve, in spite of the fact that there is no proven evidence of corrosion of pressure vessels caused by NCGs.



The document is very comprehensive, including also conditions for interlock~~control logic functions~~ and examples of the PI and interlocking diagrams of incineration and collection systems.



12 DEVIATIONS FROM PRACTICES DESCRIBED IN OTHER INSTRUCTIONS

The Finnish Recovery Boiler Committee has endeavoured to comply with other countries' recommendations and instructions concerning incineration of NCGs in the recovery boiler. At least the following factors deviate from those presented in other recommendations.

12.1 Deviations compared to BLRBAC's instructions

The Finnish Recovery Boiler Committee's instructions differ from those of BLRBAC (<http://www.blrbac.org/>) for example in the following respects:

- Chapter dedicated to the handling and incineration of chip silo vents
- Chapter dedicated to the handling and incineration of stripper off gases (SOG)
- Examples of the PI and interlocking diagrams of incineration and collection systems
- The recommendation for relative humidity of DNCGs and vents after condensers 50%, corresponding to a temperature of < 43.3 °C (110 °F). Heating at least 22.2 °C (40 °F) above condensation temperature.
- The incineration of DNCGs can be started when the steam flow greater than 30% of the steam flow at MCR.
- The incineration of CNCGs can be started when the steam flow greater than 50% of the steam flow at MCR
- Incineration of methanol/turpentine/soap/tall oil mixed with black liquor.



13 EXAMPLES OF ACCIDENTS

In the following, a number of examples of damage and risk situations known to the Finnish Recovery Boiler Committee are described. The damages are described to give the reader of this recommendation a concrete picture of possible danger situations related to the collection or incineration of NCGs.

Similar events have also been recorded in other mills.

13.1 Example 1:

Causes:

~~In connection with the start-up of the pulp mill~~, the chip feeding system to the digester stopped due to problems with the black liquor filters during mill start-up. Thereafter the chip binsile level was raised to 95% and the chips were presteamed with fresh steam, generating a surface temperature of 90 °C in the chip silo.

Half an hour after the shut-down, chip feeding to the digester was restarted and flash steam was led to the chip silo, because the CNCG system could not keep sufficient underpressure in the collection line to the CNCG burner. The presteamed chip bed was unable to condensate and to absorb the steam which therefore passed through the chip bed, quickly increasing the temperature of the upper part of the chip silo, and further to the DNCG collection system.

The temperature interlock after the chip silo vent condensers reached the interlock limit of > 55 °C and the gases were diverted to the bypass pipeline. Before this safety interlocking, gases got into the collection system of DNCGs for approximately 40 seconds, causing an explosion about 1.5 minutes after the flash steam valve was opened. The explosion occurred close to the NCG boiler burner in the collection pipeline.

Measures taken after the explosion

Before starting up the mill again, the following measures were taken:



- The opening speed of the flash steam valve was slowed down (5%/min in range 0-10%), so that the amount of flash steam is controlled at start-up.
- The chip conveyor to the silo must be in operation at least five minutes before the flash steam valve to the chip silo can be opened, to control that there are fresh chips in the silo to condensate the flash steam.
- When the chip silo temperature rises above 90 °C, the cooling water valve of the condenser opens immediately 100%.

13.2 Example 2:

An explosion in the mill concrete stack damaged the DNCG steel stack across its whole length, a part of the combustion gas stack of the lime kiln and a part of the DNCG pipeline. The explosion was the result of several interacting factors. The operation conditions preceding the explosion increased the concentration of the gas mix in the bypass stack to ignition and explosion concentration. The prevailing weather conditions forced the hot combustion gases from the lime kiln into the outgoing DNCGs causing ignition.

Causes:

- DNCGs of the evaporation plant had reached explosion concentration during normal operation and shut-down.
- Gases from the ~~firing~~ combustion liquor tank were led to the collection system of DNCGs. There were heavy gas ~~formation~~ leaks from the ~~firing~~ combustion liquor tank 22 hours before the explosion.
- ~~Fibre line~~ The DNCG fan ~~in the fibre lines~~ was stopped and therefore there was no dilutive effect from the DNCGs in the fibre lines.
- Because of the prevailing weather conditions the hot combustion gases (approximately 260 °C) from the lime kiln were directly mixed with the outgoing DNCGs which were above the lower explosion limit. The bypass stack of the DNCGs and lime kiln combustion gas stack were at the same level.



Measures taken after the explosion

- The fan of the DNCGs in the fibre lines shall be kept in operation during shut-downs while simultaneously conducting dilution air to the suction side of the fan to ensure a low enough concentration.
- The gases from the ~~firing~~combustion liquor tank shall be conducted to the collection system of CNCGs or ~~to the group of~~ evaporators, not to the collection system for DNCGs.
- The temperature and solid content of the ~~strong~~ liquor in the ~~firing~~combustion liquor tank shall be decreased to 117 °C respectively 67-69% for the shut-down to prevent gas formation reactions.
- The circulation pump of the DNCG scrubber shall be kept in operation during ~~outage~~shut-downs.
- Clear instructions shall be given for the operation of DNCGs during ~~outage~~shut-downs.
- Circulation water ~~from~~ evaporation NCG scrubber shall not be pumped from to the ~~spill~~leakage liquor tank, from where the gases are sucked back into the DNCG collection system.

Other recommendations

- The concentrations of methanol, TRS compounds and turpentine in DNCGs, liquors and condensates shall be identified.
- A concentration measurement or flow measurement shall be installed in the DNCG collection system for monitoring.
- A simulation programme shall be made to specify in a reliable way the amount of dilution air in different operation situations.
- The capacity of the fan for the closed collection of DNCGs in the evaporation plant shall be increased and the suction side equipped with an intake for dilution air.
- The lower part of the bypass pipe for DNCGs shall be redesigned.
- The use of the fan shall be stopped and the gases conducted to their own pipe on the suction side, if DNCGs reach explosion concentration.



- A hazard and operability study (HAZOP) shall be made of DNCGs concentration under normal operation and in shut-down situations.

13.3 Example 3:

~~In connection with the start up of the pulp mill and recovery boiler a~~ fire was started in the DNCG duct damaging the front wall of the recovery boiler and approximately 2 m of the duct during boiler start-up. The fire was possibly started by boiler backfire igniting the deposit accumulated at the bottom of the DNCG duct. The deposit formed at the bottom of the DNCG duct normally originates from the tall oil plant.

The Durability Sub-Committee of the Finnish Recovery Boiler Committee pointed out the following matters that should be taken into account after the accident:

- The functioning of the scrubbers shall be improved to ensure a good scrubbing result. The liquid distribution, drop separation and temperature after the scrubber shall be checked for each scrubber.
- The control of the dilution of the DNCG system shall be checked.
- There should be a droplet separator ahead of the preheater.
- The composition of NCG shall be checked regularly according to incineration performance.
- Ducts shall be regularly checked and cleaned.
- A hazard and operability study (HAZOP) shall always be made when NCG system is modified ~~in connection with a modification of the system~~ together with the supplier of the NCG system.

Measures taken after the explosion:

- The DNCG fan was renewed.
- The preheating of air was intensified by adding a heat exchanger for fresh air, thus raising the temperature of the DNCGs.
- The capacity of the collection system was increased.
- Interlockings relating to incineration were checked.
- Dissolving tank vent gases were included in incineration together with DNCGs.



13.4 Example 4:

Causes:

There have been two explosions after the start-up of mill. The first happened in the chip ~~binsile~~ and the second in a pressurised ~~firingcombustion~~ liquor tank.

When flash steam was conducted to the chip silo during a shut-down, CNCGs were discharged into the collection system for DNCGs through the chip bed or concentrated methanol in the chip ~~beds~~ was released. There was no condensatione for the flash steam so it had to be led to the chip silo. The explosion took place in the recovery boiler's DNCG duct. The amount of recovery boiler secondary air had been manually reduced.

The second accident occurred in the pressurised ~~firingcombustion~~ liquor tank, which exploded when its circulation pump was left functioning during shut-down and a heavy formation of NCG compounds started inside the tank. There was no relief valve in the tank and it was not connected to the collection system for CNCGs.

Measures taken after the explosion

As a result of the explosion of CNCGs ~~from the chip silo~~ in the chip silo's DNCG collection system, the mill invested in a reboiler where the chips are presteamed with clean steam. This improvement in the cooking plant eliminated the problems related to the collection and handling system for DNCGs. The use of reboiler steam in the chip silo keeps the concentration of DNCGs at a permanently low level. The acquisition of a backup incineration system was also considered because of odour problems, but the new reboiler solved the worst problem of the odorous gas source and the investment was abandoned.

The ~~firingcombustion~~ liquor tanks were connected to the CNCGrecollection system ~~for NCG~~ and a relief valve was installed in the tanks.

13.5 Example 5:

Causes:



During start-up of the mill Turpentine and NCG compounds with a concentration within the explosion limit escaped ~~at the start-up of the mill~~ from the chip silo and started to burn explosively in the flare. The fire front advanced backwards to the DNCG fan, where a second explosion occurred.

The explosion was the result of several interrelated factors in the chip silo, where the DNCGs safety system was unable to control the exceptional conditions. The safety system functioned well under normal operation conditions, but was insufficient under exceptional conditions with partially unknown effects. As to the other tanks within the collection system, the changes in the DNCGs are slower and the mill's safety systems have been enough.

The separate circumstances leading to the explosion are as follows:

- Long chip presteaming time in the chip silo (low-pressure feeder stuck)
- ~~Higher than normal~~ concentrations of TRS and VOC in the chip silo and its gases at start-up
- Delay in the temperature measurement in the vent line from the chip silo
- Delay in the LEL analyser
- The NCG scrubber primarily condensates water vapour, resulting in an increased amount of NCG in the air volume. The LEL analyser was positioned ahead of the condensing process, resulting in a lower measured concentration.
- Operating failure of the dilution valve located on the suction side of the collection fan for DNCGs.
- Automatic reopening of the collection valve for digester DNCGs ~~from the digestion plant~~ based on the measurement of the LEL analyser.
- Orifice plate~~Throttle disk~~ in the vent pipe of the DNCGs of the digestion plant.
- ~~Going out of the F~~flare flame out.

The amount of dilution air has been sufficient under normal operating conditions. The results are based on the measurements carried out in 1998 and 2000. However, the measurements do not cover exceptional conditions.



The amount of dilution air is increased at three points in the DNCG collection and handling system:

- The underpressure of the DNCG fan via the chip silo and vent pipe of other tanks sucks dilution air at different collection points. The system, however, does not function if the system is pressurised and gas leaks out of it. In the case of the chip silo, a orifice plate has been installed in the dilution air outlet.
- The LEL non-condensable gas analyser opens the dilution valve on the suction side of the NCG gas fan, if the concentration rises over 30% above the LEL limit.
- DNCGs are led to the flare, where the necessary amount of additional air dilutes the non-condensable gas.

Measures taken after the explosion:

- The operation mode of the chip silo and its control are developed.
- ~~In connection with the m~~Modifications the NCG handling system and the related security systems are examined as a whole.
- A sufficient supply of dilution air under exceptional conditions is ensured.
- A ~~security-safety~~ system is built up according to safety automation principles.
- The follow-up of DNCGs concentrations is improved and interlocking limits checked (development and location and possible duplication of the analyser).
- Exceptional situations are recognised and instructions for such cases are worked out.
- Dangers are systematically identified and risks assessed.
- ~~Supervision is~~~~Controls are~~ harmonised.

13.6 Example 6:

Causes:

There were disturbances in the chip silo due to stopping of the chip conveyor and leakage of flash steam into the silo vent gas system. The vent gases of the chip silo were connected to the DNCG collection system through the white liquor scrubber after the start-up of the mill. The connection was made because the chip silo was the primary individual odour nuisance at the mill. During the disturbance, the



amount and concentration of chip silo vents strongly increased and combustible gas containing turpentine, methanol and TRS compounds got into the recovery boiler tertiary air system.

Non-condensable gas measurements after the explosion showed that exhaust gas from the chip silo had exceeded the lower explosion limit concentration, and the gas could have ignited by backfire from the recovery boiler or by a spark from the fan. ~~The propagation velocity of a turpentine flame is 150 m/s, the velocity in tertiary air nozzles being approximately 85 m/s.~~

Luckily, the explosion did not cause any personal injuries.

Measures taken after the explosion

In the investigation report it was recommended that the exhaust gases from the chip silo shouldn't be taken to the DNCG collection system. The report, however, pointed out that if the vents are taken there for odour reasons, the vent line of the chip silo has to be equipped with turpentine condenser and scrubber where the gases are cooled down with cold water. Moreover, measurement of the lower explosion limit and bypass in danger situations were recommended.

13.7 Example 7:

The non-condensable gas system may involve an explosion risk if the gas concentration is within the explosion range. The gas concentration has either to be kept high enough or low enough. Especially shut-down and start-up situations are problematic.

Generally, explosions in recovery boilers caused by NCGs have been caused by DNCG systems.

The concentration of the gas mixture in the DNCG incineration system had risen too high during the start-up phase following a shut-down of the pulp mill.

The gas mixture, whose concentration probably was within the explosion range, had escaped via a partly open shut-off valve and/or a water drain pipeline to the DNCG nozzles at the back wall of the recovery boiler's tertiary level during the



start-up phase following a shut-down of the pulp mill. The mixture was ignited at the nozzles and the flame then propagated along the DNCG ducts to the front part of a heat exchanger, where an explosion occurred.

At present, mill keeps the NCG collection system running also during ~~outages~~
~~shutdowns~~.

13.8 Example 8:

If water condenses in a DNCG line, there is a risk of smelt-water explosion, if the water gets into the recovery boiler.

Special attention should be paid to water drains which shall cover all lines comprehensively. The bypass stack should be situated as close to the boiler as possible in order to minimise the length of the line between the bypass stack and the recovery boiler building. For practical reasons, the bypass stack is situated close to the recovery boiler, because the ejector is often installed inside the recovery boiler building or in its vicinity, and the stack is always on the pressure side, after the ejector.

A serious explosion occurred at the mill because a relatively large amount of water entered the furnace via the DNCG gas burner. As a result of the explosion, four persons died and 13 were injured. Another smaller smelt-water explosion occurred 15 minutes later.

The ultimate reason for the explosions was probably the water that entered the furnace, putting out the fire on the char bed. The water apparently entered the recovery boiler through the DNCG burner, when the automation system switched off the combustion system, purging the gas line with steam, while at the same time flushing the condensed water in the line into the furnace.

13.9 Example 9:

The risk of poisoning by NCGs needs to be pointed out to all those working with or close to NCG lines.



In the 1980s, there was an accident where two members of the recovery boiler's operating crew were poisoned by non-condensable gases. The accident occurred when black liquor stopped flowing into the black liquor mixing tank as a result of plugging of the vent pipe. The mixing tank's level indicator was a pressure transmitter, and because of the increased pressure the transmitter's level indication was wrong. In a normal situation, residual liquor was conducted to the bottom of the mixing tank, about 2 m below the liquor surface, thus preventing hydrogen sulphide from spreading into the room.

In the accident situation, the liquor level dropped too much, and the residual liquor then caused a reaction releasing hydrogen sulphide. The gas pressure generated by the reaction closed the incoming black liquor valve, causing the formation of hydrogen sulphide to increase strongly. An operating crew member who was cleaning the vent gas line inhaled hydrogen sulphide coming from the opening. The person in question felt dizzy and went outdoors, after which he temporarily lost consciousness. Another member of the operating crew went to inspect the operation of the mixing tank, noting that black liquor-containing foam was streaming out from the inspection hatch of the mixing tank. He too felt dizzy, and went to the control room, where he informed the control room operator about the situation and then temporarily lost consciousness. The persons concerned had a check-up at the hospital and did not suffer any permanent injury. After this accident, the pumping of extra residual liquor to the mixing tank was stopped to eliminate the risk factors involved.



14 CLOSING WORDS

The members of the Finnish Recovery Boiler Committee have called for a clear stand on the collection and incineration of NCGs in chemical pulp mills. The reason for this is the general concern over safety and the functionality of the overall process.

Depending on their ignition or explosion range, NCGs are divided into dilute gases, whose concentration is below the explosion range, and concentrated gases, whose concentration is above the explosion range.

The incineration of CNCGs in the recovery boiler started attracting strong interest during the 1990s. DNCGs have been incinerated in the recovery boiler before.

There have been numerous problems with the incineration of dilute and CNCGs. Incineration involves certain risks, in the same way as incinerating other combustion gases, which is why a recommendation is needed. ~~Sweden and North America already have recommendations, though incineration of CNCGs is much less common in these countries than in Finland.~~ There are also different schools of thought, which has caused confusion and a lively debate on this subject.

There are numerous instructions for recovery boiler operation. The incineration of NCGs requires a recommendation of its own to confirm safe incineration of NCGs in the recovery boiler. The relevant legislation does not deal with NCGs to any major extent; it is primarily governed by the general principles of the Chemicals Act. The purpose of this recommendation is to allow safe incineration of NCGs in the recovery boiler.



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APPENDIX I

Example of how to calculate furnace~~base~~ load

EXAMPLE OF HOW TO CALCULATE FURNACEBASE LOAD

To confirm a standard procedure for calculating the minimum load for incineration, two ways to calculate the minimum load are presented in the following. In the first, the load is tied to the steam flow, in the second to the liquor flow. The unit for furnace heat loading is the net heat to steam generation divided by the floor~~bottom~~ area of the furnace.

	100	%	43	%
<u>Boiler floor</u> a Area	100	m ²	100	m ²
Steam generation	80	kg/s	34.3	kg/s
<u>HP</u> Fresh steam pressure	80	bar	80	bar
<u>HP</u> Fresh steam temperature	480	°C	480	°C
<u>HP</u> Fresh steam enthalpy	3349.5	kJ/kg	3349.5	kJ/kg
Feedwater pressure	100	bar	100	bar
Feedwater temperature	115	°C	115	°C
Feedwater enthalpy	489.6	kJ/kg	489.6	kJ/kg
<u>Steam drum</u> Cylinder pressure	90	bar	90	bar
Sootblowing steam enthalpy	2942.9	kJ/kg	2942.9	kJ/kg
Blow-out enthalpy	1363.7	kJ/kg	1363.7	kJ/kg
Blow-out	1.5	%	1.5	%
Sootblowing	2.0	%	2.0	%
Heat to steam	2.922	MJ/kg	2.922	MJ/kg
	233.8	MW	100.2	MW
Net heat per <u>floor</u> bottom area (<u>furnace heat</u> loading)	2.34	MW/m ²	1.00	MW/m ²

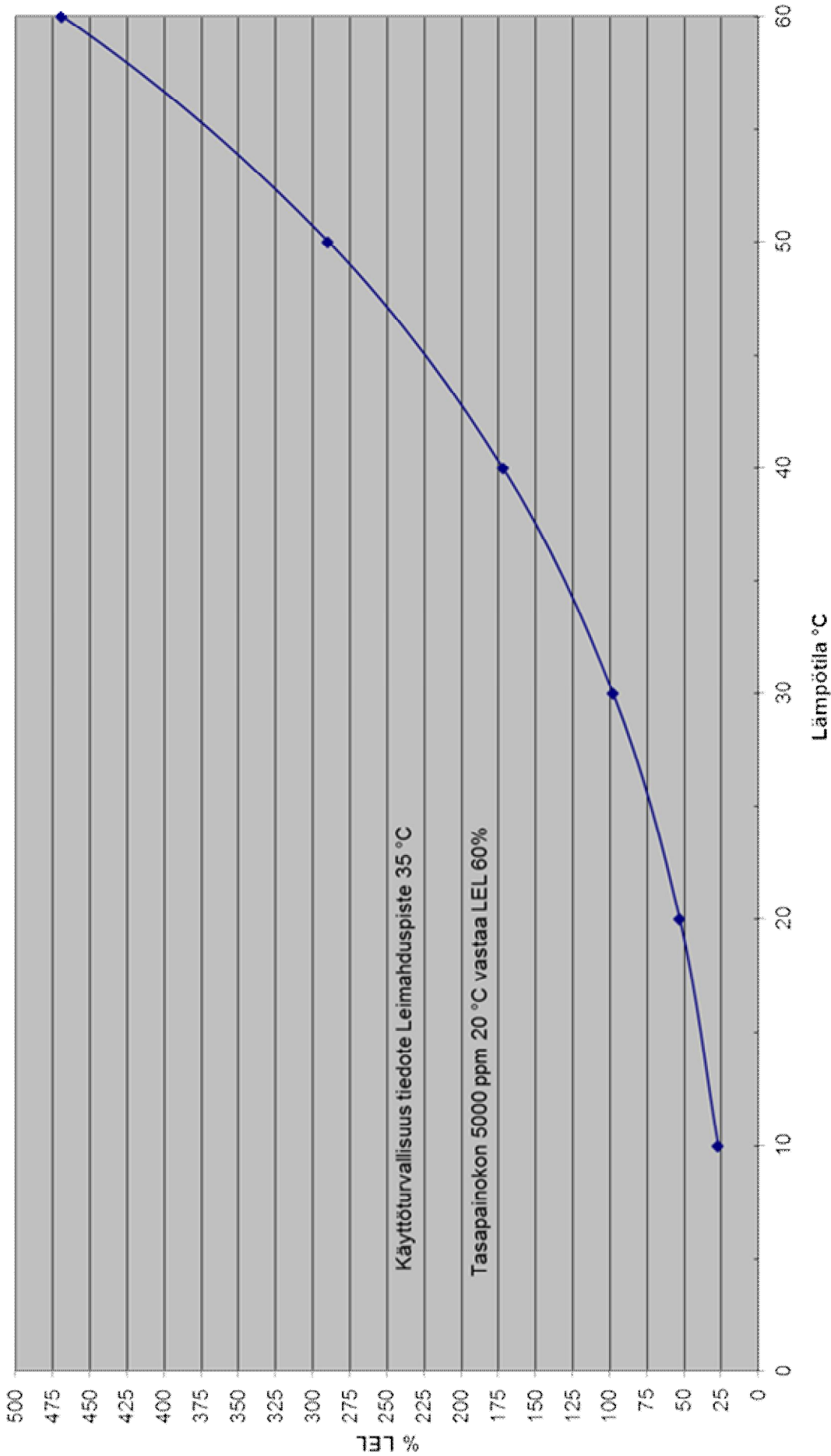
or

	100	%	43	%
Black liquor flow	2200	tka/24h	943	tka/24h
	25.5	kgds/s	10.9	kgds/s
Net heat (from balance)	9200	kJ/kgds	9200	kJ/kgds
	234.3	MW	100.4	MW
Net heat per <u>floor</u> bottom area (<u>furnace heat</u> loading)	2.34	MW/m ²	1.00	MW/m ²

APPENDIX II
Alfa pinene
Turpentine balance curve



LEL %LEL Tärpätintasapainoilmassa



APPENDIX III
Methanol balance curve

