# Syngas Cleaning by Wet Scrubber

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*Abstract*: This work deals with syngas wet scrubbing and tar properties. Wet scrubbers are a technologies that helps clean the pollutants in the gas so that the gas may be used in combustion engines. Gas has been generated from biomass in the atmospheric fluid gasifier and has high temperature (500-800 °C). Research focuses mostly on tar elimination and dust removal.

Key-words: biomass, gasification, gas cleaning, wet scrubber, tar reduction,

# **1** Introduction

Gasification is a very old technology, historic records about its use date back to first half of the 19th century. Mass use of this technology began during World War II when automobiles combusted wood gas. Renewed interest in the technology continued at the turn of 1980s and in the early 1990s. Current research in the area of gasification is fuelled by need to replace fossil fuels with renewable energy sources due to expected unavailability of the fossil fuels in the future and increased impact of greenhouse effect.

Gas generated by gasification contains several impurities which impede its direct use. There are several methods to modify actual gas quality so that it complies with requirements for final gas quality. Primary methods, which positively affect gasification process itself, and secondary methods, which clean the produced gas, are researched. Both of these methods have their pros and cons. Unless their principles, boundary conditions, particulars and restrictions of use are sufficiently described, the methods cannot be compared.

This paper deals with gas wet scrubbing which cleans the impurities in the gas so that the gas may be used in combustion engines. Gas is contaminated and has high temperature (500-800°C) as it leaves atmospheric fluid gasifier.

Research focuses mostly on tar elimination and dust removal.

Core of our work lies in research of tar elimination, which is a mixture of higher hydrocarbons and their compounds. Water and organic liquid may be both applied in tar elimination from gas. Boiling point (volatility), availability, and price of organic liquid are major criteria for selection of a proper material. In past, scrubbing oil was often applied for the purposes of extensive research in the area of coal gasification and production of town gas. Biodiesel and/or rapeseed oil methyl ester (RME) are alternatively widely used nowadays. Both liquids were assessed; however, since RME comes from renewable sources of energy and seems to be more environmentally friendly, most of the paper deals with RME technologies. Scrubbing of gas generated by biomass gasification (or a different organic material) has its own specifics. Gas contains very fine little wettable dust (ash, unburned residues, and soot), tar (mixture of higher hydrocarbons and their compounds) and water vapour. Tar condensates in a wide temperature range from ca. 180 to 300°C and in temperatures below zero. As the tar condensates, it creates adhesive liquid coating on cool surfaces which, together with dust, forms deposits that are very difficult to remove. If the temperature drops below condensation point, fine tar mist is formed; this mist cannot be removed from gas by common separators. Condensation temperature and tar in gas are bound by phase equilibrium.

Different mechanisms apply when water and organic liquid are used in gas scrubbing since the tar has different solubility. This concerns condensation and subsequent capture of aerosol droplets in water applications, and absorption and diffusion in organic liquid applications. Both these mechanisms are described in the paper, and comparison of tar concentrations for various temperatures of scrubbing liquid before after the gas scrubbing is given. and Temperature range for both liquids during experiments overlaps only partially (water: 0-40°C, RME: 30-85°C). Temperature range was selected so that both types of liquid may be later practically applied. In case of scrubbing using organic liquid, the process will be conducted below water condensation point so that amount of waste water polluted with PAH and its derivates is minimized.

In addition to studying efficiency of the scrubbing, we also analysed other properties of scrubbing liquids and important technology aspects. This includes particular transport and thermodynamic properties (heat capacity, viscosity, and vapour tension), disposal and regeneration of waste liquid, risks of liquid treatment, its availability, and price.

# 2 Gas impurities

Potential of gas generated by biomass gasification for subsequent production of electrical energy is restricted by difficulties related to purification of this gas. Gas impurities cause several operating problems to facilities. Corrosion, fouling of supply routes and tarring of engines and turbines work areas may lead to severe accidents. Impurities may be classified as:

- Dust (solid particulate matter)
- Alkali compounds
- Nitrogen compounds
- Tar
- Light hydrocarbons (methane, ethane)
- Heavy hydrocarbons

Chlorine, fluorine, and silicon may also be dangerous for certain pieces of equipment. Silicon is monitored in combustion engines especially as it has negative impact on quality of lubrication oil.

#### 2.1 Solid particulate matter

Solid particles in raw gas leaving the gasifier comprise inorganic fuel residues (ash), nonreacted biomass in the form of charred material (the so called coalite), and inert material of gasifier bed. Soot is also present in the dust. Gas with high concentrations of solid particles is typical of gasifiers with bubbling and circulating fluidized bed (turbulent conditions inside the reactor). Thermal treatment of biomass may form aerosol that has negative impact on human health and causes greenhouse effect.

Together with tar, it fouls pipe walls and causes abrasion as well as fouling of equipment modifying and using the gas.

Cyclone separators, barrier filters, electrostatic separators, and wet scrubbers are used and/or tested for elimination of solid particles.

# 2.2 Alkali compounds

Biomass is low in dust. Certain components of dust may cause several operation problems, and include alkali metals, i.e. potassium (K) and sodium (Na). These are present in stalk culm plants. Salts of potassium and sodium evaporate below 700 °C; they are easily meltable and form deposits of ash on cooler surfaces of equipment (below 650 °C). Deposits concern mostly heat exchangers, combustion engines and gas turbines positioned beyond generator because unless alkali compounds are removed from the gas, they pass through the whole system, and condensate and deposit beyond the filtering devices. Another issue is a hightemperature corrosion of metal materials, which may be caused by alkali salts, especially if catalyses vanadium the corrosion (cogasification of biomass and waste) [1].

Deposition of alkali salts is usually treated by cooling the gas and eliminating fine particles where salts condensate at temperatures below 600 °C. These technologies are highly efficient, e.g. electrostatic and bag filters, or wet scrubbers.

#### 2.3 Nitrogen compounds

Most of nitrogen in the produced gas is in the form of  $N_2$ , and its concentration depends on type of the gasifying medium. Nitrogen also forms compounds where ammonia (NH<sub>3</sub>) is the main nitrogen component in the raw gas from biomass gasification. NH<sub>3</sub> is created by conversion of protein and other nitrogen-containing biomass components. Fuel rich in protein (such as animal waste) are therefore a source of high amounts of ammonia. Cyanide (HCN) is a less significant compound.

Elimination of nitrogen compounds may be performed via standard catalytic methods for  $NO_x$  reduction. It is even better to use a catalyst for destruction of  $NH_3$  prior to combustion or wet scrubbing, provided gas temperatures are low enough. Nitrogen compounds may cause various operation problems in the scrubbers due to high solubility in water since their elimination from the water is very difficult (water is saturated with HCN and  $NH_3$  and is not capable of collecting them anymore).

#### 2.4 Sulphure

Considering the reducing environment in the gasification reactor, sulphure in the raw gas is in the form of  $H_2S$  (93-96%); carbon disulphide CS<sub>2</sub>, COS and SO<sub>x</sub> concentrations are minimum. These compounds are formed from sulphure in the fuel. In general, concentrations of these compounds in biomass are very low (hundredth and tenth of weight %) and thus H<sub>2</sub>S concentrations are commonly below limits of traceability. Yet, elimination of sulphure from the gas is highly desirable for majority of gas applications becuase it may various operation complications. cause Sulphure together with chlorine, fluorine, and alkali salts is corrosive for steel constructions. Sulphure is also potentially hazardous for certain types of catalysts, even in low concentrations (ppm). Technology of methanol production from syngas employs catalysts which may be deactivated (poisoned) by sulphure. Particular tar elimination catalysts are also sensitive to sulphure. Therefore it is important to thoroughly eliminate sulphure from the gas.

Wet scrubbing using additives (very costly), reactions of suitable sorbents or adsorption on metal catalysts (see chapter 6.7.3) are field-tested technologies, readily available.

### 2.5 Chlorine

Concentrations of chlorine in biofuel are rather low. Despite this fact, chlorine may cause formation of HCl and persistent organic compounds substances (PCDD, PCDF) which are created by reactions of carbon and chlorine, and catalysed by copper oxides. HCl is the most common form of chlorine in the gas.

Wet scrubber may eliminate certain chlorine compounds, especially HCl.

#### 2.6 Tar

Tar is a by-product of pyrolysis (main source is a volatile combustible). Definitions of tar differ and there had not been a unified scientific view on what constitutes tar, what its composition is, and how to analyse it. Energy Research Centre of the Netherlands (ECN) has researched tar for a long time and initiated a change in studying tar, its components, properties, tar collecting, and analyses. ECN published the so called Tar Protocol whose definition and methods are recognized both in Europe and in the world [2]. Tar designates a group of organic substances with various structures and chemical properties with boiling point higher than boiling point of benzene (80.1 °C). [3]

Concentrations of tar in gas from biomass gasification ranges from 1 to 15 g.m<sup>-3</sup> (in rare cases up to 75 g.m<sup>-3</sup>) depending on a type of gasifier, temperature in the gasifier bed during gasification, temperature and retention time of gas in freeboard, and fuel properties (granulometry, moisture, etc.). [4][5][6]

Tar in gas is in the form of vapors or persistent aerosols. Tar condensates in lower temperatures. Tar, condensed on cool surfaces, creates tar deposits and, together with solid particles, fouls pipe and other equipment. Initiation of tar condensation is related to concentrations of tar in the gas and its composition. If temperatures are below 400 °C, tar components may be dehydrated and start to create charred materials and coke. Not all tar components are dangerous and undesired, certain compounds may have positive impact on gas lower heating value and cause no operational difficulties. Despite this fact, tar is considered to be an underbelly of thermal treatment of biomass. Therefore, if the gas from biomass is to be further used, tar has to be removed.

# **3** Requirements on gas purity

Requirements on final equipment differ among various types of this equipment as well as among various manufacturers. Some of the requirements for operation of the equipment using syngas are not stipulated in internal standards of the manufacturers but are defined by researchers dealing with gasification. However, even these requirements often significantly differ. Differences gradually disappear and an unwritten, commonly accepted standard emerges.

Combustion engine manufacturers commonly define maximum allowed concentrations of sulphur (H<sub>2</sub>S), halogen (Cl), ammonia (NH<sub>3</sub>), silicon (Si), residual oil, solid particles and condensate in the gas. Manufacturers further stipulate requirements for gas inlet temperature (10-60 °C) and moisture content (max. 80 %), pressure and methane number and their changes in time. Required concentrations are related to a specific amount of energy, e.g. mg/MJ or mg/10kWh (related to gas in normal state), so that absolute amount of undesired substances entering the engine may be taken into consideration. Requirements for gas quality are rather stringent due to very low calorific value of the gas generated by gasification of biomass using air  $(3.5-6 \text{ MJ/m}_n^3)$ . Overview of requirements of several manufacturers of gas engines is given in Tab. 1.

Tab. 1 The requirements of manufacturers of internal combustion engines

	Caterpillar		Jenbacher		Deutz AG	
S compounds (H <sub>2</sub> S)	(mgH <sub>2</sub> S/ MJ)	57	(mgH <sub>2</sub> S/ 10 kWh)	2000	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	2340
halogenes (Cl)	(mgCl/ MJ)	19	(mgCl/ 10 kWh)	400	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	100
NH <sub>3</sub>	(mgNH <sub>3</sub> / MJ)	2.81	(mgNH <sub>3</sub> / 10 kWh)	55	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	30
Si	(mgSi/ MJ)	0.56	(mgSi/ 10 kWh)	0.2	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	10
oil	(mg/MJ)	1.19	(mg/ 10 kWh)	5	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	3250
particles	(mg/MJ)	0.8	(mg/ 10 kWh)	50	(mg/ m <sub>n</sub> <sup>3</sup> CH <sub>4</sub> )	10
particle size	(μm)	1	(μm)	3	(μm)	3–10

Manufacturers of engines combusting wood gas do not define any limit values for tar as there is no practical experience with this type of engines (or manufacturers have only negative experience). Unwritten, commonly accepted standard slowly emerges. This standard stipulates that tar concentrations should be decreased below 100 mg/m<sub>n</sub><sup>3</sup> in order to minimize problems related with tar; however, continuous operations of the equipment necessitate 50 mg/m<sub>n</sub><sup>3</sup> of tar.

# 4 Decrease in concentrations of tar from gas

There are generally two types of methods to decrease tar concentrations in gas: primary and secondary methods. Secondary methods further distinguish between dry and wet procedures.

Primary methods concern procedures performed inside the reactor. This is a desired practice because it has a potential to increase a total efficiency of energy conversion; concentrations of tar in produced gas are reduced and subsequent tar disposal is eliminated. In general, two procedures are applied - thermal destruction and catalytic destruction. More in [7].

Secondary methods for decrease in tar concentrations are applied outside of the reactor like are barrier filters, catalytic filter [8] and wet scrubbers.

Wet scrubbers droplets of separated matter and scrubbing liquid precipitate, and thus decrease tar concentrations. Formation of tar droplets has to be preceded by cooling of the gas to temperatures below 100 °C (usually from filtration temperature of ca. 250 °C - separates

solid particles). Exhaust temperature of gas is required to range from 35 to 60 °C. Scrubber must comprise a cooler, fine tar mist separator, and occasionally a solid particle separator, at the same time [9]. High concentrations of solid particles may have a negative impact on efficiency of the process of tar elimination. In addition to condensation and precipitation of particles, other physical phenomena may be used for tar elimination, e.g. diffusion, solubility, and absorption. Therefore, we should focus on properties of scrubbing liquid that not only collects the relevant particles but also cleans the gas. This issue is discussed further in the text. Disadvantage of gas scrubbing systems is their decreased efficiency, which may be influenced by applying organic liquids. Highcapacity sensible heat of the gas passes into a scrubbing liquid but cannot be utilized due to low output temperature of the liquid. Unless a different liquid is used, polluted water is produced and has to be pre-treated before entering the sewage system.

#### **5** Condensation and solubility of tar

Condensation of tar in gas is a complicated state, and the course of the process is affected by pressure in the assembly, temperature, proportionate composition, and content of particular tar components in a gaseous phase, mutual solubility of all components in the assembly, cold surfaces, condensation nuclei, and other factors. Following part of the paper approaches simplifying deals with and assumptions for understanding the condensation of mixtures; sources of basic data necessary for the calculation are presented.

Rapid cooling causes oversaturation of tar vapours, formation of condensation nuclei, and formation of undesired mist. If gas saturated with vapour cools slowly so that the gas temperature oscillates around condensation point for a sufficiently long time, small amount of droplets nuclei are formed and later develop into small droplets, as the gas continues to cool. If the gas is cooled suddenly, lot of droplet nuclei are formed and later form small droplets. The higher the vapour concentrations in the gas are, the more difficult it is for the droplet nuclei and droplets to increase in size. Highly diluted vapour form mist more easily than concentrated vapour. If dispersed droplets are very small, mist is stable.

Thanks to low tar concentrations in gas, a scrubber cooling the gas by injections of cold scrubbing liquid is an ideal environment for formation of fine persistent tar mist.

On the other hand, liquid or solid particles may, in addition to increasing in size thanks to condensation, coagulate in the stream of gas and form droplets or larger clusters.

Several literature sources are provided for better illustration of aerosol particle sizes. Perry et al., 1997 [9] say that mist particle sizes range from nanometres up to 10  $\mu$ m. According to more detailed overview given in Hemerka, 1995 [10], tar mist interval ranges from 0.1 up to 2  $\mu$ m. Perna and Riedl, 1944 [11] identically state that dispersed tar mist particles have 0.25  $\mu$ m in diameter. For comparison: Water mist particle size ranges from 1 to 40  $\mu$ m, and coal ash grain size ranges from 2 to 200  $\mu$ m.

In terms of thermodynamics, scrubbing liquid is considered a solvent in a tar scrubber, and tar is considered a soluble substance. However, solubility as a physical property always differs for a particular solvent-soluble substance pair.

If the solvent is polar, i.e. contains polar molecules (e.g. water), it may dissolve both polar and ionic compounds. If the solvent is not polar, i.e. contains non-polar molecules (e.g. benzene), it does not dissolve ionic compounds but may dissolve non-polar covalent compounds.

Tar, being part of hydrocarbon compounds, form covalent bonds predominantly. Nonetheless, other compounds containing ionic bond keep part of the covalent bond.

Tar and scrubbing liquid may be:

- 1. Fully soluble
- 2. Partially soluble (certain concentrations): This concerns tar scrubbing
- 3. Insoluble

# 6 Comparison of scrubbing liquids

Elimination of tar from gas using a wet scrubber technique differs for water and for RME. Although the scrubber may remove dust from the gas, dust should be removed prior to entering the scrubber as it may cause several difficulties in operation of the equipment.

#### 6.1 Scrubber using organic liquid

Several organic liquids dissolve tar compounds very easily. The two substances may be mutually miscible. Scrubber using organic liquid separates tar by absorption and the process is driven by differences in actual and equilibrium concentrations. Temperatures of scrubbing should be higher than condensation point of water contained in the gas because properly selected scrubbing liquid provides satisfactory results even for high temperatures (e.g. 70-90 °C) and the scrubbing liquid should be easily regenerated. Process of tar elimination from the gas using organic liquid may be divided into several stages:

– Direct gas-to-liquid contact cooling – gas inlet temperature should be higher than initial condensation temperature of tar so that fouling of connection pipes and equipment located prior to absorber is prevented. Certain amount of tar is to be absorbed by scrubbing liquid during a cooling phase. Tar vapors may be oversaturated and aerosol formed due to sudden temperature drop; this does not occur on large-scale basis, though. Outlet gas temperature should not be lower than water condensation point so that scrubbing liquid is not spoiled.

- Tar absorption and tar mist separation - mass transfer by diffusion, driven by concentration differences, occurs when gas gets in contact liquid under relatively with stabilized temperatures (gas may be further cooled in second stage of tar elimination). Mass flow dominates over mechanical actions in aerosol separation. Scrubber for mass transfer is preferred in a second stage of tar elimination. Efficient droplet separator should be arranged beyond the scrubber to avoid scrubbing liquid or coagulated aerosol droplets leaving the scrubber.

- Cooling and treatment of gas – temperature of gas leaving the scrubber is too high and cannot be lead directly into combustion engine; therefore the gas has to be further cooled. Gas is not saturated with tar vapours due to absorption of tar by scrubbing liquid and may be further cooled to a certain level without the risk of condensate formation. Or another stage of gas cleaning may be alternatively located beyond the scrubber, e.g. for elimination of hydrogen sulphide, hydrogen chloride or nitrogen compounds.

#### 6.2 Scrubbing using water

Water absorbs PAH only to a certain degree. It saturates quickly and tar starts to form a separate liquid phase – sub-micron type of aerosol (tar mist). This dispersion has to be separated so that the gas is pure enough. Tar separation in water-scrubber consists of following steps:

- Direct gas-to-liquid contact cooling – gas inlet temperature should be higher than initial condensation temperature of tar so that fouling of connection pipes and equipment located prior to absorber is prevented. Minority of tar may be absorbed by water during the cooling stage; tar vapours may be oversaturated and aerosol formed due to sudden temperature drop; water condensates in low temperatures, too.

- Tar mist separation - expected dispersity of tar mist reaches 0.1 to 2  $\mu$ m, according to literature. Tar droplets may be separated using certain types of scrubbers or other separators, see below. Their use is limited by a potential of micron and submicron particles to be separated.

- Gas heating – gas leaves the equipment saturated with tar, it is therefore recommended to slightly increase its temperature to prevent further condensation. Final outlet temperature has to be lower than maximum allowed temperature of the gas at the engine entrance.

Gas may be cooled to low temperatures using organic liquids, e.g. scrubbing oil. Outlet tar concentrations may be much lower, too. Disadvantage of this solution is the mixing of condensing water and organic substances. Used liquid is then hard to regenerate. If water is the cooling medium, NH<sub>3</sub>, H<sub>2</sub>S and HCl may be absorbed the gas, from too. However, absorption process is limited by water saturation, and successful separation of these gases is conditioned by regeneration of the scrubbing liquid.

Efficient droplet separator should be positioned beyond every scrubber so that final effect of the

method is not spoiled by droplets driven by scrubbing liquid and aerosol.

# 7 Methodology of measurement at Biofluid 100 gasification fluid generator

Research was performed at Biofluid 100 stand (see Fig. 1) which is equipment with stationary fluidized bed.

Simplified scheme of experimental equipment is presented at Fig. 2. Fuel is supplied from fuel storage tank equipped with shovel and is introduced via dosing screw with frequency convertor into reactor. Primary supply of blower compressed air is lead into reactor under the bed, secondary and tertiary supplies are located at two high-rise levels. Produced syngas is stripped of its solid particulate matter in cyclone. Output gas is combusted in burner equipped with stabilization burner for natural gas and individual air supply. [10] Ashes from reactor can be removed from tank located beneath bed. Power based heater for primary air supply is placed behind blower so that impact of air preheating may be monitored. In recent years, filters for research of efficiency of various methods of gas cleaning were attached to basic part of stand.



Fig. 1 Experimental equipment Biofluid 100 Reactor parameters:

- Capacity (in produced gas) 100 kW<sub>t</sub>
- Fuel demand (consumption, requirement) 150 kWt

Wood consumption
Air flow rate
40 kg.h<sup>-1</sup>
50 m<sup>3</sup>.h<sup>-1</sup>

Basic characteristics of operation at fluid generator are described in following respect:

• Operation of fluid generator – after ignition, fluid generator is operated in combustion mode so that its heating is quick. After achieving required gasification temperatures, secondary and tertiary air is supplied into generator and thus produced gas is immediately combusted and consequently heats up the generator. Air supplies are then shut off and generator is introduced into stable mode for specific and preset gasification temperature. Stable mode is achieved when amount of dosed fuel is not altered, amount of gasified air is even and temperature swings in middle section of gasification generator are stable within narrow range given by gasification temperature.



Fig. 2 Scheme of Biofluid gasifier

• Data entry of gasification process – monitored data are continuously recorded by computer in time interval of 10 seconds for each measurement. Following values are monitored:

• Frequency of convertor of dosing screw so that mass flow rate is determined;

• Temperature in various parts of equipment which is measured by thermocouples; position of thermocouples is given in detail in scheme at Fig. 2. There are 3 thermocouples along generator top, 1 thermocouple in cyclone and 2 thermocouples in in semi-coke pipe, 1 thermocouple in output gas pipe and 1 thermocouple measures temperature of primary air supply.

• Pressure difference between upper and lower sections of fluid generator (fluid bed);

• Pressure difference at orifice plate so that gas flow rate is determined;

• Pressure of the generated gas at the generator outlet and at the fuel storage tank.

Other values such as temperature and air moisture, primary air flow rate and its temperature have to be recorded manually.

Aim of the research was to determine concentrations of tar eliminated from the generated gas. Tar samples were collected using a methodology of Tar Protocol [11] and evaluated by weight spectrometer of a gas chromatograph.

# 8 Results and discussion

Several experiments were performed to assess theoretical requirements of gasification. Results are given in the graph below.



Fig. 3 Graph dependency of scruber efficiency and condensation point temperature on temperature

• Graph shows a strong dependence of scrubber efficiency on scrubbing water temperature and thus also on condensation point of tar after scrubber. If RME is used as a scrubbing liquid, this dependence is not obvious. This may be attributed to the fact that efficiency of tar absorption by RME is very high and no dependence can be observed.

Paper deals with elimination of tar from gas generated by biomass gasification. Tar is especially dangerous due to formation of thick condensate. Engine manufacturers do not provide limit concentrations of tar but they do require there are no condensates in the suction area. Problems related to corrosion (H2S, HCl. and NH4Cl) and fouling (alkali compounds, tar, dust, NH4Cl) may arise during cooling of gas from gasifier outlet temperature (ca. 700 °C) to engine inlet temperature (10-60 °C). It is recommended to eliminate dust at high temperatures (ca. 180-300 °C, depending on tar concentration and composition) and then lead the gas into the scrubber; this procedure helps prevent formation of sticky deposits (mixtures of tar and dust).

• Sudden cooling of gas creates fine persistent aerosol: tar mist. Size of aerosol droplets ranges from 0.1 to 2  $\mu$ m. If gas saturated with vapour cools slowly, small amount of droplets nuclei are formed which develop into small droplets, as the gas continues to cool. If the gas is cooled suddenly, lot of droplet nuclei are formed and later form small droplets. If the diffused droplets are very small, mist is stable. Efficient elimination of aerosol or prevention of its formation is a key factor in gas scrubbing.

• Combustion engines require gas free from mist with temperature sufficiently exceeding condensation point of the tar. Temperature of cool walls and external pressure in suction must be considered.

• There are vast differences between water scrubbing and organic liquid scrubbing.

#### 8.1 WATER

- Solubility of most tar components in water is low (except for heterocyclic compounds). Absorption eliminates by pure water heterocyclic compounds and highest PAH, especially. Polycyclic hydrocarbons with two to three rings remain in the gas and their concentrations allow for subsequent condensation. Separate liquid phase is either driven by water or leaves with gas in the form aerosol. Water supplied to real-life of equipment is assumed to be partially saturated with tar. Tar scrubbing is therefore based on elimination of aerosol.

- Disadvantage of gas scrubbing using water is a transfer of applicable heat of the gas into lowpotential production of environmentally unfriendly waste water; another disadvantage is the need to have a source of cool water or equipment for its production. During summer, facilities may utilize spray tower, sprinkled cooler or chiller. Chiller is very expensive.

- Method of scrubbing using water is relatively cheap.

#### 8.2 ORGANIC LIQUID

- Most of the tar components are fully soluble in the organic liquid. Ability of organic liquids to collect tar is therefore high, and the equilibrium concentrations corresponding with the selected temperature are much lower than equilibrium concentrations of tar condensation. Elimination of tar using organic liquid is based above all on diffusion, and therefore it is necessary to use an efficient mass exchanger. Similar to the technology of water scrubbing, the gas must be cooled. Gas output temperature should not drop below 75-80 °C (depending on gas moisture) so that scrubbing liquid is not impaired by the condensing water, and that no condensation of water disrupts the heat and mass flows in the scrubber.

- The advantage of organic scrubbing is high efficiency and ability to reduce tar condensation point well below the scrubbing temperature. Other positive aspects include: virtually wastefree facility and the ability to use gas heat for cooling of scrubbing liquid.

- Disadvantage of scrubbing using organic liquid is especially the high cost of scrubbing liquid, which must be constantly replenished due to evaporation.

- Type of organic liquid should be selected with respect to price, in accordance with Ewell classification and vapour pressure. Considering the need to minimize the liquid loss, scrubbing liquid pressure at operating temperature should be as low as possible.

• For economic reasons, it is desirable that the tar concentrations, or concentrations of other impurities in the gas, are minimized within the scope of primary methods (as a part of optimization). This reduces consumption of scrubbing liquid and requirements for gas scrubber efficiency.

Organic dust dispersed in the generated gas is difficult to wet with water and contains a large proportion of micron and submicron particles. Its removal in water scrubber is therefore very difficult. Alternatively, direct water cooler and electroseparator may be used. Elimination of dust using organic liquid is, from the physical point of view, relatively simple, and it is therefore possible to combine the process with tar scrubbing. It is necessary to be careful and take measures to prevent the formation of sticky deposits in the scrubber and find solution to liquid regeneration problems. Elimination of dust using textile filter seems to be a much easier solution for both tar elimination methods.

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