Analysis of biomass use in the energy sector in cooperation with the chemical industry

Master of Science Thesis in the Master’s Programme Sustainable Energy Systems and Industrial Ecology

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Department of Energy and Environment
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CHALMERS UNIVERSITY OF TECHNOLOGY
Göteborg, Sweden 2011
T2011-361
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Chalmers Reproservice
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ABSTRACT

The use of renewable raw material has increased every year and climate change is becoming a widespread issue in both public sector and enterprises. For both energy and chemical industry today’s most prominent alternative for fossil feedstock is biomass. Vattenfall plans to expand its use of biomass and for this reason cooperation between the energy and chemical industry with respect to biomass use could be of interest.

This Master’s thesis will look at biorefinery technologies giving solid residues, and further see if these residues can be used in co-combustion with coal. Another perspective will be to see how biorefinery technologies can be combined in order to produce “right” target products and residues. A carbon cycle and energy balance has been performed to see if integration between these industries can be sustainable out of a biomass supply perspective.

To evaluate different biorefinery technologies a classification and a ranking system has been developed. Categories in classification are base on Cherubini (2010) and chapter 3, 4 and 5 in this report. The ranking system is designed by seven criteria’s with different weighting factors, these are closely connected to the objective of this thesis. The c-cycle and energy balance will show how much biomass is needed to replace crude oil within chemical industry. Both calculated values are then compared with available farmland in EU-27.

This thesis discusses ten different technologies that give eight valuable solid residues, five of these are of special interest for co-combustion with coal. The ranking shows that fermentation 2nd is the most promising technology, both due to target product production and valuable residue. Further promising technologies are pyrolysis, gasification and hydrolysis. The c-cycle and energy balance shows that there is enough arable land to cultivate biomass and replace the use of crude oil.

Today there are good technical possibilities for cooperation between chemical and energy industry and there are several suitable residues for co-combustion with coal. Out of a sustainability perspective it is theoretical possible to exchange crude oil with biomass, but the complex land use question has to be further investigated. Also more research has to look into process integration of both different biorefinery technologies and process integration between biorefinery and coal fired power plant.

Key words: Biorefinery, biomass, chemical industry, residues
Analys av biomassa användningen inom energisektorn i samarbete med kemiindustrin
Examensarbete inom mastersprogrammen Sustainable Energy Systems och Industrial Ecology
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SAMMANFATTNING

Användningen av förnyelsebara råmaterial ökar för varje år och klimatförändringar
har blivit ett känt problem både bland offentlig sektor och privata företag. Både för
kemiindustrin och energiindustrin är biomassa ett av de bästa alternativen för att
ersätta fossila råmaterial. Vattenfall planerar att utöka sin användning av biomassa
signifikant, därav skulle ett eventuellt samarbete mellan kemiindustri och energi
sektorn kunna vara av intresse.

Detta ex-jobb avser att undersöka vilka bioraffinaderi teknologier som ger fasta
restprodukter och vidare vilka av dessa som kan sameldas med kol. En annan aspekt
är att undersöka hur olika bioraffinaderi teknologier kan kombineras för att producera
"rätt" produkter och restprodukter. En kolcykel och en energibalans är gjord för att se
om integration mellan dessa två industrier är hållbar utifrån ett biomassaperspektiv.

Utvärdering av de studerade teknologierna har gjorts genom att utveckla ett
klassificerings och rankning system. Kategorierna i klassificeringen är baserade på
Cherubini (2010) och kapitel 3, 4 och 5 i denna rapport. Ranking systemet är designat
av sju olika bedömningskriterier med viktnings faktorer, dessa är stark kopplade till
examensarbetets frågeställning. Beräkningar i kolcykeln och energibalansen visar hur
mycket biomassa som behövs för att ersätta råolja användningen i kemiindustrin.
Båda dessa värden jämförs sedan med odlingsbar mark i EU-27.

Detta examensarbete diskuterar tio olika grupper av bioraffinaderi teknologier som
ger åtta värdefulla restprodukter med fast karaktär. Fem av dessa är intressanta för
sameldning med kol vid värme- och elproduktion. Rankingen lyfter fram andra
generationens fermentering som den mest lovande teknologin, detta beror både på
passande målprodukt och en värdefull restprodukt. Andra lovande teknologier är
pyrolys, förgasning och hydrolys. Kolcykeln och energibalansen visar att det är
teoretiskt finns tillräckligt mycket land areal för att ersätta råolja med biomassa.

Det finns goda tekniska möjligheter för ett samarbete mellan kemiindustrin och
energiindustrin. Arbetet har visat att det finns flera restprodukter från bioraffinaderi
teknologier som passar för sameldning med kol. Utifrån ett hållbarhets perspektiv är
det teoretiskt möjligt att ersätta råolja med biomassa, men hur odlingsbar mark skall
användas är en komplek fråga som behöver studeras mer ingående. Även mer
forskning på processintegration både för bioraffinaderi teknologier och
processintegration mellan bioraffinaderi och kolkraftverk är nödvändig.

Nyckelord: Bioraffinaderi, Biomassa, Kemiindustrin, Restprodukter
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Preface

This report presents a master thesis project carried out in cooperation with Vattenfall Europe New Energy GmbH. The project was done from March 2011 to August 2011 at the Department Energy and Environment, Energy Technology at Chalmers University of Technology, Göteborg, Sweden. The project has been achievable due to finance by Vattenfall Europe New Energy GmbH.

The aim has been to give Vattenfall GmbH a future prospect of possibilities for biomass utilisation in EU-27. Focus has been on different biorefinery technologies and their residues, with a sustainability systems perspective. The project has been carried out with Dr. Jan Grundmann (Vattenfall GmbH) as supervisor and Assoc. Prof. Erik Ahlgren (Chalmers) as examiner. We would like to thank both of you for your time, patience and feedback.

We highly appreciated Andreas Beyer and Michael Klysz at Vattenfall GmbH for their assistance and advice throughout the project. Special thanks to Energy Technology Department for hosting us during our work period. We would also like to show our gratitude to all people we have interviewed for their time, patience and valuable knowledge.

Göteborg August 2011

Martin Johnsson and Anna Åhlund
Notations

Abbreviations

AHR  Solid dilute-acid hydrolysis residue  
CCS  Carbon Capture and Storage  
DM  Dry matter  
DR  Concentrated liquid distillation residue  
ETBE  Ethyl tert-butyl ether  
ETS  European Emission Trading Scheme  
EU  European Union  
FAME  Fatty acid methyl ester  
FBC  Fluidised bed combustion  
HHV  Higher Heating Value  
HTL  Hydrothermal Liquefaction  
LHV  Lower Heating Value  
LPS  Lignin Precipitation System  
PBC  Pulverised bed combustion  
PVC  Polyvinylchloride  
RME  Rape methyl ester  
SHF  Separate Hydrolysis and Fermentation  
SNG  Synthetic Natural Gas  
SSF  Simultaneous Saccharinification and Fermentation  
SSF R  Solid SSF residue  
WC  Wood chips  
WWT  Waste water treatment

Elements

C  Carbon  
Ca  Calcium  
Cd  Cadmium  
Cl  Chlorine  
Co  Cobalt  
K  Potassium  
H  Hydrogen  
Mg  Magnesium  
N  Nitrogen
\( Na \) Sodium
\( O \) Oxygen
\( P \) Phosphorus
\( Pb \) Lead
\( S \) Sulphur
\( Zn \) Zink

**Chemical compounds**

\( CH_4 \) Methane
\( CH_3OH \) Methanol
\( C_2H_4O_2 \) Acetic acid
\( C_2H_5OH \) Ethanol
\( CH_2O_2 \) Formic acid
\( C_3H_6O_2 \) Lactic acid
\( C_3H_8O_3 \) Glycerol/Glycerine
\( (C_6H_{10}O_5)_n \) Cellulose
\( C_9H_{18}O_2(OCH_3)_n \) Lignin
\( CO \) Carbon monoxide
\( CO_2 \) Carbon dioxide
\( H_2 \) Hydrogen gas
\( HCl \) Hydrochloric acid
\( HMF \) 5-hydroxymethylfurfural
\( H_3O_4P \) Phosphoric acid
\( H_2SO_4 \) Sulphuric acid
\( KOH \) Potassium hydroxide
\( N_2 \) Nitrogen gas
\( NaOH \) Sodium Hydroxide
\( SO_2 \) Sulphur dioxide

**Words**

*Feedstock* Stream of raw material used to produce valuable products

*Residue* Process waste that can be both valuable or non valuable

*Target product* The main product leaving a biorefinery technology
Units for calculations

- **g**: gram
- **G**: Giga, $10^9$
- **ha**: hectare
- **J**: Joule
- **kg**: kilogram, $10^3$ gram
- **kg_{dm}**: kilogram of dry matter material
- **M**: Mega, $10^6$
- **tce**: tonne of coal equivalent
- **toe**: tonne of oil equivalent
- **t_{dm}**: tonne of dry matter material
- **P**: Peta, $10^{15}$
- **ppm**: Parts per million
- **wt%**: weight percent

Symbols
- Operation
- Intermediate product
- Input/Feedstock streams
- Output/Product streams
- Process streams
- Recirculation streams
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1 Introduction

The Bruntland commission, the Kyoto Protocol and the Rio conference is a few examples of attempt to achieve climate mitigation. One of the main objectives for the European Union within the environmental field is to increase the use of biomass. The region is committed to increase the proportion of renewable energy to 20% of total energy consumption by 2020 (ENCROP, 2010).

To meet these goals, Vattenfall plans to increase the use of biomass to 5-10 million ton per year and aims to control and invest in the whole value chain from cultivation to end use. This will lead to new areas for cultivation of biomass, new areas for food production and new logistic solutions for biomass transportation (Grundmann, 2011). Due to the fact that many energy enterprises will become players within the biomass market, to fulfill their climate ambitions, cooperation’s between energy sector and the chemical industry with respect to biomass use could be of interest. Energy producers do have an interest in how the biomass can be used and which technologies in chemical industry that gives the best residues as energy carrier back for use in energy services. One example of cooperation can be an energy enterprise producing the biomass and sells it to a chemical industry. The chemical industry refines the biomass to high value chemicals and the residues go back to power producer for co-combustion in a coal fired power plant. The excess heat from this plant can be used in the chemical refinery if they are placed closely (Grundmann, 2011). In this way coal power plants and chemical industry create an industrial cooperation site, where waste heat and waste products from biomass are used. To give Vattenfall the prospect to make a relevant forecast for the potential demand for biomass within the chemical industry in EU-27, including pharmaceuticals, a carbon cycle and an energy balance is performed.

1.1 Objective

Both energy and chemical industry shows more interest in alternatives for their fossil feedstock, the most prominent alternative today is biomass. The purpose with this project is to investigate different biorefinery technologies and the future market for target product from these technologies. The aim is to see if Vattenfall and other actors in energy industry could benefit from cooperation with other branches, such as the chemical industry.

Due to purpose of this project and the interest of the constituent, Vattenfall Europe New Energy GmbH in Hamburg, four main questions has been formulated as the framework for this project.

- What solid residues do different biorefinery technologies give?
- Do the residues from biorefineries fulfil the quality requirements for the combustion processes within coal fired power plants?
- Which combination of biorefinery technologies is best for creating synergies between the chemical and energy sector?
- Is cooperation between chemical industry and energy sector sustainable out of a biomass supply perspective?

By elaborate a classification system and a ranking system of biorefinery technologies the results will be presented, together with demand for biomass if crude oil would be exchanged.
1.2 Limitations

The master thesis limitations are here defined, this to prohibit the work from being too broad or incomprehensible to handle within the timeframe.

The geographical area is set to EU-27; other different geographical areas are sometimes discussed briefly to be able to give “hands on” examples. This area is not valid for the biomass market which is a global market with no geographical restrictions. The two industries looked into are energy and chemical, with a focus on chemical industry regarding future markets and technologies of use. Regarding infrastructure this project does assume that existing infrastructure in EU-27 is intact and no development has been taken into account.

Different types of technologies to refine biomass will be investigated, the focus will be on technologies giving solid residues. The technologies will be studied with the focus on input and output. The quality requirements for fuel of the coal fired power plants will only be discussed directly with Vattenfall professionals.

In evaluation of process integration this master thesis only looks at which processes that can be combined to produce desired products. Quality of excess heat leaving specific processes and more exact process integration recommendations will not be a part of this project.

Regarding the carbon cycle and energy balance the result will be affected by a number of parameters. Limitations for calculations regard yield, conversion efficiency and type of raw material used in chemical industry. The yield of energy crops vary a lot with both geographical location and spices cultivated. This is taken into account by doing calculations on three different yields. The fact that biomass yield most likely will increase in the future with optimised and effective cultivation standards, has been neglected to simplify calculations. All calculations will be performed with today’s crop yields. Assumptions around conversion efficiency are complex due to different technologies, in this master thesis the conversion rate is set to when syngas is produced from crude oil or biomass, respectively 97% and 87% (Dinjus, 2009). The crude oil used for energy and fuel in chemical sector are in these calculations neglected. Only the crude oil used as feedstock is included.

1.3 Outline

To be able to investigate different biorefinery technologies and to find adequate background information a literature review and qualitative interviews has been carried out.

1.3.1 Literature review

The literature studies will focus on scientific articles and books available in databases at library of Chalmers University of Technology. There are lot of articles and books written in the field of biorefinery, much of this literature focus on the biomass process and chemical/biochemical reactions and processes. Some of the articles describe the way of biomass through the plant and type of products that are produced in different steps of a biorefinery. Most of the articles are written from a chemical or a biomass production aspect and just a few from the energy point of view. Only a small number of articles discuss the residues composition from biomass used in biorefineries.

Books where used when information about well known technologies and subjects where searched for. For more recent research result scientific articles contributed with
adequate information. Research in the open web has also been implemented to find adequate company and organizations reports; here the search engine Google has mainly been used.

European Chemical Industry Council, Cefic, and ENCROP where mainly used to find data on crude oil use, energy content in energy crops and energy crop yield/ha/year. Cefic represents 29,000 chemical companies in Europe. The council try to embody its members and interacts on their behalf with international and EU institutions, non-governmental organizations and the international media (Cefic, 2011). ENCROP is a European project to promote the production and cultivation of energy crops and to increase the knowledge of different actors along the supply chain. The project was carried out during a three year period, from October 2007 until August 2010 (ENCROP, 2010).

1.3.2 Qualitative interviews

To fill information gaps from literature studies interviews with experts have been valuable for important data collection. Interviews with concerned companies, power plants and institutions have been performed.

The interviewer together with the interviewee created the result through a dynamic conversation. During the literature study four different areas of interview objects of special interest were been obtained, these are following:

- Chemical industry to understand the residues leaving the chemical plant
- Chemical industry to understand their requirements for future industry cluster
- University experts within field of biomass use and biorefinery technologies
- Vattenfall for information about the coal fired power plants requirements in Germany

A list of connected people can be found in Appendix A, each person’s is displayed with title, contact details and date for interview.
2 Background

This chapter will give the reader a short introduction to biomass as raw material, what is defined as biorefinery, how does a general combustion process works and what does it mean to co-fire biomass and coal. These subjects are necessary in order to assimilate the following chapters of this report. This chapter will also include two examples, one of an exciting biorefinery and one of an integrated industrial site.

2.1 Biomass

Biomass is a renewable energy source and the demand is expected to increase over the coming decades. Interest in biomass as an energy source and raw material is rising for several reasons. The use of biomass is CO₂ neutral due to the photosynthesis which converts sun energy and CO₂ to biological material. It is also an alternative for fossil fuels which is a limited resource (non-renewable) and several energy crops are also alternative agriculture products where other crops are not suitable. The four types of biomass discussed are: wood, recycled material (for example demolition wood and waste paper), agricultural by-products (for example straw and tops) and energy crops (for example willow and Miscanthus) (de Mol, Jogems, van Beek, & Gigler, 1997). Biomass lignocelluloses materials refer to plants that are composed of cellulose, hemicelluloses and lignin (Doherty, Mounsavioun, & Fellows, 2010), see Figure 1. Compared to fossil raw material, biomass material generally has a lower fraction of carbon, more oxygen and less hydrogen (Cherubini, 2010). Cellulose gives trees and woods their strength; these fibers are long, flexible and natural linear homopolymer (Lui, 2010). The chemicals structure of cellulose is \((C_6H_{10}O_5)_n\) and is made by long chains of glucose molecules, \((C_6\text{ sugar})\) (Cherubini, 2010). Lignin \((C_9H_{10}O_2(OCH_3))_n\) is the natural phenolic glue that holds together cellulose fibers and act as a natural protection against biological degradation of cellulose and hemicelluloses (Lui, 2010).

Although biomass is stated as CO₂ neutral the future use of biomass has several limitations, both concerning limited availability of land and logistic problems. If explored biomass really is CO₂ natural depends on cultivation methods and previous use of land to mention a few parameters. The costs for biomass as a fuel and raw material are divided into three parts: the purchase price, the logistic costs for the collection and the costs for establishing and running the energy plant (de Mol, Jogems, van Beek, & Gigler, 1997). Costs of logistics may determine a major part of feasibility, especially when biomass costs are relatively low (Annevelink & de Mol, 2006). Today the only commercial biofuel, here referring to only transportation fuel, products comes from plant oils esters (FAME) of long-chain fatty acids or ethanol from the enzymatic digestion and fermentation of starch or sucrose. These raw materials are competing over arable land for food production (Doherty, Mounsavioun, & Fellows, 2010). As for all agriculture products biomass has losses during storage, these losses can be positive effects (moisture losses) or negative losses (dry matter losses) (de Mol, Jogems, van Beek, & Gigler, 1997).
In Figure 2 the product refinery chain of biomass within chemical industry is illustrated, here it can be seen that there are four main steps towards final application. Today there are only a few biorefineries operating but this will have to change if climate goals are going to be met and if industries want to secure a steady supply of raw material.

![Figure 2: Pathways for biomass from raw material to chemical applications (Luo, van der Voet, & Huppes, 2010). Building Blocks are the pre steps before end use application is produced.](image)

2.2 Biorefinery

To get a more efficient utilization of biomass the users must exhaust as much of the biomass as possible. This means that both the energy sector and the chemical industries can process the same biomass, but different parts of it. The concept for this cogeneration is named biorefinery. In the biorefinery all necessary biological, chemical and energy processes will be handled at the same site but in different processes. These processes will give different chemicals with possible utilisation in the chemical industries instead of fossil products but also energy services in different ways. The International Energy Agency (IEA) describes the biorefinery process as: “Biorefining is the sustainable processing of biomass into a spectrum of marketable products and energy” (IEA Bioenergy, 2010). The concept needs a lot of different methods to separate the biomass into their components (i.e. carbohydrates and proteins).

There are different processes to separate the valuable parts from the biomass feedstock i.e. mechanical, chemical, bio chemical and thermochemical. This structure with separation in different steps or processes is similar with the oil refinery and that’s the reason of the similar name. A major difference between the biomass feedstock and the fossil feedstock is that the fossil compounds most often are homogeneous, but the biomass is not (Cherubini, 2010). This variety of the feedstock gives disadvantages which means that the site needs a lot of different equipments and technologies but also seasonal problems connected to that the refinery perhaps can’t produce the
maximal capacity during the winter. But there are also advantages, these are often connected to the range of different products and the biorefinery can easier change their production due to the raw material and technologies than the petroleum based can do (Dale & Kim, 2006). Another advantage is the fact that biorefinery uses biomass that do not affect the environment as negative as coal (Brown, 2003). There are possibilities to produce gaseous and solid fuels from the lignin part. This fuels are often used for the own plant but there are also possibilities to connect the system to a bigger power plant i.e. co-fire the lignin part with coal in a coal fired power plant.

### 2.3 Combustion processes

Even if Europe has a heavy climate debate and the European Union (EU) works with ambitious climate targets, coal is still one of the two most important energy sources in the EU. Together coal and nuclear produces a clear majority of the electricity and heat in EU and the coal alone stand for around 30% of the electricity production (Eurostat, 2009). The situation is similar in Germany and here around 43% of the produced electricity comes from coal (24.6% lignite and 18.3% hard coal) and coal is also an important source for heat (Bundesministerium für Wirtschaft und Technologie, 2010). There are also a lot of new plants with coal especially in “new economies”. China and other developing countries are building coal fired power plants due to that these are quiet cheap, easy to operate and quick to build, which is an advantage due to the fast increasing energy demand in these regions (Breeze, 2005).

There are some problems with this big consumption of coal; first the coal is an ending resource even if coal has a long expire date, approximately 150 years with today’s reserves and extraction rate (British Petroleum, 2010). The second problem is the formation of carbon dioxide in the combustion processes using coal (Miller, 2005). Only in Germany around 40% of the CO$_2$-emissions come from coal (Bundesministerium für Wirtschaft und Technologie, 2010). The third problem is the increasing price of coal as a fuel. The increase in fuel price together with a forecasted price increase of the European Emission Trading Scheme (ETS) will give a higher marginal cost for coal as technology (Erdmann & Zweifel, 2008). These three reasons will give good incitements to find other solutions or improvements. These questions are also important for Vattenfall due to that they have 18 coal power plants (11 in Germany) producing heat and electricity (Vattenfall AB, 2010).

Changing directly to other sources from coal is very hard due to the big share of the market. There are a lot of different possible improvements such as efficiency improvements of existing plants, carbon capture and storage (CCS) and co-firing with biomass (Miller, 2005), (Termuehlen & Emperger, 2003). Switch the fuel in some parts to biomass is difficult and it’s important that the biomass follows the quality requirements. The size of the particles is very important due to that the combustion processes can handle the particles or not.

The basic idea of a coal fired power plant and all other combustion power plants is to let the fuel burn in an excess of air. The air will be heated and in a heat exchanger system the air heats water to steam. This steam drives a steam turbine, which is connected to a generator for producing electricity. After the turbine the steam is cooled to water again and returns back to the heat exchanger system again. The releasing heat from the cooling of the steam can be connected to a process or a district heating net (Miller, 2005).
There are possibilities to burn biomass directly as the humans have done for many years. In electricity production unfortunately this gives a poor efficiency, only around 25-30%. So direct firing of biomass is not a good idea, especially not in the case of electricity production. For this production steam is necessary to drive the turbine and to produce good steam quality, high temperatures are important. However it can still be high enough temperatures for heat production. Another option using biomass for electricity production is to use co-firing, this increase the efficiency close to the normal efficiency for a coal power plant (35-40%) but is only possible to add a smaller fraction of biomass in the process (Breeze, 2005).

2.4 Biomass co-firing

The term co-firing or co-combustion means combustion of two or more fuels parallel in the same boiler for energy services (Leckner, 2007). Co-firing technology is one of the options for EU-27 countries to partly meet its climate mitigation goals. This type of technology also have an economic advantage, it is 2-5 times less cost intense than other bio-electricity generating technologies (Berndes, Johnsson, & Kjärstad, 2010) and the technology is less complex than other biofuel technologies (Leckner, 2007).

For co-combustion with coal different type of boilers can be used here five different types are mentioned, but co-combustion can be applied on any type of boiler. A few examples of boilers are pulverised bed combustion (PBC), fluidised bed combustion (FBC) and combustion on added grates inserted in pulverized coal boilers (Leckner, 2007). To make co-firing possible it is important to take quality of fuel into account. Here are four important factors obtained (Grundmann, 2011).

- Particle size
- Chlorine content
- Sodium and Potassium content, ash melting point
- Sulphur content

These factors has to be right for all fuels going into the boiler, but the main fuel is the one determine the quality properties of other fuels coming in.
There are both positive and negative synergy effects with biomass/coal co-combustion. Some of the positive synergies are the reduced consumption of coal and therefore also mitigation of fossil CO₂ emissions (Leckner, 2007). Using biomass in co-firing has a relatively low impact on the conversion efficiency. Neither an uncertainty with fuel supply is put on risk, the plant is then depending on two different types of fuel. Another advantage is that using biomass in coal fired power plant, existing infrastructure can be used. This give incentive to use local fuel (Leckner, 2007) and an expanded use of biomass is not constrained (Berndes, Johnsson, & Kjärstad, 2010). The disadvantage of co-combustion can be additional costs for equipment and drying of biomass, lack of experience is also seen as a negative aspect. The mix of fuels may also cause negative synergies in combustion process (Leckner, 2007).

In Figure 4 a pulverised fuel boiler is scheduled, the different combustion zones can be viewed. Originally the fuel injected in a PBC boiler at re-burning zone was natural gas, but now is has been seen that it is here possible to inject solid fuels in this zone as well (Leckner, 2007). This solid fuel could be biomass if the grinding technology can give sufficient small particles. It is also important that sufficient residence time and temperature in the furnace for burnout of the char coal particles is kept and at the same time as ash deposition is avoided (Harding & Adams, 2000).

According to Leckner there are four main groups of the impact of co-firing, these are (1) energy content and volatiles, (2) precursors to gaseous emissions, (3) ash-forming elements and (4) trace elements. Often do biomass fuels have lower heating value than coal and higher moisture content than coal so total energy content is differenced. By drying biomass in an external facility the heating value can be increase and moisture content can be lowered (Leckner, 2007).

Plants size restrictions are not theoretical rather economical due to the limitation on how far biomass can be transported. A feasible level of biomass input is probably 5 % to 10 % of the maximum boiler capacity (Leckner, 2007), (Berndes, Johnsson, & Kjärstad, 2010). Both for PBC and FBC this amount of biomass input can be handled without any major changes of technology or problems with corrosion (Leckner, 2007), (Berndes, Johnsson, & Kjärstad, 2010).

Conclusions out of this is that co-firing biomass with coal can have a positive impact on the lignocelluloses supply system, which would have a positive spin off on infrastructure for biomass logistic and storage. Coal co-fired with biomass can clearly increases the near-term demand for biomass in many countries (Berndes, Johnsson, & Kjärstad, 2010).

2.5 Paper Mill

One example of a today existing biorefinery is a modern chemical paper mill. There are two existing main process in the pulping process, either the mill use mechanical
pulping or chemical pulping. The pulp consists of cellulose fibres and the mechanical or chemical process names describe how these fibres are extracted from the wood.

The mechanical process grinds wood or wood chips with electrical power. The fibres are separated from one another with a series of rotating discs. This process has a very high efficiency, 90-98% (Brown, 2003). Mechanical pulp fibres are stiff and mostly intact. The pulp also includes a large amount of a smaller material called fines. This fine consists of fragments from the fibre wall and broken fibres. The fines have an important role for optical skills which give smooth printing surface. This is one of the reasons mechanical pulping is used for production of newsprint (Brännvall, 2009). Another important difference due to the chemical pulping is that the lignin is not removed from the fibres. The lignin is the reason why newsprint is weaker and also the reason for darken the newsprint over time. The mechanical process is very energy intensive and uses everything from the tree, therefore it is not interesting out of a biorefinery perspective (Brown, 2003).

In the chemical pulping process the fibres are released chemically. The lignin that acts like glue between the fibres is removed by chemicals and thereby releases the fibres. The problem with these chemicals is that they are strong and not specific for lignin, so even other components like carbohydrates are released. There are two different main processes in the chemical pulping; kraft and sulphite (Brännvall, 2009). The sulphite process gives a lighter, easier to bleach pulp but the kraft process have a god established system for chemical recycling and use the lignin as a fuel. Due to that fact that even some hemicelluloses are damage in the chemical process the pulping yield is only around 35-60% of input biomass. The fibres are stronger than the mechanical and around 70% of the total pulp production is based on a chemical production (Brown, 2003).

Around 20% of the chemical pulp is from the sulphite process which is often connected to that the pulp from kraft process has better properties. But still the sulphite process is important in several countries and for some special paper qualities. The advantages for this sulphite process are: higher brightness of unbleached pulp, lower odour problems, in some situations higher yields and lower investment costs (Alén, 2000).

As described earlier the kraft process is the dominating chemical pulping process around the world. The chemicals the process use (named cooking chemicals) are sodium hydroxide and sodium sulphide. The process can also work with only sodium hydroxide and in that case the process is named soda cooking (Brännvall, 2009). In a short description three different liquors are used in the kraft process; white, black and green. The white liquor is the clean one going into the cooking process together with wood chips. After the cooking the black liquor comes out and contains dissolved organic substances such as lignin and inorganic cooking chemicals. The black liquor is evaporated and burned to smelt. The smelt is solved in water and named green liquor and with a process called causticising the green liquor is converted to white (Brännvall, 2009). The black liquor is separated from the pulp with washing and after evaporation the solid content is around 65-80%. The lignin from the process contains around 6% sodium of the dry mass (Alén, 2000).

Before the developments of the chemical recycle system the kraft process wasn’t competitive with the sulphite process, due to high price of sodium hydroxide. The high cost was connected to the use of sodium carbonate as make up chemical to
produce sodium hydroxide. Instead the mills started to use the inexpensive sodium sulphate for this. The produced pulp from the kraft process is strong and that’s the reason for the name kraft, which means “strong” in German and Swedish (Brown, 2003).

The black liquor is the key point in this process related to the energy issue. The black liquor contains lignin and degraded polysaccharides. This lignin has limited applications but has a high heating value compared to raw wood and is for that reason a good boiler fuel. Also the degraded polysaccharides are used as boiler fuel even if the heating value is low. In the pulp mill the black liquor is used in a recovery boiler (Brown, 2003).

2.6 Stenungsund chemical industry cluster

Chemical industry cluster in Stenungsund, Sweden, shows what an industry cluster can look like and what benefits that can occur in such cluster. Since 1960 Stenungsund, positioned 50 kilometres north of Gothenburg, has been a central location for Swedish petro chemical industry (Berglund, 2010). The companies in Stenungsund are now deeply integrated with many synergy effects and a lot of to-know-how is available in this cluster, this type of cluster also has a competitive advantage (Zhao, Watanabe, & Griffy-Brown, 2009). Such industry cluster often do operate in the same market, share suppliers, education institutes and will most often share the same challenges and fears (Zhao, Watanabe, & Griffy-Brown, 2009). This is also the case in Stenungsund.

The first three companies involved in making Stenungsund a chemical industry hub was Esso, Modo and Unifos (Borealis, 2010). During the 1960’s further expansion of the cluster was needed due to international market outlooks, Sweden didn’t want to be dependent on imported basic petro chemical products (Berglund, 2010). Not only did Stenungsund expand, the chemical industry also wanted total integration between refinery plants, steam cracking and polymerisation (Berglund, 2010).

Today Stenungsund chemical industry is one of the major integrated industry cluster which has developed to be one of its kind worldwide. Here six chemical companies and two energy companies operating together to optimize energy and raw material resources. What characterizes this cluster is the hard niche on high performing products and materials, the electricity supply to the cluster comes from Nordic energy mix which is nearly CO₂ free, and this is the first cluster taking clear actions towards green chemistry (Onsander, 2011). Figure 5 shows how complex the cluster is and the “pipes” can be followed to see how the resource flows are connected.

Danish Dong and Swedish Vattenfall is the gas, water and electricity supplier to the plant. Since a couple of years Dong delivers gas with a small part of biogas mixed in the natural gas, this part could increase if biogas production would be put into focus (Onsander, 2011). Vattenfall provide the plant with electricity and runs a water treatment plant which supplies the industry and part of Stenungsund municipally with water (Stenungsund Kommun, 2011). AGA Gas produces nitrogen, oxygen, carbon dioxide and argon from air. Both nitrogen and oxygen is delivered through pipelines to other industries in the cluster, AGA Gas is not included in the cluster group and is treated as a normal supplier (Onsander, 2011). Nitrogen is here used to prohibit fire and explosions (Stenungsund Kommun, 2011).
The Stenungsund cluster “starts” with Borealis cracker facility which produces ethylene and propylene from butane, ethane, naphtha and propane. Switches between these feedstock’s can quickly be made (Onsander, 2011). Ethylene and propylene is transferred to the own polyethylene plant, using a self developed technology, producing plastic products (Stenungsund Kommun, 2011). The oil cracker produces hydrogen gas at 110bar which is used by actors in cluster. Fuel gas and hydrogen is mainly used by the industries for steam production. Many of the industrial processes in Stenungsund requires steam at very high pressures, this implies own production of steam to fulfil the right requirements (Borg & Stralström, 2007). Fuel gases and hydrogen are residues from Borealis production site in Stenungsund (Borg & Stralström, 2007) and ETBE, ethyl tert-butyl ether, leaving the Borealis cracker is sold direct and mixed into petrol (Onsander, 2011).

AkzoNobel produces special chemicals like ethylene oxides, surfactants and amines from ethylene, oxygen and ammonia. Through conversion of saline to chlorine with electricity Ineos crates their base product. Chlorine together with ethylene forms the plastic material polyvinylchloride (PVC) (Stenungsund Kommun, 2011). Both AkzoNobel and Ineos import parts of their ethylene demand (Lindkvist, 2010).

Perstorp Oxo is the youngest and smallest member of Stenungsund industry cluster. They produce aldehydes, alcohols and organic acids which are used in colour, coatings and safety glass. Raw materials used are ethylene, propylene and natural gas (Lindkvist, 2010). Perstorp Oxo excess heat is transferred into Stenungsund municipally district heating system, the excess heat stands for 90 % of the total heat in the system (Borg & Stralström, 2007).
The chemical industry in Stenungsund has together started an initiative called "Hållbar kemi 2030 – Kemiföretagen i Stenungsund" (Lindkvist, 2010). Their common vision for the future states “In 2030 Stenungsund will be the hub for the manufacturing of sustainable products within the chemical industry. Our operation is based on renewable feedstock and energy and contributes towards a sustainable society” (Lindkvist, 2010). Behind this statement lie plans to have an integrated industry site which uses raw materials and produce products that are a part of a natural ecosystem, in addition have efficient energy consumption, develop existing system further to take care of excess heat, recycle plastic materials and use locally produced transportations fuels. This to avoid furthers dependence on fossil feedstock (Tysklind & Söderberg, 2010). The industry cluster can’t achieve these changes on their own, they also need system service from municipally and both national and EU-27 policies have to change to favour use of renewable raw materials in the chemical industry. Today there are three main value chains for biomass feedstock, these are energy generation, vehicles fuel and materials/chemicals. EU-27 policies of today support the first two of these values chains through tax reduction and cap and trade scheme. For chemical industry to become competitive for biomass feedstock this picture has to change. According to Robert Onsander Hållbar Kemi 2030 is a vision and not a detailed action plan (Onsander, 2011).

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1 English translation – Sustainable Chemistry 2030 – Chemical industry in Stenungsund
3 Biorefinery technologies

The different parts of the lignocelluloses feedstock can be used for different purposes and therefore is it necessary to separate it in different steps. The different parts can as described before create different valuable products such as chemicals or fuels. The technical processes have the aim to depolymerise, break up polymers to monomers, and deoxygenating (removal of oxygen) the biomass components (Cherubini, 2010). The first part is primary refinery which divides the raw material after decided specifications such as size (de Jong, van Ree, & Elbersen, 2006). The second part is described well by Cherubini and includes four different main categories; Thermochemical processes, Biochemical processes, Mechanical processes and Chemical processes (Cherubini, 2010).

Due to the objective of this Master thesis the focus will be to find technologies that give solid residues which can be used for co-firing in coal fired power plants. Other biorefinery technologies, giving non solid residues, will also be presented together with their possibilities for the future, but in a shorter context (Clements & Van Dyne, 2006). Chapter 3 is a summary and explanation of all for this master’s thesis relevant biorefinery technologies. Information about target products and residues will be found in chapter 4 and 5.

3.1 Mechanical processes

Cherubini describes the mechanical processes as “Processes which do not change the state or the composition of biomass, but only perform a size reduction or a separation of feedstock components” (Cherubini, 2010). In the biorefinery system these processes often are first in line because a size reduction is necessary to continue with other processes. Examples of these other processes can be drying, pelletizing and granulating. Another common mechanical process is the barking process. This is a process especially used in the paper mills but also in other plants to bare the hemicelluloses part of the tree (Cherubini, 2010). The residue is often bark or sawdust.

3.2 Thermochemical processes

These processes use high temperatures and have short reaction rates converting biomass into energy and/or heat and power but also in some of the processes produce chemicals (Cherubini, 2010). In this report are pyrolysis, gasification and liquefaction described. A regular combustion for heat and power is also a thermochemical process with the purpose to produce heat and power. The residues after such process are mostly ash, which is no valuable residue (Miller, 2005). The difference between pyrolysis, gasification and combustion is the temperature. Pyrolysis work with the lowest temperature (between 400 and 700°C), gasification operates at 700 to 900°C and finally combustion between 800 and 950°C (Neves, Thunman, Matos, Tarelho, & Gómez-Barea, 2011). The yield from thermochemical processes is around 60% and that is high compared with other biorefinery processes (Eriksson, 2009).

3.2.1 Pyrolysis

Pyrolysis is a technology that treats biomass with moderate temperatures around 400-700°C. The heat is transferred from surrounding gas in absence of oxygen to the biomass particle (Dermibas, 2009). The particle size is depending on the pyrolysis
model, the heating rate and residence time. The heating rate, the speed of temperature increase, is important due to the production rate of volatile gases. The main product types from pyrolysis are volatile gases and liquids, with the residue char coal. The char coal is a solid product similar to coal, see chapter 5.2. Volatile gases are a mix of CO, CO₂, H₂, Methane and Ethylene. These are interesting gases to condensate in the production of chemicals (Clements & Van Dyne, 2006), (Neves, Thunman, Matos, Tarelho, & Gómez-Barea, 2011). The liquid is called bio-oil and consists of complex mixture of different compounds, see chapter 4.2. If the target is to produce as much gas as possible in the pyrolysis process a low heating rate and a high temperature is used, the opposite will favour char coal. (Dermibas, 2009). It is possible to use pyrolysis with both hemicelluloses directly and separated lignin.

There are different types of pyrolysis, slow, fast and flash. Slow pyrolysis means pyrolysis with a low heating rate, approximately 0.1-1°Cs⁻¹ and reach a maximum temperature around 500°C (Bahng, Mukarakate, Robichaud, & Nimlos, 2009). This heating rate corresponds to traditional pyrolysis and will favour production of char coal. This technology has been used for thousands of years in production of char coal as a fuel and the residence time is between 5 to 30 minutes.

The fast model is more efficient for extraction of liquid products and uses a higher heating rate in combination with smaller particles (<1mm) (Dermibas, 2009). The heating rate is approximately between 10 to 200°Cs⁻¹ and the residence time is only up to a couple of seconds (Bahng, Mukarakate, Robichaud, & Nimlos, 2009). After the fast heating, a rapid cooling of the pyrolysis vapour is necessary to get the bio oil (Bridgewater & Peacock, 2000). Even with cooling some gaseous products will be formed and the process will give around 60-75wt% liquid bio-oil, 15-25wt% char coal and 10-20wt% non condensable gases (Bridgewater, 2003).

Flash is an improved model of the fast pyrolysis with higher heating rate, up to over 1000°Cs⁻¹ and a residence time of only a few seconds and uses very small particles, under 0.2mm (Dermibas, 2009). Due to the short residence time this process requires smaller particles than fast and slow pyrolysis (Bahng, Mukarakate, Robichaud, & Nimlos, 2009). If the biomass raw material contains heavy metals, most of these will be in the char and the bio-oil is almost heavy metal free (Kuppens, et al., 2009).

### 3.2.2 Gasification

Gasification is also a form of pyrolysis but in this report, as in much of the literature, gasification will be treated separately. In this technology the solid biomass will be treated with higher temperatures (700-900°C) with an oxygen supply around 35% of the demand for combustion. This treatment converts the biomass to a combustible gas mixture. This gas mixture is called producer gas or fuel gas (Faaij, 2006) and consists of carbon monoxide (CO), hydrogen (H₂), methane (CH₄), nitrogen (N₂), carbon dioxide (CO₂) and some smaller fractions of higher hydrocarbons (Brown, 2003). The gas can be used as fuel for heat and power services, which is most common, but can also be used as feedstock for production of liquid fuels and chemicals. The process is endothermic and requires heat, this heat can be produced either from the produced gas or from burning some parts of the biomass directly (Brown, 2003). Also the gasification process will give some char coal (around 10%) even if the high temperatures favour gas production (around 85%) (Zhang, Xu, & Champagne, 2010).

There are a several advantages with the gaseous fuel from gasification of biomass compared to a fossil fuel; lower emissions of toxic by-products to the atmosphere and
efficient usage of solid by-products in form of char coal. The fact that the greenhouse
gas comes from a renewable source is another advantage. Most common is to use the
char coal direct in the gasification process. Char coal could be extracted technically,
but is normally not economically feasible.

Depending on how the gases will be treated, for example cleaning, after the
gasification different heating values can be obtained on the gas in a range from 3.4 to
over 35 MJ/m$^3$. 35 MJ/m$^3$ is similar to natural gas and the gas with this high values
are also named Substitute natural gas or Synthetic natural gas (SNG). This gas is
suitable for replacing or as compliment to natural gas and is already in use (Brown,
2003), (Zhang, Xu, & Champagne, 2010).

3.2.3 **Liquefaction**

Another thermochemical conversion method is liquefaction. There are two different
types of liquefaction techniques, direct and indirect, both technologies produce as
much liquids as possible and the residue is mainly tars. (Brown, 2003).

3.2.3.1 **Direct liquefaction**

Direct liquefaction takes place in an aqueous phase with a temperature around 350°C
and a pressure around 203bar. The residence time is long, around one hour (Brown,
2003). One benefit of this technology, also called Hydrothermal liquefaction (HTL),
is that the technology can handle waste streams from other processes which often are
in aqueous solutions (Dermibas, 2009). The technique produces, as from fast
pyrolysis, a lot of different liquid products such as alcohols, phenols and other
hydrocarbons. During the process the oxygen content of the organic material is
reduced richly to carbon dioxide (Dermibas, 2009). The research is not as developed
as for pyrolysis and gasification but there is some interesting results and heating
values around 36 MJ/kg is obtained together with a thermal efficiency of 80-90%
(Brown, 2003).

3.2.3.2 **Indirect liquefaction**

This is a technology where low-value organic materials produce a liquid with a gas
mixture of syngas (H$_2$ and CO) with a high heating value. The liquid will afterwards
continue to a catalytic or biological synthesis to produce ethanol or other chemical
products or compounds. The technique is developed for the process of coal into
methanol or synthetic fuel but can also work with biofuels. There are two ways of
production: the traditional process with moderate-temperature, high pressure and a
catalytic chemical reaction and a more modern way is to use microorganisms growing
on coal atoms to convert syngas to alcohols or acids (Brown, 2003).

3.3 **Biochemical processes**

In these processes the temperature is lower, down to 30°C, and the reaction rate is
also lower than for the thermochemical processes. The two most common types are
anaerobic digestion and fermentation (Cherubini, 2010).

3.3.1 **Anaerobic digestion**

This technique decomposes mainly organic wastes of different types into biogas, a
combination of mainly methane and carbon dioxide. The process takes place in an
oxygen free environment with help from bacteria. The process is in different steps and
in every step a special bacteria have a special task to break down the organic
compounds. This process is good for wet biomass and examples are manure, organic industrial waste sludge’s and organic domestic wastes. This is an old technology in the food and beverage industry but is increasing also in other branches to take care of waste and to produce a useful product, such as fuel. If the produced fuel is used directly in electricity production this process has a low efficiency. Biogas needs upgrading to give the process similar efficiency as with natural gas. The residue is often wet slurry with different energy and nutrient content (Faaij, 2006), (Brown, 2003).

### 3.3.2 Fermentation

Fermentation is a biochemical process which uses microorganisms and/or enzymes to create reactions. The reactions take place in an aqueous solution with the final products in modest concentration. One of the most used substrates is glucose but other examples are glycerol, pentoses and other hydrocarbons (Clements & Van Dyne, 2006). Fermentation is used for ethanol production from sugar, corn or other starch rich products, this is called 1st generation of fermentation. Besides the ethanol different numbers of other chemicals can be produced from fermentation. Most of them are produced in a small amount as specialty chemicals in example aliphatic acid and lactic acid (Brown, 2003). The fermentation process is chemically described as:

\[ C_6H_{12}O_6 \rightarrow 2C_2H_5OH + 2CO_2 \]

Fermentation 2nd generation uses cellulose based raw material and therefore gives different residues, these are mainly lignin based with good energy content.

### 3.4 Chemical processes

These processes conduct a change in the chemical structure of the molecule with a reaction to other substances. The most common processes are hydrolysis and transesterification. There is a group of other chemical reactions, which will not be discussed in this Master thesis due to their complexity. These chemical reactions are standard chemical process in the industry today, but with other raw materials, which also can work with biomass.

#### 3.4.1 Hydrolysis

Hydrolysis is a process where water splits a larger reactant molecule into two smaller molecules. The development of the technology started in the beginning of the 19th century when Napoleon started an economic continental blockade which stopped the import of sugar to Europe. Some researchers, especially in Germany, started different projects and Kirchhoff found that potato starch converted into sugar when it was cooked in dilute acid. In 1835 the Swedish scientist Berzelius found out that an enzyme with a catalyst could speed up the process and this developed the sugar production from sugar beets (Kamm, Kamm, Gruber, & Kromus, 2006).

This technology is used in the production of sugar from sugar beets or in extraction of starch from a lignocellulosics feedstock for ethanol production. The operating temperature is around 150-200°C and the process time is between a couple of seconds up to some minutes. The process can be one of the important technologies to extract starch from cellulosic feedstock for ethanol production (Brown, 2003).
3.4.1.1 Dilute-Acid Conversion

Hydrolysis with acid can be done by different acids, for example sulphuric and hydrochloric acids. But due to economic reasons the most common acid is sulphuric. The temperature of the steam is very important for the process and a low temperature favour xylose and a high will favour more furfural and tars. Xylose comes from xylan which is a complex polysaccharide in plant cell walls. There is an option for lower temperatures but in that case the concentration of the acid must be higher and a good chemical recycling system must be used to reach economical feasibility. Most of the technical development was done around the Second World War to secure some necessary production of sugar. An important factor here was to use a cellulosic feedstock, which was large in both USA and Germany. After the war some pilot plants have been built, none of these was commercialised because of the relatively low petroleum price the last decades. It was more economic feasible to use the “cheap” petroleum feedstock than develop a new and more expensive technology (Katzen & Schell, 2006).

3.4.1.2 Enzymatic Hydrolysis

Also the enzyme process started to develop after the Second World War. Research was done especially during the late seventies and early eighties. Up to the nineties some pilot and demonstration plants was built, but still no commercial plants. The reason for this is the same as for acid plants; it was cheaper with petroleum based products (Katzen & Schell, 2006).

Since a couple of year more research is performed and some projects are started and there is now a working technology. This has been possible due to the fact that the oil fields are smaller and the oil price is increasing but also because of the environmental debate. The technology is commercialised for sugar canes and sugar production but not for cellulosic feedstock at the moment (Katzen & Schell, 2006).

The enzymatic hydrolysis converts cellulose with a higher efficiency and has been tested in bench-scale and pilot scale. There is two ways to do the enzyme process either in separated reactors called Separate Hydrolysis and Fermentation (SHF) which has the problem that glucose formed in the hydrolysis process reduces the enzyme performance. The other possibility is to do the hydrolysis and fermentation in the same reactor and here the glucose can be consumed directly by yeast. This process is named Simultaneous Saccharification and Fermentation (SSF). Drawbacks are a compromise between efficient hydrolysis and efficient fermentation due to process conditions. Another drawback is that some of the yeast is removed together with the lignin and can therefore not be recycled (Eriksson, 2009). Even the hydrolysis residues are mainly lignin based, such as AHR and SSFR.

3.4.2 Transesterification

Transesterification is a process where triglyceride reacts with alcohols like methanol or ethanol to produce methyl ester or ethyl ester (also known as FAME) and the residue is glycerol. Triglyceride is extracted from the seed from different plants (soybean, sunflower or rape). If the seed have oil content over 20% mechanical pressing can be used and the seeds are crushed. If the seeds have a lower content a solvent extraction can be necessary. The extracted triglycerides, known as “fats and oils”, are esters of glycerol and fatty acids. The fatty acids have an even number of carbon atoms and the
degree of saturation can vary a lot. Fats are often solids in room temperature and have a high percentage of saturated acids. Oils have on the other side a high percentage of unsaturated acids and are liquid in room temperature.

Triglycerides from plants are often oils and therefore they have a high number of unsaturated fatty acids. The oils can be extracted from many different plants but today around 50% of the world production comes from soya beans which have a high yield in comparison. There are also future possibilities with a tree called Chinese tallow tree that is growing in the south of United States. The estimated potential of this plant is close to ten times higher than for soybeans (Brown, 2003)

The transesterfication process is a simple process between the triglyceride and methanol. The reaction takes place at 60°C if sodium hydroxide is used as catalyst or at room temperature with potassium hydroxide as catalyst. But in the last case the reaction time is longer. The two products (glycerol and RME) will be in two separate phases so the RME can be decanted of. The glycerol layer also contains an excess of methanol and the catalyst. The methanol is separated for reuse with distillation and the catalyst can be separated due to the use of the glycerol. Sometimes the catalyst need to be naturalised and for that purpose a phosphoric acid can be used in that case also another by-product is formed. This by product is a phosphorus salt which can be separated and use as fertilizer, the produced volume is very small. At the end the RME is washed to remove the last amount of methanol and after that the RME is vacuum dried (Batchelor, Booth, & Walker, 1995).

3.5 Cellulose separation technologies

The cellulose separation technology is also a chemical separation technology, but is here presented separated due to many different methods. The lignin is separated in the pulp process due to the fact that the lignin give bad properties connected to the colour of the paper. To avoid this, the lignin is separated to some level and the pulp is bleached to get a perfect paper. The separated lignin is used for internal heat and power services or extracted if it’s economical feasible to do that. The separation from the today’s most widely used pulp system, the Kraft mill, can recover lignin with a very high yield by acidification and filtration (i.e. LignoBoost) (Brunow, 2006). Of course there is also isolation of lignin from other types of production with lignocelluloses biomass. These two categories can be described based on the lignin’s sulphur content and in some way the sulphur content also describes what the lignin can be used for (Borges da Silvaa, et al., 2009), to see this complex system see Figure 6:

- Sulphur free lignin based on biomass conversion technologies for biofuel production.
- Sulphur containing lignin based on production from kraft and sulphite pulp production.

It is important to note that the sulphur containing processes also can be used for the production of biofuel i.e. ethanol.

Today the paper mills use the black liquor in the recovery boiler to get electricity and steam but also in the recycle of cooking chemicals. There is an excess of lignin in the black liquor which can be extracted for other services. This is also good for the paper mills where the recovery boiler often is the bottleneck for increasing pulp production. If some lignin is extracted more black liquor can pass through the recovery boiler and more pulp can be produced (Wallmo, 2008). After the separation of lignin in the pulp production the product called lignin consist of both organic and inorganic materials.
The organic materials consist of around 50% sugars which give this product a potential as animal feed additive (Pye, 2006).

3.5.1 Lignin Precipitation System (LPS)

The pulp mill of the soda model do not use sulphur in the chemical processes which will give a possibility to separate sulphur free lignin. Instead this lignin has special thermal and solubility properties, in combination with the sulphur free this give lignin for high value applications. The separation process starts with filtration of the black liquor to separate contaminated pulp fibres. After that will the filtrated liquor be acidificated to create lignin slurry. Next step will be to condition the slurry in a maturation step and then filter to remove the created lignin solids. The last step will be to wash and dry the filter cake to a high purity powder with only around 5% moisture. The rest of the black liquor which will not be returned to the mill will rather be used in an anaerobic digestion or by wet oxidation. The LPS technology is quite new and the first mill to use the technology was a French mill in Thonon and there are also building projects in India (Pye, 2006).

3.5.2 LignoBoost

The LignoBoost process is simple in just a few steps. The first step after the black liquor leave the evaporator system is precipitation vessel. In this vessel carbon dioxide is used to reduce the pH of the black liquor to around 9-10. Around 75% of the lignin is precipitated as a sodium salt (Borges da Silvaa, et al., 2009).

![Diagram of biorefinery site for kraft lignin and vanillin](Borges da Silvaa, et al., 2009).

STFI – Packforsk and LignoBoost AB have a demonstration plant in Bäckhammar, Sweden. This plant uses between 260 and 280 kgCO₂/tonne lignin (Tomani, 2008). In the next step this lignin will be suspended in water and the pH will drop more, the drop will be a result of acidification with sulphuric acid (H₂SO₄) (Borges da Silvaa, et
al., 2009). There are possibilities to use sulphuric acid in both steps, but it can damage the sulphur balance in the paper mill. The CO₂ process is more complicated but better in an economic perspective (Wallmo, 2008). The final step is a filtration process and washing where the lignin is taken out in different steps before the black liquor returns to the recovery boiler at the paper mill. The separated lignin can then be used for energy purposes or in production of different chemical products such as vanillin, see chapter 5.2.

The demonstration plant in Bäckhammar has in cooperation with Chalmers University of Technology tested two different lignin fuels with good results. One fuel as powder and one as filter cake, properties of lignin as fuel are presented in Table 4, in chapter 5.2.

3.5.3 Near-Neutral Pre-Extraction

There are also other technologies based on a paper mill, one of them is the Near-Neutral Pre-extraction. This technology is based on extraction of hemicelluloses with focus on production of acetic acid and raw material for fermentation to ethanol. The system is similar to the LignoBoost and the pH is decreased with H₂SO₄ after the evaporation. In the described model from van Heiningen et al will the lignin return within the plant. The advantages are that the process off loaded the recovery cycle (like LignoBoost), the quality and quantity of the pulp is also unchanged (van Heiningen, Mao, Genco, Yoon, Zou, & Pendse).

3.5.4 Organosolv

Organosolv is a very old technique and have been used for over hundred years to separate wood in different components. In the beginning it was to study lignin and carbohydrates and to do this separation organic chemicals (solvents) where used. Later in the history research was done to find out if the technology was possible to use to replace short comings in the traditional pulp industry. Just a few of these technologies were good enough to compete with kraft pulping. These are often regular chemical processes with addition of organic solvents. Depending on which solvents used the process name can change i.e. Alcell (ethanol), Acetosolv (acetic acid) or Formacell (formic acid). The Alcell are the most common and the aqueous pulping solution consists of ethanol, here pulping liquor will be cooked in relatively high temperatures and pressure (around 195°C and 28 bar) (Pye, 2006). The organosolv follow the mechanism for regular chemical pulping, but is more selective due to the alkali-alcohol reactions that occur to the organic solvents. The result of this is that organosolv is more selective to lignin and therefore more lignin can be separated. Organosolv have the following advantages due to regular chemical pulping (Kadia & Dai, 2006):

- Higher separation factor of lignin
- Milder conditions during production; free of strong acids or alkalis and low temperatures
- Reduction or elimination of sulphur containing compound emission
- A simplified recovery chemical recovery system
- Recovery of lignocelluloses chemical by-products

There is one major problem with the organosolv technique; the quality of the pulp is not god enough. The pulp can be used for some special paper application but the most promising for this technology now is the development around biofuel production.
residues (lignin) from the organosolv technique are clean and that is a reason way it can be interesting for the biofuel and chemical market (Kadia & Dai, 2006).

3.6 Development status of technologies

In the following table the development status of the different involved technologies are summarized. As can be seen many of the technologies are existing (commercialised) or on their way towards market introduction. It is important to mention that some of the technologies are commercialised as technology but are sometimes not commercialised as biorefinery technology or as a technology used for production of green chemicals. Gasification and liquefaction works with coal, but both technologies need further development before use with biomass. According to gasification the technology is ready but only demonstration plants up 10 MW_{th} is existing.

*Table 1: Development status of biorefinery technologies.*

<table>
<thead>
<tr>
<th>Technology</th>
<th>Status</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaerobic Digestion</td>
<td>Existing</td>
<td>World wide</td>
</tr>
<tr>
<td>Barking</td>
<td>Existing</td>
<td>World wide</td>
</tr>
<tr>
<td>Fermentation (1\textsuperscript{st})</td>
<td>Existing</td>
<td>World wide</td>
</tr>
<tr>
<td>Fermentation (2\textsuperscript{nd})</td>
<td>Demonstration/Existing</td>
<td>Kalundborg, Abengoa</td>
</tr>
<tr>
<td>Gasification</td>
<td>Demonstration/Existing</td>
<td>Güssing, Chalmers</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>Existing</td>
<td>World wide</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>Pilot plant</td>
<td>Bäckhammar</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>Pilot plant</td>
<td>USA/Europe</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>Existing</td>
<td>USA/Europe</td>
</tr>
<tr>
<td>LPS</td>
<td>Existing</td>
<td>France/India</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>Demonstration/Existing</td>
<td>USA</td>
</tr>
<tr>
<td>Organosolv</td>
<td>Existing</td>
<td>World wide</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>Demonstration/Existing</td>
<td>USA/Europe</td>
</tr>
<tr>
<td>Transesterfication</td>
<td>Existing</td>
<td>World wide</td>
</tr>
</tbody>
</table>
4 Target products from biorefinery technologies

The different biorefinery technologies produces several main products, in this chapter the target products with large potential for future use are presented. These target products also gives interesting residues during production.

4.1 Syngas

Syngas and biogas is used for different purposes and have different compositions. Carbon dioxide, carbon monoxide and hydrogen is the mix building up syngas and this synthetic gas is produced though gasification or pyrolysis. Other outputs from these technologies are bio-oil and char coal. Both the fraction of bio-oil and char coal is very depend on temperature in the process and char coal is most often used for direct combustion within the own biorefinery plant.

Syngas energy density is 50% of natural gas and it can be used as combustion fuel or a feedstock for various chemical products, but is mostly used for production of methanol. Due to the short carbon chains it is easy to rearrange syngas to desired carbon chains in chemicals. Syngas contains a small part of nitrogen that has to be separated to get a useful product. This is a hard process due to the similar boiling points of nitrogen and carbon monoxide (Biofuel org. U.K., 2011).

4.2 Bio-Oil

Bio-oil from pyrolysis is a dark brown fluid with a viscosity close to medium fuel oil and will not mix with any other hydrocarbon liquid. A problem with the oil is the temperature sensitivity; due to this the bio-oil cannot be distilled in a further chemical process without being other products. Bio-oil can act as raw material in the chemical industry and fulfil the same requirements in many cases as the traditional crude oil (Bridgewater & Peacock, 2000). The higher heating value for bio-oil is around 17 MJ/kg in comparison with crude oil 42-44 MJ/kg.

4.3 Biogas

With a decomposition of organic materials (including wastes) with bacteria in anaerobic digestion (see chapter 3.3.1) gaseous fuel will be produced. The bacteria break down the organic material into simpler organic compounds. Biogas mainly consists of methane and carbon dioxide but also a number of trace gases (Brown, 2003). To achieve a good gaseous fuel as clean methane as possible is necessary. This clean stream is reached by separating carbon dioxide with different methods in a step called upgrading. If sulphur needs to be separated, this can be done with a water scrubber.

If biogas is upgraded and compressed it can be used as transportation fuel and it can through upgrading reach natural gas values. Biogas upgraded to 97% of methane has a caloric value of 9.67 kWh/m$^3$, slightly less than natural gas that has 11.0 kWh/m$^3$, although this upgrading is energy intensive and can be costly as well (Erdmann & Zweifel, 2008)

4.4 Ethanol

Ethyl alcohol or ethanol is an organic compound with the molecular formula C$_2$H$_5$OH. Ethanol has quite broad application possibilities within chemical industry as a solvent in the synthesis of other organic compounds or as transportation fuel. The most
common production model is fermentation (see chapter 3.3.2) which gives a dilute aqueous solution. This solution needs afterwards a fraction distillation to obtain an acceptable quality, after the distillation even dehydration can be necessary. Final product is a colour less flammable liquid (Encyclopædia Britannica, 2011). Production can be done either with a starch based feedstock (the first generation) or with a lignocelluloses feedstock (second generation). If the lignocelluloses feedstock is used than some pretreatment is necessary either with hydrolysis or with different lignin separation methods (Faaij, 2006). The second generation only has energy efficiency around 35%, so to be competitive the residue must be used. These residues are often lignin rich and are therefore energy rich (Eriksson, 2009).

4.5 RME

FAME is formed by a reaction called transesterification between triglyceride and methanol. Normally in Europe the fatty acids are from Rapeseed and therefore the most common product name is Rape methyl ester (RME). There are different ways to sell this fuel either it can be sold as B100 which is 100% of RME and then 100% renewable fuel. Another way to sell the RME is to blend it with diesel and sell this blended product. The positive aspect with RME is that it can directly or with very small modifications replace diesel in vehicles. In Europe, especially in Germany and France, large amount of this fuel is used. The problem with RME is that it is still a quite expensive fuel and needs subsidies to be competitive. In Europe this is solved by a subside system in two parts. The first part of the system is according to rape production includes subsidies to farmers that produce these plants for example instead of crops for food production. The second part is in the tax system and tax benefits are given to producers and users of RME (Faaij, 2006).

4.6 Paper

The first product is from the pulp and paper mill is pulp and with further process paper can be produced. Pulp is not an end consumer product but there are cases with mills that only produce pulp and sell to paper producing sites. Paper production is a traditional industry with a large energy demand connected to different steps of drying and pressing. This because the pulp is a wet slurry when it leaves the mill. The paper making process also includes the bleaching which is necessary for a white paper with good contrast. The chemicals used for this is chlorine or in more modern case oxygen. Paper production is an important industry in some of the EU-27 countries, especially in Finland and Sweden. The two biggest categories of produced paper are printing/writing and packaging, together they stand for around 60% of the worldwide production. The paper production worldwide increase every year and there is a clear connection of paper use and income in a country, higher income means more paper (Brännvall, 2009).
5 Valuable residues

Output from biorefinery technologies are target products, by-products and valuable residues. In this chapter the solid residues coming from different biorefinery technologies in chapter 3 is presented.

5.1 Bark

The bark proportion of biomass feedstock often plays an important role in energy conversion properties, this due to the fact that removal of bark sometimes is not feasible and in other cases bark is separated and sold on a separate market (Kenney, Sennerby-Forsse, & Layton, 1989). Biomass is usually classified by parameters energy purposes, humidity, density, heating value and ash content (Werther Guidi, 2008). Compared with wood, bark has higher moisture content, higher ash and alkali metal content which may cause corrosion in boilers and furnaces (Werther Guidi, 2008). Compounds like sulphur and chloride in the ash are corrosive and can reduce the oxide layer in boilers and furnace, these compounds can also reduce the melting point of the ash. Although in woody biomass there are low concentrations of sulphur and chlorine (Lehtikangas, 2000). Since bark acts as a protection and nutrients barrier for the wood it contains higher concentrations of N, P, K, Mg, Ca, Cd, Pb, Co and Zn, see Table 2 (Adler, Verwijst, & Aronsson, 2005). Because of these minerals the ash content of the whole tree is largely depending on the proportion of bark (Werther Guidi, 2008).

Bark also has a higher content of lignin and a lower percentage of cellulose, see Figure 7. This induce that biomass with high bark content is a valuable energy source due to the high content of lignin, but to get maximum energy output this type of biomass has to be stored properly and bark properties has to be taken into account (Adler, Verwijst, & Aronsson, 2005). Bark is more sensitive to temperature and pressure than wood because of the relatively high content of lignin and alkali metals. The proportions of lignin in bark products increase during storage time (Lehtikangas, 2000).

For a willow stand assessed in 2003 the bark proportion of biomass standing in the field where on average 19%, the small twigs had around 56% bark content and 5 year old plants had lowest with 15% bark content. Bark content in willow where seen to be relatively constant when plant had grown a diameter of 20 mm at a height of 55 cm. Due to the special properties of bark, the bark content in energy crops have direct impact on the quality of fuel and ash content in bark is seen as high (4.1%-4.8%) (Adler, Verwijst, & Aronsson, 2005). Higher ash content in biomass depends on alkali metals and lowers the heating value (Lehtikangas, 2000).
Table 2: Properties for bark pellets (Berglin, Tomani, Salman, Herstad Svärd, & Åmand, 2008)

<table>
<thead>
<tr>
<th>Unit</th>
<th>Bark pellets</th>
</tr>
</thead>
<tbody>
<tr>
<td>C %wt of dry solids</td>
<td>51.90%</td>
</tr>
<tr>
<td>H %wt of dry solids</td>
<td>6.10%</td>
</tr>
<tr>
<td>N %wt of dry solids</td>
<td>0.40%</td>
</tr>
<tr>
<td>S %wt of dry solids</td>
<td>0.04%</td>
</tr>
<tr>
<td>O %wt of dry solids</td>
<td>37.90%</td>
</tr>
<tr>
<td>Cl %wt of dry solids</td>
<td>0.02%</td>
</tr>
<tr>
<td>Ca mg/kg dry solids</td>
<td>9324</td>
</tr>
<tr>
<td>K mg/kg dry solids</td>
<td>2138</td>
</tr>
<tr>
<td>Na mg/kg dry solids</td>
<td>310</td>
</tr>
<tr>
<td>Ash %wt of dry solids</td>
<td>3.60%</td>
</tr>
<tr>
<td>LHV MJ/kg dry wt</td>
<td>17.7</td>
</tr>
<tr>
<td>HHV MJ/kg dry wt</td>
<td>21</td>
</tr>
<tr>
<td>Moisture content % dry wt</td>
<td>10.30%</td>
</tr>
</tbody>
</table>

There has been found that bark from poplar crops contain important pharmaceuticals (Barton, 1984). This bark can also be used for acid treatment of leather, and in this way replace today use of hazard acid liquids (Grundmann, 2011). This could maybe lead to a higher value on this type of bark.

From the pulp and paper industry of today a lot of wooden bark is produced. One modern mill can separate and sell 242 GWh/yr. The electricity production in EU-27 was in 2008 3.4 million GWh (Eurostat, 2009), (Jönsson & Berntsson, 2011). Considering that there are 209 pulp mills or combined pulp and paper mills in EU ETS, a significant amount of bark is available on the market (EcoFys, 2009).

5.2 Char coal

Char coal is formed then wood is slowly heated in a chamber with limited and controlled amount of air, it can either be extracted as a residue for further combustion process at another plant or it can be combusted directly at the own plant. If it’s economically feasible to extract char coal for co-combustion with coal is uncertain, this depends on for instance transportation distance and cooperation with owner of plant. Char coal has several advantages among the solid residues coming from biorefinery technologies. One of them is the small amount of minerals and alkali content (Brown, 2003). A chemical biorefinery plant has shown that char coal from biomass is a promising fuel with heating values around 25.6 MJ/kg and ash content of 15%. Combustion of this char coal can yield more energy than combustion of the entire originally feedstock (50% moisture content) at pilot plant (Hayes, Fitzpatrick, Hayes, & Ross, 2006). The pilot plant discussed by Hayes et al. estimated that the energy potential coming from char coal is greater than the energy needed at the plant when scale of operation is equal or bigger than 270 dry tonnes of feedstock per day. This surplus could be sold as electricity on the grid or the char coal could be sold for co-combustion with coal (Hayes, Fitzpatrick, Hayes, & Ross, 2006).

Char coal from low and medium temperature processes with biomass or waste as feedstock material is one of the most reactive carbon materials, more reactive than char from coal (Di Blasi, 2008). This is due to a highly disordered structure together
with the porosity (Henrich, Bürkle, Meza-Renken, & Rumpel, 1998). If the temperature in process is increased the yield of char coal from biomass decreases, so if high yield of biomass char should be obtained the pyrolysis process should work with low temperature and low heating rate. Increasing particle size of biomass and higher lignin content of biomass also affects the yield of char coal in a positive way (Demirbas, 2004). Figure 8 shows the trend of char coal yields, figures for drawing this diagram are average values from Di Blasi. Here fast pyrolysis is performed in a fluidized-bed pyrolysis and slow pyrolysis refers to pyrolysis of cylinders 40 mm in diameter (laboratory scale), both fast and slow pyrolysis is performed with softwood (Di Blasi, 2008).

Due to that higher pyrolysis and gasification temperatures give more valuable products from biomass it is unlikely to believe that pyrolysis at low temperature will be preferable at commercial scale. In most processes studied the char coal is directly combusted within the own plant both due to simplifications and for char coals high heating value. During literature study no valuable products coming from char coal used as feedstock have been obtained, only char coal own value as fuel is discussed in literature.

Char coal is a very important energy source in developing countries, and is often used in households, but also for larger applications. The briquetting of char coal improves the efficient of the energy used and therefore also the efficiency of the biomass based energy. As example a wood based char coal contains 80% carbon, 1-3% ash and 12-15% volatile gases (Demirbas, 2009).

5.3 AHR and SSFR

Ethanol can be produced in several different ways and before fermentation dilute-acid hydrolysis or enzymatic hydrolysis can be used to separate out cellulose. These two processes are described in Figure 9 and Figure 10, here it can be seen that there are some possibilities to take out solid lignin rich residues which can be pelletized and used as fuel (Wingren, Galbe, & Zacchi, 2008).
In the dilute-acid hydrolysis residue (AHR) is taken out by filtration before the distillation. The data for the residue will be presented in Table 3. The hydrolysis process leads in both cases to a reduction in the alkali content which is good for the boiler (Eriksson, 2009).

![Figure 9: Dilute-acid hydrolysis for ethanol production (Eriksson, 2009).](image)

In the SSF process the final ethanol concentration will be low, only around 4wt%. To increase this it is necessary to concentrate the ethanol stream to a higher level. The most common operator to use is distillation. After the distillation a solid phase residue named SSFR is separated out with filtration. The liquid phase/slurry can be further processed in either an evaporator system or in an anaerobic digester. The evaporator system will use a lot of energy but will give a condensate that can be recycled within the system to save water. The digester on the other hand gives methane/biogas that can be used or sold. The both systems also give residues back to the pellet making process, but with different properties. The solid stream from the filtration and the concentrated steam from the evaporator system will be mixed together and dried to 85% DM. The drying process will most likely use steam on different steam levels (around 12 and 20bar). Dried solids can be used as fuel to create the necessary steam and the excess can be sold. Both processes are operated as batch process and the total residence time for the digester will be around 25 days and for the evaporator system 84h and the fermentation takes around 72h (Wingren, Galbe, & Zacchi, 2008).

![Figure 10: Simultaneous Saccharification and Fermentation enzyme hydrolysis process for ethanol production (Eriksson, 2009).](image)

The residues from the anaerobic digester case are good and consist of 80% lignin which also gives a good heating value. The pellets from the evaporator consist of only 55% of lignin. This is due to that the concentrated syrup from the evaporators contains of other substances as glycerol, ash and inorganic compounds. This will decrease the
heating value of the pellets. The lower heating value in combination with the higher ash content will do the evaporator residue, DR, to a lower quality fuel. There are also some risks for higher sulphur content in the evaporator syrup (Wingren, Galbe, & Zacchi, 2008).

Table 3: Properties of AHR and SSFR (Eriksson, 2009), (Wingren, Galbe, & Zacchi, 2008).

<table>
<thead>
<tr>
<th>Unit</th>
<th>AHR</th>
<th>SSFR</th>
</tr>
</thead>
<tbody>
<tr>
<td>C %wt of dry solids</td>
<td>59.10%</td>
<td>46.80%</td>
</tr>
<tr>
<td>H %wt of dry solids</td>
<td>5.90%</td>
<td>6.20%</td>
</tr>
<tr>
<td>N %wt of dry solids</td>
<td>0.15%</td>
<td>5.82%</td>
</tr>
<tr>
<td>S %wt of dry solids</td>
<td>0.10%</td>
<td>0.30%</td>
</tr>
<tr>
<td>O %wt of dry solids</td>
<td>34.60%</td>
<td>33.10%</td>
</tr>
<tr>
<td>Cl %wt of dry solids</td>
<td>&lt;0.01%</td>
<td>0.03%</td>
</tr>
<tr>
<td>Ca %wt of dry solids</td>
<td>0.00074%</td>
<td>0.72%</td>
</tr>
<tr>
<td>K %wt of dry solids</td>
<td>0.00044%</td>
<td>1.32%</td>
</tr>
<tr>
<td>Na %wt of dry solids</td>
<td>0.00026%</td>
<td>0.013%</td>
</tr>
<tr>
<td>Ash %wt of dry solids</td>
<td>0.50%</td>
<td>0.19%</td>
</tr>
<tr>
<td>LHV MJ/kg DS</td>
<td>20.9</td>
<td>19.9</td>
</tr>
<tr>
<td>HHV MJ/kg DS</td>
<td>22.7</td>
<td>21.5</td>
</tr>
<tr>
<td>Moisture content %</td>
<td>-</td>
<td>15.0%</td>
</tr>
</tbody>
</table>

5.3.1 Concentrated liquid distillation residue (DR)

DR is a wet residue from the second generation ethanol production and the composition of this residue is not completely known but part of it is similar to glucose. The heating value is under 20 MJ/kg and it can be dried to pellets. There is some ongoing research about combustion of DR, but there is a major problem with alkaline. This alkaline comes from products which are used to neutralize the stream after pretreatment process and will also give a higher ash content of the residue. There are possibilities to use the DR in digester for biogas production or as fertilizer (Eriksson, 2009).

5.4 Glycerol

Glycerol or glycerine (C₃H₈O₃) is a residue from the production of rape methyl ester (RME) or FAME (from other plant-oils) a fuel substitute for diesel engines. The oilseed can be used in three ways in the engine; pure rape oil, blended with fossil diesel and as RME. The fluid properties of pure rape oil make it unsuitable, because of large droplets which give incomplete combustion. Both the blended version and RME can be used in a diesel engine with no or very small modifications (Batchelor, Booth, & Walker, 1995).

Currently the glycerol market is nearly saturated. The market consists of cosmetics, chemical, food and beverage industries, after the entry of the RME glycerol price has decreased. Due to the low price the market at the moment tries to find out which valuable products can be developed from glycerol to get a higher value out of this residue (Bohon, Metzger, Linak, King, & Roberts, 2011). The price policy of the glycerol products is complex due to different purity of the products. The traditional industries use glycerol at purity of around 98%, from transesterification the purity is around 80-88% (Rahmat, Abdullah, & Mohamed, 2010). There are other examples;
one of the Swedish RME producers, Perstorp AB, in Stenungsund produce glycerol with a purity of at least 97% and sell this product to other chemical companies (Perstorp AB, 2008).

The good thing for the environment with the RME glycerol production is that this can replace fossil based glycerol even if it’s not used as fuel. Further more important factor for the environmental friendliness is how the methanol is produced. The methanol can be produced either from natural gas or from gasification of biomass. If the methanol comes from the gasification the total process can be more sustainable (Eriksson, 2009).

The technical problem with glycerol is that the compound has a melting point around 17.8°C and therefore is syrup in room temperature (CHEMnetBASE, 2011). So the glycerol is more similar to oil but still possible to use as a fuel.

5.5 Lignin

Lignin (C₉H₁₀O₅(OCH₃)ₙ) is the natural phenolic glue that holds together cellulose fibers and act as a natural protection against biological degradation of cellulose and hemicelluloses (Lui, 2010). It is a three-dimensional macromolecule with a carbon to oxygen atom ratio higher than 2:1. Hemicelluloses and cellulose have approximately the carbon ratio 1:1. This is the reason why lignin is a more energy dense substance than sugar (Lui, 2010). Cellulose and hemicelluloses can be hydrolyzed to sugars and then fermented into high value products such as ethanol. Lignin cannot be fermented but it may be useful for chemical extraction and energy generation (Cherubini, 2010). First in the 1940’s engineers learned how to use lignin and hemicelluloses for energy production, prior lignin was a waste product with severe environmental problems (Lui, 2010). There has been shown that there is a significant difference between the hardwood and softwood lignin structure which gives different physical and chemical properties of these two lignin types (Lui, 2010). Lignin molecules weight distributions do to certain extend affect the reactivity and physicochemical properties of lignin (Doherty, Mounsavioun, & Fellows, 2010). There are large research projects going on to find out and evaluate what chemicals that can be extracted from the lignin. But at the moment this research is on a basic level in many cases. So besides today’s use for production of vanillin it is hard to say how the future uses will be (Lui, 2010). To see properties of lignin see Table 4.

There is an ongoing process to find a possible ways to use the isolated lignin in an efficient way connected to both resource and environmental questions. This is hard due to the complex molecule structure of the lignin. Already today there are some commercial products for the lignin i.e. vanillin and dimethylsulfoxide. Product classification of different lignin products (Brunow, 2006):

- Combustion products
- Utilization of surface products
- Condensation of lignin so that becomes an integral part of a product

So there is competition between different usages of the lignin. This competition has both an environmental and economical aspect. Also the properties of the lignin residue will be important and the value can be depending on the contamination from the separation processes (Pye, 2006). This future processes will mainly be based on the focus to get fermentable sugars from cellulose and hemicelluloses. An important
factor to keep in mind is that the lignin in many cases will be given in wet slurry. This slurry will afterwards pass through a drying process and later solid lignin or pellets are produced.

Table 4: Properties of lignin (Berglin, Tomani, Salman, Herstad Svärd, & Åmand, 2008).

<table>
<thead>
<tr>
<th>Unit</th>
<th>Lignin</th>
</tr>
</thead>
<tbody>
<tr>
<td>C %wt of dry solids</td>
<td>62.70%</td>
</tr>
<tr>
<td>H %wt of dry solids</td>
<td>5.70%</td>
</tr>
<tr>
<td>N %wt of dry solids</td>
<td>0.15%</td>
</tr>
<tr>
<td>S %wt of dry solids</td>
<td>0.18%</td>
</tr>
<tr>
<td>O %wt of dry solids</td>
<td>27.60%</td>
</tr>
<tr>
<td>Cl %wt of dry solids</td>
<td>&lt; 0.01 %</td>
</tr>
<tr>
<td>Ca mg/kg dry solids</td>
<td>1085</td>
</tr>
<tr>
<td>K mg/kg dry solids</td>
<td>543</td>
</tr>
<tr>
<td>Na mg/kg dry solids</td>
<td>1666</td>
</tr>
<tr>
<td>Ash %wt of dry solids</td>
<td>1.4%</td>
</tr>
<tr>
<td>LHV MJ/kg dry wt</td>
<td>17</td>
</tr>
<tr>
<td>HHV MJ/kg dry wt</td>
<td>26.3</td>
</tr>
<tr>
<td>Moisture content % dry wt</td>
<td>29.30%</td>
</tr>
</tbody>
</table>
6 Method

This chapter describes the different methods that are used in the Master thesis and explains how chapter 7 and 8 are performed. The research method used in this project is analytic research through classification and ranking of biorefinery technologies.

6.1 Market potential for target products

To see what market potential different target products can reach an analysis based on statistical data and scenario analysis has been encompassed. Here the fossil feedstock market and renewable resources market were analysed from different views such as, market share of today, production of today and possible future demand.

Data for today’s production of target products based on biomass, see chapter 4, where collected and this production was compared to production of similar products from fossil feedstock. After these two steps calculations have been done to see how much production has to increase in order to substitute products from fossil feedstock. Different fractions of substitutions up to 50% where used to calculate the market share of bio based products. This has been performed by calculating the amount bio based ethanol that is necessary to replace 10, 20, 30, 40 or 50% of today’s production of ethylene and how large increase in ethanol production this will give. The same type of calculations has been carried out for syngas and biogas.

Information about today’s production in EU-27 is collected from Eurostat 2009 production database and calculations where performed in Microsoft Office Excel. Calculations on ethanol were done on tonne bases, due that energy content in renewable ethanol is the same as for fossil ethanol. For syngas the energy content is lower than of fossil natural gas, here energy content was used as bases.

For RME, lignin and paper a theoretical market potential assessment where performed. This theoretical assessment is based on literature and market situation today. Calculations where here excluded because the constituent do have low or no interest for these markets. Instead a short description of the market situation is done.

6.2 Classification

Classification is used to systematically order a larger group of research objects and it works as a useful tool in decision making process. To be able to classify research objects one or several categories are chosen, each and every object should fit into one of these categories. In order to do so the process of classification is divided into two parts, one static for order of steps in value chain and one dynamic for type of biorefinery process (Welter, 2006). The static process is fixed and does not change throughout the classification work. The dynamic part aloud the classification system to expand there is necessary.

The categories for type of biorefinery process are chosen from Cherubini (2010) the categories for value chain are closely connected to objective and chapter 3, 4 and 5. The input in category “Technology/process” represents chapter 3, category “Target product” is based on chapter 4 and “Residues” column is based on chapter 5. The order for these categories represents biomass process way through a biorefinery

---

2 Value chain is describing the biomass path from gate to gate.
system. In total this classification has seven categories forming a 4x3 matrix, see Figure 11.

![Classification System](image)

To easy follow the steps in value chain the objects placed in each box will be connected with arrows. If there are any interconnections between different boxes cross section arrows show these connections. All seven categories can also be found in chapter 3, 4 and 5, with these texts as background the matrix is filled with information.

### 6.3 Weighted ranking

To get an overview over which biorefinery technologies that can be interesting to focus on in the future a weighted ranking is performed. 14 different technologies are ranked by seven different criteria’s, with a score from 1 to 14, where 14 is the best. In this ranking it is not possible that two technologies get the same score in each criteria even if they seem to be similar. This was done in order to have distinct differences between all technologies. So in total a score of 105 points ($\sum 1+2+3+\ldots+14$) will be given in each criteria. The maximum score for a technology is therefore 98 points (7 criteria x 14 maximum score). Each criteria will be weighted and is given a weighting factor based on the importance of the criteria connected to the objective of the thesis. This weighting process will be described more deeply in chapter 7.3 and the maximum weighted score is 26.6 points. The ranking is based mainly on the fact review in the report. In Appendix B a short text describes each technology in all criteria and these texts creates a judgement table. The way from the judgment table to the allocated score in the ranking table is based on Table 8 that describes which factors that are important in each specific criteria. The ranking is performed in Microsoft Office Excel.

### 6.4 Combination of technologies

In the section Combination of technologies the review of different biorefinery technologies and their properties together with literature of planned and existing plants are put together in possible plant solutions. These plants are suggested combination of different process steps to achieve interesting target products and at the same time valuable residues. Some of the processes options comes from Eriksson
(2009) but are after that modified. The suggested plants are described with process schemes to give the reader a clear view and explanation of symbols can be seen at page IX. Different inputs and outputs are marked in these schemes and can except from raw material, products and residues as well as other chemicals or heat. The focus for the scheme is based on interesting target products and how their system can be built up and at the same time give interesting residues. This chapter don’t include calculations of process integration which not are included in the objective. The technologies used in these chapters do not need temperatures that are higher than the excess temperatures from coal power plants.

6.5 C-cycle and energy balance

The aim here is to calculate the amount of carbon coming from crude oil that is used as feedstock, in EU-27 chemical industry. Calculated demand for carbon will be used to estimate how much farmland is needed to cultivate energy crops to cover the demand for carbon. To make the result more transparent an energy balance is also performed, here the energy coming from crude oil in chemical industry is calculated. Then calculations are performed to determine the amount of biomass corresponding to the energy coming from crude oil. For biomass and crude oil used in energy services such calculations should be based on exergy content rather than energy content, although for chemical industry this is of less importance. If energy content is used instead of exergy content, there is most likely an underestimation of demand for raw material.

The calculations of carbon content and energy balance will be done on a mix of energy crops, this master thesis will exclude wooden biomass and agricultural by-products as biomass raw material. The energy crop mix includes willow, hemp, reed canary grass and poplar, the term energy crop mix refers to these species. Miscanthus is excluded because no appropriate data has been found on specific properties.

To carry out the carbon cycle analysis calculations and data collections where divided into four steps:

1. Find data for crude oil consumption in EU-27 chemical industry, with pharmaceuticals included. Calculate the total amount of carbon in this amount of crude oil.
2. Find data and calculate the carbon content in a mix of energy crops cultivated in EU-27.
3. Calculate how much energy crops in kg that is needed to replace the total carbon coming from crude oil.
4. Calculate how much cultivated area that is needed to cultivate the amount of energy crops from point 3, compare these numbers with available farmland in EU-27.

The calculations on energy balance where performed in three stages:

1. Calculate the total energy coming from crude oil within chemical industry. Use the same crude oil consumption as in calculations for carbon cycle.
2. Calculate the mass of biomass in t\text{DM} corresponding to energy coming from crude oil.
3. Calculate how large cultivated area is needed to produce this amount of biomass. These numbers are then compared with available farmland in EU-27.

Calculations for both cases are performed in Microsoft Office Excel software. Before doing calculations unit analysis for each point above where performed, here beneath the unit analysis for the carbon cycle and energy balance is presented.
Equation 2: Unit analysis for calculations of total gC coming from crude oil.

\[
\frac{C \text{ content in crude oil}}{\text{Energy unit}} \times \frac{\text{Amount of crude oil used in chemical industry}}{\text{Energy content Oil equivalent}} \times \frac{\text{Conversion efficiency for crude oil to useful product}}{\text{Conversion efficiency for crude oil to useful product}} = \frac{gC}{MJ} \times \frac{M}{\text{toe}} \times \frac{MJ}{\text{toe}} \times \% = M \ gC
\]

By performing calculations due to Equation 2 the total amount of carbon coming from crude oil in chemical industry is obtained. The unit tonne of oil equivalent is often used to compare energy content in different fuels. The second step is to calculate the carbon content in energy crop mix.

Equation 3: Unit analysis for calculation of carbon content in energy crop mix.

\[
\frac{C \text{ content in energy crop mix mean value}}{\text{Energy content energy crop mix DM (mean value)}} \times \text{multiplied with 1000 to get g} = \frac{kg \ C}{kg_{DM}} \times 1000 = g \ C / MJ
\]

Here the mean net calorific value in energy crop mix and mean percentage of carbon content in dry matter material, DM, where used to calculate the carbon content per energy unit for biomass. Equation 2 and Equation 3 is needed to move forward and perform calculations on how much biomass the amount of carbon corresponds to.

Equation 4: Unit analysis for calculations of need for biomass to cover total amount of carbon from crude oil.

\[
\frac{\text{Amount of carbon coming from crude oil used in chemical sector}}{\text{Energy content in energy crop mix DM (mean value)}} \times \frac{C \text{ content in energy crop mix per energy unit}}{\text{Conversion efficiency of biomass to useful product}} = \frac{gC}{MJ} \times \frac{M}{kg_{DM}} \times \% = kg
\]

Equation 4 gives the total amount of biomass in dry weight needed to cover the necessity for carbon. In order to calculate how many kg of dry matter biomass that is needed, the energy content in energy crop mix is used.

Equation 5: Unit analysis for calculation of need for cultivated land.

\[
\frac{\text{Tonnes of dry weight material needed}}{\text{Yield energy crop mix mean value}} = \frac{t_{DM}}{\text{ha/year}} = \text{ha/year}
\]

The last equation is performed to obtain the size of the cultivated area that is required to produce the annual amount of carbon calculated from Equation 2. This figure is then compared with the total area of farmland available in EU-27.

Following three equations shows the unit analysis performed to calculate the energy balance.

Equation 6: Unit analysis for calculate total energy coming from crude oil within chemical industry.

\[
\frac{\text{Amount of crude oil used in chemical industry}}{\text{Energy content Oil equivalent}} \times \frac{\text{Conversion efficiency crude oil to useful product}}{\text{Conversion efficiency crude oil to useful product}} = \frac{t_{DM}}{\text{ha/year}} = \text{ha/year}
\]
\[
= M_{\text{toe}} \times \frac{GJ}{\text{toe}} \times \% = PJ
\]

Equation 6 is used to calculate the total energy coming from crude oil within chemical sector. Equation 2 and Equation 6 is differentiated by one factor, in Equation 6 the carbon content in crude oil is excluded. In calculations the same crude oil consumption as in calculations for carbon cycle is used.

**Equation 7: Unit analysis to calculate the amount for biomass in DM that is needed to cover energy demand.**

\[
\text{Amount of total energy coming from crude oil divided by 1000 to get tonne} / \text{Energy content in energy crops mean value of DM} = \frac{PJ}{Mj/kg} /1000 = t_{DM}
\]

After calculating how much energy that comes from crude oil, calculations are done to see how much biomass this amount of energy corresponds to. The figure from Equation 7 will probably differ from Equation 4 which shows the need for biomass to cover the amount of carbon coming from crude oil.

**Equation 8: Unit analysis to calculate the cultivated area needed to produce the necessary amount of biomass.**

\[
\text{Amount of biomass corresponding to energy coming from crude oil divided by 1000000 to get Mha} / \text{Yield energy crop mix mean value} / \text{devide by 1000000 to get Mha}
\]

\[
\frac{t_{DM}}{Mj/kg} /1000000 = Mha/year
\]

The calculations on energy balance make it possible to compare the cultivated area from two perspectives; the first having the carbon content as preference and the second having the energy content as preference. This can give a deeper understanding for the complexity of exchanging crude oil as raw material within chemical industry.

![Figure 12: EU-27 chemical industry, incl. pharmaceuticals, energy consumption by source on annual basis (Cefic, 2010).](image)
Figure 12 shows the energy consumption within chemical industry in EU-27. Here energy used as feedstock and energy used for transportation and processes are separated and displayed with different colours. For calculations on carbon cycle and energy balance in chemical industry the amount of crude oil used as feedstock, 69.1 M toe has been used.

To easy display variables from data collection a table of figures has been composed, see Table 5. Figures in italic has not been used in calculations for carbon cycle or energy balance, but can be used as reference value for further comparison. Other figures have been used in calculations done with equations above.

Table 5: Basic figures for calculations on C-cycle.

<table>
<thead>
<tr>
<th>Basic facts</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-content in crude oil (Johansson, 2010)</td>
<td>20</td>
<td>gC/MJ</td>
</tr>
<tr>
<td>C-content in energy crop mean value (calculated)</td>
<td>25.7</td>
<td>gC/MJ</td>
</tr>
<tr>
<td>Arable land EU-27 (ENCROP, 2010)</td>
<td>111</td>
<td>Mha</td>
</tr>
<tr>
<td>Permanent pasture land EU-27 (ENCROP, 2010)</td>
<td>69</td>
<td>Mha</td>
</tr>
<tr>
<td>Cultivated land Europe, inclusive Ukraine (Berndes, 2011)</td>
<td>164</td>
<td>Mha</td>
</tr>
<tr>
<td>Permanent pasture land Ukraine (Berndes, 2011)</td>
<td>76</td>
<td>Mha</td>
</tr>
<tr>
<td>Energy content in energy crop mix DM (mean value) (ENCROP, 2009)</td>
<td>18.234</td>
<td>MJ/kg</td>
</tr>
<tr>
<td>Conversion efficiency from biomass to useful product (Dinjus, 2009)</td>
<td>87%</td>
<td></td>
</tr>
<tr>
<td>Conversion efficiency from crude oil to useful product</td>
<td>97%</td>
<td></td>
</tr>
<tr>
<td>Yield energy crop mix mean value (ENCROP, 2010)</td>
<td>10</td>
<td>tDM/ha/year</td>
</tr>
<tr>
<td>Reed canary grass (ENCROP, 2010)</td>
<td>5</td>
<td>tDM/ha/year</td>
</tr>
<tr>
<td>Energy willow (Salix) (ENCROP, 2010)</td>
<td>6.0-10.0</td>
<td>tDM/ha/year</td>
</tr>
<tr>
<td>Poplar (Italy) (ENCROP, 2010)</td>
<td>20</td>
<td>tDM/ha/year</td>
</tr>
<tr>
<td>Aspen (ENCROP, 2010)</td>
<td>5</td>
<td>tDM/ha/year</td>
</tr>
<tr>
<td>toe = energy from burning 1 tonne of crude oil</td>
<td>42</td>
<td>GJ/toe</td>
</tr>
<tr>
<td>Crude oil use in chemical industry (Cefic, 2010)</td>
<td>69.1</td>
<td>M toe</td>
</tr>
</tbody>
</table>

6.6 Sensitivity analysis

To see if the results are validated different sensitivity analysis has been carried out. For most systems and analysis this project uses variables that have an uncertain character, such as biomass yield, weighting factors or future market potentials. To understand the result in a border aspect these types of variables has been varied to show what effect they have on the out coming result.

6.6.1 Ranking

To analyse how the chosen score system and the selection of weight factors affect the result of the ranking, sensitivity analysis is done. This is carried out in Microsoft Office Excel together with the ranking, both the scoring system and the weighting factor is varied in different cases.
The analysis is performed in three different cases:

1. The winner in each criteria is awarded one bonus points which give a score of 15 points.
2. The score system is not linear and a better placing is awarded “higher” score. The used score system is 1, 2, 3, 4, 6, 8, 10, 12, 15, 18, 21, 24, 28, 32.
3. Variations of the weighting factors. Here different changes are done, but the one that is presented in the report have higher weighting for the criteria target product and lower for combination of power plant and independence.

The second option is chosen to spread out the score and award better technologies a higher score in comparison with not so good technologies. In the original case the difference between two or three places is not so large. But in this case the consequences between couples of places can be large. There is also an extra focus on the top four technologies, to see the sensitivity of the best ranked technologies to the chosen parameters.

**6.6.2 C-cycle and energy balance**

Due to large variations in energy crop yield three different (5, 10 and 20 t dm ha⁻¹ ha/year) yields has been used during calculations. Yield of energy crops vary a lot with both geographical location and spices cultivated. In addition yields of biomass will increase throughout the years when more optimised and effective cultivation standards are applied on energy crops (Grundmann, 2011). The three different yields are determined from literature source of different yield in different geographical areas, climate and spices (ENCROP, 2010). For the c-cycle and energy balance the sensitivity analyse in incorporated in the results, see chapter 7.5.
7 Results

This chapter is divided into four parts, these mirror the subchapters in chapter 6 Method. First the classification system is presented where the target products and solid residues from different biorefinery technologies are shown, this leads onto the second part, the ranking system for technologies presented in classification. The third part present example on how different technologies can be combined in order to produce right target products and valuable residues. Last subchapter give the reader the results of the carbon cycle and the energy balance.

7.1 Market potential for target products

As discussed previous in chapter 3 the biorefineries give both target products and residues, here the future market potential for the most important biorefinery products are presented. The future market potential for different target products is performed as a sensitivity analysis, concerning both technology of use and residue. The target product that will gain market shares in the future will effect development of technologies and therefore also what residues that will be available. Text here beneath is based on statistical data and future scenario analysis.

7.1.1 Syngas and biogas

In 2009 EU-27 consumed almost 500 Gm$^3$ of natural gas, which responds to 0.27 ppm of EU-27 total energy consumption (Eurostat, 2009). As seen today the fraction of gas as energy supplier is very small, but with increasing oil prices this fraction can easily increase both in EU-27 and worldwide. Natural gas can be replaced both by syngas or biogas after upgrading, due to different energy content in biogas it is preferable as transportation fuel and syngas is appropriate for production of chemicals. Table 6 shows what volume that is needed to replace different fractions of natural gas. Today 11.8 Gm$^3$ of biogas is produced, this number can be used as reference value to understand that there has to be a large increase in production of syngas and biogas, if these are suppose to replace the today’s use of natural gas. If it is added to the picture that gas consumption will probably increase with increasing oil price, production of renewable gases has to increase further. Only to replace 10% of natural gas with biogas the production has to increase 639%.

Table 6: Percentages of EU-27 gas utilisation replaced with biogas or syngas

<table>
<thead>
<tr>
<th>Fraction replaced</th>
<th>Syngas G m$^3$</th>
<th>Biogas G m$^3$</th>
<th>% increase in biogas production</th>
</tr>
</thead>
<tbody>
<tr>
<td>10%</td>
<td>98.35</td>
<td>75.40</td>
<td>639%</td>
</tr>
<tr>
<td>20%</td>
<td>263.86</td>
<td>150.80</td>
<td>1278%</td>
</tr>
<tr>
<td>30%</td>
<td>295.04</td>
<td>226.20</td>
<td>1917%</td>
</tr>
<tr>
<td>40%</td>
<td>393.39</td>
<td>301.60</td>
<td>2556%</td>
</tr>
<tr>
<td>50%</td>
<td>491.74</td>
<td>377.00</td>
<td>3195%</td>
</tr>
</tbody>
</table>

7.1.2 Ethanol

One of the most important chemicals from ethanol at the market is ethylene, this is an important part of many product categories. Such as polymers, oligomers and ethylene oxide which can play an important role in different chemical production. Ethylene can be produced in different ways and most common today is production from a fossil base in a cracker. It can also be produced with dehydration of ethanol from biomass
fermentation or gas from gasification or digestion. The ethylene production is very important for the plastic industry and the most important role for the ethanol today is the transport industry.

Table 7: Market share of bioethylene according to today’s ethylene production in EU-27 (Eurostat, 2011), (ePure, 2010)

<table>
<thead>
<tr>
<th>Part Bioethylene</th>
<th>M tonne Bioethylene</th>
<th>M tonne Ethanol</th>
<th>% of capacity case 2009</th>
</tr>
</thead>
<tbody>
<tr>
<td>10%</td>
<td>1.72</td>
<td>2.82</td>
<td>40%</td>
</tr>
<tr>
<td>20%</td>
<td>3.43</td>
<td>5.64</td>
<td>79%</td>
</tr>
<tr>
<td>30%</td>
<td>5.15</td>
<td>8.46</td>
<td>119%</td>
</tr>
<tr>
<td>40%</td>
<td>6.86</td>
<td>11.27</td>
<td>159%</td>
</tr>
<tr>
<td>50%</td>
<td>8.58</td>
<td>14.09</td>
<td>198%</td>
</tr>
</tbody>
</table>

Table 7 shows in the first column the percentage part of the ethylene production that will be changed to bioethylene and the second column shows the necessary amount of bioethylene in mega tonne. The third column gives the necessary quantities of bioethanol and the last column shows a comparison with the production capacity of 2009. As can be seen in Table 7 a start of a bioethylene production will take a big part of the ethanol production capacity even at such as low parts as 10%. It is important to keep in mind that the ethanol production for transport services has increased with several of hundreds percent the last five years and the last year with around 30% (ePure, 2010). With focus only at the transport sector there is an over capacity in production at the moment. But if the increase of consumption will continue with the same rate, this over capacity will turn over to a shortage in a few years. In total this means that there is a huge potential of production of ethanol in EU-27 and the situation looks similar in Germany. Ethanol can also be used for other purposes in chemical industry than fuel and plastics.

There is some ongoing bioethylene research at some of the big sites in Europe, in example Stenungsund. Another good example is Braskem, in Brazil, that have started the first bio polyethylene site in the world. In this project Braskem cooperates with some of the largest packing producers (i.e. Papier-Mettler) in the world to produce sustainable packing applications with sugar canes as base (Braskem, 2011).

7.1.3 RME

There are some different ways to sell this fuel either it can be sold as B100 which is 100% of RME and a 100% renewable fuel. Another way to sell the RME is to blend it with diesel and sell this blended product. In Europe especially in Germany and France large amount of this blended fuel is used. The problem with RME is that it is still a quite expensive fuel and needs subsidies to be competitive.

Germany is one of the examples of countries with subsidies for RME but now the government decreased the tax subvention and the market increase flattened out (EurObserv’ER, 2009). But still there is an increasing market in EU-27 and in addition to that the European Union has a target for 2020 that 10% of the transportation fuel should be covered by “renewable energy” (European Union, 2009). This means that there must be a huge potential for different renewable transportations fuels. According to Eurostat around 3.5% (6.5% in Germany) of the transportation fuel is renewable (Eurostat, 2008). At the moment there is an ongoing discussion
about the sustainability of RME as biofuel which can result in a future decreased interest in this fuel.

7.1.4 Lignin and paper

Lignin is, as described, a very interesting residue that is necessary to separate from the cellulose. The residue is interesting because of the high energy content, but also because of a future potential in the production of chemicals such as aromatics. There are some well-known chemical products extracted from lignin as lignosulfonate, examples from these are vanillin and pesticides (Doherty, Mounsavioun, & Fellows, 2010). The chemical industry is very interesting for companies producing lignin because of high prices of chemicals which can give higher prices for high quality lignin. At the moment the chemical extraction of chemicals from lignin is in the research phase. There is no big competition over the lignin between the energy sector and the chemical industry today, but this can change in the future. There are also possibilities to use the lignin as raw material in gasification to produce syngas and energy.

The main product from cellulose separation today is paper and the two biggest categories of produced paper are printing/writing and packaging, together they stand for around 60% of the worldwide production. The paper production worldwide increases every year and there is a clear connection of paper use and income in a country, higher income means consumption of more paper (Brännvall, 2009). This means that the paper branch is quite safe at the moment due to the volumes but there can on the other hand be new and cheaper producers in the future. Connected to the concurrence situation it is important to the European companies to keep the quality and learn to use as much of the wooden raw material as possible.
7.2 Classification of biorefinery technologies

During this Master thesis ten main groups of different biorefinery technologies have been reviewed and classified. Arrows in Figure 13 helps reader to follow steps in value chain, interconnections between different sections are marked with diagonal arrows. Out of these ten processes, five technologies give five different valuable solid residues; see black boxes under “Residues” in Figure 13. These residues have properties suitable as fuel for co-combustion and they are cellulose based. Some other “solid” residues such as glycerol, rape meal and sludge have shown to be inappropriate for co-combustion with coal. Among these five different three residues have shown to be most fitting, these are lignin, AHR and SSFR, see tables Table 4 and Table 3. It has been hard to estimate the output of specific residues from each technology. This is partly due to the fact that there are hardly any commercial biorefineries to study today, most activities in this field are still in a research phase and not commercialised. It is also important to mention that as combustion fuel some of these solid residues do have competition with other valuable products. Lignin for example is used for producing vanillin, glycerol is used in food and cosmetic industry and rape meal can be used as animal feed. Bark can be suitable for co-combustion if it
is pre-treated and dried to pellets before, but this is not believed to be economical feasible. As for char coal it is often not extracted, rather used for direct combustion within the own biorefinery plant.

7.3 Ranking of biorefinery technologies

The different criteria’s and their explanation are described in the following table, see Table 8. This table is then used to allocate the score to the technologies in each criteria. The feedstock in to the system is cellulose based, except from anaerobic digestion and fermentation (1<sup>st</sup>).

*Table 8: Explanation of criteria’s in ranking.*

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Explanation</th>
<th>Score comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target product (TP)</td>
<td>Possible use of TP. Quality of TP. Amount of product and final or intermediate product.</td>
<td>Good applications give high score and vice verse.</td>
</tr>
<tr>
<td>Market potential (TP)</td>
<td>How is the market potential today and in the future?</td>
<td>High score for good market possibilities.</td>
</tr>
<tr>
<td>Development status technology</td>
<td>How is the development status for the technology?</td>
<td>See ch. 3. Existing technology gives a high score.</td>
</tr>
<tr>
<td>Independence</td>
<td>Can the technology stand alone or is it a part in a bigger system.</td>
<td>High score for good stand alone possibilities.</td>
</tr>
<tr>
<td>Residue</td>
<td>Properties of residue: state, energy content and chemical composition.</td>
<td>Solid residues with co-combustion properties give a high score.</td>
</tr>
<tr>
<td>Combination with power plant</td>
<td>Are there possibilities to use streams from a coal fired power plant?</td>
<td>High score for good integration possibilities.</td>
</tr>
<tr>
<td>Residue market potential</td>
<td>Are there other valuable markets for the residue?</td>
<td>Low score for competing markets.</td>
</tr>
</tbody>
</table>

The weighting process is based on different weighting factors which are connected to the objective. The most important object for this thesis is to find an interesting residue that can be used in a co-combustion process. This is the reason why the criteria “Residue” have the highest weighting factor. To get as much good residue as possible it is important to have an interesting target product that companies want to produce and customers want to buy. That is why the criteria connected to target product is the second and third most important. The market potential for the target product, weighted as third criteria, is hard to predict and that is the reason why “Market potential target product” have a lower weighting than “Target product”. Development status is given a low weighting and that is mainly based on the fact that the development status can change fast if right research focus is chosen. The fact that it can change fast and that the criteria have minor importance are the reason for low weighting in the other cases. Also criteria’s with minor importance and a low weighting factor is important for the whole process. In Table 9 the weighting factors used in the ranking are presented.
Table 9: Weighting factors for ranking.

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Weighting (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target product</td>
<td>25%</td>
</tr>
<tr>
<td>Market potential target product</td>
<td>15%</td>
</tr>
<tr>
<td>Development status technology</td>
<td>5%</td>
</tr>
<tr>
<td>Independence</td>
<td>5%</td>
</tr>
<tr>
<td>Residue</td>
<td>30%</td>
</tr>
<tr>
<td>Combination with power plant</td>
<td>10%</td>
</tr>
<tr>
<td>Residue market potential</td>
<td>10%</td>
</tr>
<tr>
<td></td>
<td>100%</td>
</tr>
</tbody>
</table>

The result of the ranking is presented in Table 10. There are four dominating technologies in the ranking both based on score and weighted score; fermentation (2\textsuperscript{nd}), pyrolysis, gasification and hydrolysis. In the bottom of the table the two types of liquefaction and some of the lignin technologies are found. Finally in the middle some interesting technologies such as digestion. In the top fermentation (2\textsuperscript{nd}) is a bit better than the other technologies and that depends on a good product and a good residues especially in combination with hydrolysis. Those are the main reason to the high weighted score and also secure the first place of fermentation (2\textsuperscript{nd}). This ranking includes some parts that are subjective especially connected to the allocation of score in the criteria. The basis of judgement will be further presented in Appendix B.

Table 10: Results of the ranking.

<table>
<thead>
<tr>
<th>Total score</th>
<th>Ranking total score</th>
<th>Weighted score</th>
<th>Ranking weighted score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fermentation (2\textsuperscript{nd})</td>
<td>74</td>
<td>2</td>
<td>12.2</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>75</td>
<td>1</td>
<td>10.9</td>
</tr>
<tr>
<td>Gasification</td>
<td>69</td>
<td>4</td>
<td>10.7</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>73</td>
<td>3</td>
<td>10.3</td>
</tr>
<tr>
<td>LPS</td>
<td>53</td>
<td>9</td>
<td>8.5</td>
</tr>
<tr>
<td>Transesterification</td>
<td>58</td>
<td>6</td>
<td>7.9</td>
</tr>
<tr>
<td>Anaerobic Digestion</td>
<td>56</td>
<td>7</td>
<td>7.7</td>
</tr>
<tr>
<td>Fermentation (1\textsuperscript{st})</td>
<td>55</td>
<td>8</td>
<td>7.4</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>40</td>
<td>10</td>
<td>6.9</td>
</tr>
<tr>
<td>Barking</td>
<td>59</td>
<td>5</td>
<td>6.4</td>
</tr>
<tr>
<td>Organosolv</td>
<td>40</td>
<td>10</td>
<td>5.6</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>27</td>
<td>13</td>
<td>4.2</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>32</td>
<td>12</td>
<td>3.7</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>24</td>
<td>14</td>
<td>2.8</td>
</tr>
</tbody>
</table>
7.4 Possibilities for combination of technologies

To give the requested products, different technologies in chapter 3 can be used in combinations, here a number of these combinations will be described. Technologies from all four categories will be included in at least one example. The combination will also include production of the in chapter 4 presented target products and extraction of the in chapter 5 presented valuable residues. These three chapters and the review of the literature according to existing plants and ideas for plants have given input to the presented combinations in this chapter. One important factor in many of these combinations is to fulfil the energy requirements for the different processes, which need further investigations, both for the primary technology and for upgrading of the residue. This is important especially in the primary process, because the residue will not be used as a fuel directly on site which is normal otherwise. To solve this important question the biorefinery in these cases must be connected to the coal fired power plant which use the residues for combustion and can send back excess heat. The process integration and the necessary calculations for this are beyond the limitations of this thesis. Another good reason of building the plants close to each other is the transport of the residue. If the plants are close to each other the cost and environmental problem with the transport can be minimized. Produced chemicals have a high value and therefore transportation can be more accepted in an economical point of view. Also gas can be transported, in that case connection to the gas grid is necessary.

7.4.1 Syngas production

The syngas production from pyrolysis and gasification starts with size reduction to get the desired size. If it is necessary to remove bark it will be removed in this stage. Next process step will be pyrolysis or gasification step, in this step the biomass will be heated to high temperatures and then release the volatile gases. The volatile gases are often named syngas and have a lot of different application possibilities, such as fuel for power production or as platform for production of other chemicals (for example methanol) or fuels. The power production uses a gas turbine to drive the generator for electricity production. Depending on the process parameters the relationship between gas and the two other components will vary. The two other components are bio-oil, see chapter 4.2, which can be used in the chemical industry and char coal, see chapter 5.2. Char coal is an energy rich residue, but will often be recirculated to produce as much gas as possible. There are possibilities to take out parts of the char coal and use for co-combustion with coal.
7.4.2 Ethanol and Biogas plant

There are many ways to combine the production of ethanol and biogas. The main operations are fermentation and anaerobic digestion, here one model will be presented. The model has an ethanol production in the first step with both first and second generation ethanol and the second step is biogas production. These two main products have a wide range of applications both in direct use and in different upgrading processes, see also chapter 5.3.

An advantage with second generation fermentation is that lignin is separated and the lignin can then be used as a fuel to cover the plants heat demand. Or in this case be used as fuel in a coal fired power plant and excess heat from the plant can be used for different heat demands at the biorefinery plant. If the ethanol stream will be used for other services than fuel, further processing will be necessary. These processes can of course be done on site and can also use heat from the power plant. The solid separation can be done with evaporation which also needs extra heat. The pretreatment, in this case is hydrolysis, either needs some kind of acid or enzyme to release the cellulose before fermentation. There are some suggestions that the acid can be replaced with carbon dioxide which then can be taken from the power plant is CCS is used.

The residues from the ethanol production can then go further to the combustion process or the slurry phase (DR) to the digester for biogas production. As mentioned before the quality of the DR is not good enough to use for co-combustion with coal, it is better to use it in the digester. Also other wastes from farms or household can be put in the digester to increase the produced amount of gas. This gas can be consumed directly on site, upgraded and sold to the gas network or as transportation fuel. The fertilizer produced in the waste water treatment (WWT) can be used at farms, even the solid part from the fermentation can be used as fertilizer but it can also be used in the in the power plant. The process scheme can be seen in Figure 15 and the pretreatment step includes size reduction and hydrolysis.
7.4.3 RME transesterification

Figure 16 shows a theoretical process design of a biorefinery with the main product RME. The RME production is special because it does not use cellulosic feedstock and instead oil based crops. The process have two possibilities to take out fuel for a combustion process, first in the chemical extraction where rape seed meal is extracted and then in the transesterification where glycerol can be decanted of. Due to the high value for rape seed meal as animal feed this residue is of no interest for co-combustion with coal. If glycerol should be used as a fuel drying process is necessary. The temperature level for the transesterification is around 60°C so can be supplied by excess heat from power plant.

Another important part for the RME production is the use of methanol. Methanol is used in excess and everything will not react with triglyceride this gives possibilities to reuse some parts of the methanol. Even if some part is reused new methanol is necessary to add on to every batch. Methanol can be produced from syngas which can be produced from gasification of biomass. This means that also a gasification process can be integrated with the RME production and the syngas which not is necessary for the methanol production can be used for other purposes.
7.4.4 Cellulose separation

The separation between cellulose and lignin are done for different purposes such as ethanol production 2nd generation and paper production. Also production of other chemicals can be interesting in the future and there is some ongoing research on this area. The separation technologies are described in chapter 3.5. The first step to extract cellulose from lignin is to size down the wood stock and also barking of the wooden feedstock. Bark is traditionally used as a fuel internally or sold on the market and can be used in co-combustion. The second step is to separate out the lignin, also the lignin is traditionally used internally as fuel for heat but it can be extracted and used in the co-combustion process. If bark and lignin are used in the co-combustion the excess heat from the plant must go back to the separation processes. There is an excess of bark and lignin, and especially the lignin has other possible use for example vanillin or other chemical production. The chemical production from lignin is an ongoing research area.

![Figure 17: Process scheme for cellulose/lignin separation (Eriksson, 2009)](image)

7.5 Result from C-cycle and energy balance

Calculations on both carbon cycle and energy balance are performed with figures from Table 5 and equations in chapter 6.5 have been used in the same order as presented. In this master thesis conversion efficiency is set to when syngas is produced from crude oil and biomass, respectively 97% and 87% (Dinjus, 2009). Here beneath follows one example of a calculations to further simplify comprehension.

**Equation 9:** Calculations of total gC coming from crude oil.

\[
\frac{C \text{ content in crude oil}}{\text{Energy unit}} \times \frac{\text{Amount of crude oil used in chemical industry}}{\text{conversion efficiency for crude oil to useful product}} \times \frac{\text{Energy content}}{\text{Oil equivalent}}
\]

\[
= \frac{20 \text{ gC}}{\text{MJ}} \times 69.1 \text{ Mtoe} \times \frac{42,000 \text{ MJ}}{\text{Mtoe}} \times 97\% = 5.63 \times 10^{13} \text{ gC}
\]

**Equation 10:** Calculations of carbon content in energy crop willow.

\[
\frac{C \text{ content in energy crop mix mean value}}{\text{Energy content energy crop mix DM}} \times \text{multiplied with 1000 to get g}
\]
Equation 10 was done for willow, hemp, reed canary grass and poplar. All these values were then added together and divided by four to get a mean value of carbon content in energy crop mix, this showed to be 25.68 gC/MJ. Results from Equation 9 and Equation 10 are used in Equation 11 together with figures from Table 5.

**Equation 11:** Calculations of need for biomass to cover total amount of carbon from crude oil.

\[
\frac{49.80 \text{ kg C}}{18.62 \text{ MJ/kg}_\text{DM}} \times 1000 = \frac{26.75 \text{ g C}}{\text{MJ}}
\]

Equation 12 give that 14.0 Mha/year of land area needs to be cultivated in order to cover carbon content from crude oil in chemical industry. Same calculations were done for all three yields, 5, 10 and 20 t$_\text{DM}$/ha/year. Calculations on energy balance was performed in a similar way, but used energy content in crude oil and energy crop as basis for comparison, see Appendix C.

In Figure 18 the result from calculations of carbon cycle, shown by red bars, and energy balance, shown by blue bars, can be viewed. Each bar represents one of the sensitivity analyses by using different yields of biomass. Figure 18 shows that out of an energy perspective the demand for cultivated land area is higher for all three yields than for the carbon cycle.

**Figure 18:** Area of land that needs to be cultivated by energy crops in order to replace energy and carbon coming from crude oil. Number above staples show percentage of total arable land.
Total farmland includes both arable land and permanent pasture land in EU-27. In EU-27 there is 111 million ha of arable land and 69 million ha of permanent pasture land available (ENCROP, 2010). To facilitate if areas in Figure 18 are reasonable a future scenario of demand for food and energy is used. The scenario assumptions for food demand in 2030 are chosen from the book *European Energy Pathways*. Here it is shown that 44-53 Mha of cultivated land and 20 Mha of pasture land might not be needed for food production (Berndes, 2011). This corresponds to 40-47% respectively 29% of EU-27 present agricultural land area. Figure 18 shows that even with the lowest yield of energy crops the need for cultivated area is smaller than the expected “free land” (44-53 Mha), more specific 32.2 Mha for energy balance and 25.0 Mha for C-cycle is needed. Although it is important to keep in mind that result from forecasting demand for energy and food in the future always depends heavily on how the scenario assumptions are chosen.

### 7.5.1 Biomass economy

The carbon cycle and energy balance shows that theoretical there is enough land available to cultivate the biomass needed to replace crude oil in chemical industry. Another interesting question would be to see how much the crude oil price has to increase in order for biomass to be competitive with crude oil.

Assumptions regarding price and conversion level have been done to perform simplified calculations regarding biomass economy. The price of biomass used in calculations are 200 €/MWh and 78 €/barrel for crude oil (Grundmann, 2011) (Oil-price.net, 2011). Conversion level for crude oil to useful product is assumed to be 97%. Biomass is simplified to consist of 33% lignin and 67% cellulose, the chemical industry is assumed to use all cellulose and separate all lignin in the biomass feedstock.

Three different scenarios has been viewed, in all the chemical industry use all cellulose. In the first the lignin is separated but not sold, in the second case chemical industry sell the lignin part to energy industry and in the third scenario chemical industry sell both lignin and emission permits that they can avoid by using biomass as raw material. These three scenarios gives three different “future crude oil price”, this means what level crude oil price has to reach in order for biomass to be economical preferable or at least feasible. All three scenarios are calculated with the assumption that chemical industry does not get any subsidies for using biomass instead of crude oil.

In Table 11 the result for future crude oil price can be viewed. The three different “future crude oil price” are marked with italic typing. Also the percentage increase can be viewed.

As seen the three different scenarios do not give any major impact on the different future crude oil price, not even scenario three that on forehand where believed to make a major difference on “future crude oil price”. For all three cases biomass is competitive with crude oil first at levels of 300 % increase of today’s crude oil price. Conclusion can be made that with this economically situations the exchange of crude oil towards biomass will be postponed. This not only because of the price of biomass, the exchange of crude oil towards biomass is more complex. This process does have both economical, political, technical and resource problems. The calculations done here are very rough and are performed to show further one of the problems with
exchanging crude oil to biomass. Here only crude oil price is taken into account, resources like coal and natural gas are excluded, and also technical improvements for both biomass utilisation and extraction of fossil raw material are excluded.

*Table 11: Calculations on future crude oil price.*

<table>
<thead>
<tr>
<th>Calculated parameter</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy coming from crude oil</td>
<td>2815</td>
<td>PJ</td>
</tr>
<tr>
<td>Total cost for crude oil</td>
<td>38619</td>
<td>million €</td>
</tr>
<tr>
<td>Total cost for biomass</td>
<td>139979</td>
<td>million €</td>
</tr>
<tr>
<td>Today price of crude oil</td>
<td>78</td>
<td>€/barrel</td>
</tr>
<tr>
<td>Today price of biomass</td>
<td>200</td>
<td>€/MWh</td>
</tr>
</tbody>
</table>

**Scenario 1**

<table>
<thead>
<tr>
<th>Future crude oil price</th>
<th>284</th>
<th>€/barrel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increase in percentage</td>
<td>362%</td>
<td></td>
</tr>
</tbody>
</table>

**Scenario 2**

| Lignin replacing hard coal         | 840 | PJ      |
| Revenue lignin                    | 2320| million €|
| Future crude oil price             | 279 | €/barrel |
| Increase in percentage             | 356%|         |

**Scenario 3**

| Avoided g CO₂                    | 178428823 | t CO₂   |
| Avoided emissions rights          | 2378      | million €|
| Future crude oil price             | 274       | €/barrel |
| Increase in percentage             | 350%      |         |
8 Sensitivity analysis

In this chapter the results from the sensitivity analysis based on ranking and market potential are presented. The sensitivity analysis for c-cycle is presented directly in chapter 7.5.

8.1 Ranking of biorefinery technologies

As described in the method chapter (6.6.1) the sensitivity analysis of the ranking is performed in two parts, one with changes for the whole list and one for the top four technologies. The result of the first part of the sensitivity analysis can be seen in Table 12. Notice that there are no big changes between the original and the three cases in the analysis and the top four are intact. This means that with this construction of the ranking the system is stable. The ranking in Table 10 it can also be seen that the top four have high scores in comparison to the other ten technologies and as a result of this it is very hard to knock one of them out. In the table head W means position after weighting and T means total score before weighting.

Table 12: Result of sensitivity analysis for ranking.

<table>
<thead>
<tr>
<th>Technology</th>
<th>Original ranking</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>W</td>
<td>T</td>
<td>W</td>
<td>T</td>
</tr>
<tr>
<td>Fermentation (2nd)</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Gasification</td>
<td>3</td>
<td>4</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>4</td>
<td>3</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>LPS</td>
<td>5</td>
<td>9</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Transesterification</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Anaerobic Digestion</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>Fermentation (1st)</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>9</td>
<td>10</td>
<td>9</td>
<td>10</td>
</tr>
<tr>
<td>Barking</td>
<td>10</td>
<td>5</td>
<td>10</td>
<td>5</td>
</tr>
<tr>
<td>Organosolv</td>
<td>11</td>
<td>10</td>
<td>11</td>
<td>10</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>12</td>
<td>13</td>
<td>12</td>
<td>13</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>13</td>
<td>12</td>
<td>13</td>
<td>12</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>14</td>
</tr>
</tbody>
</table>

In the second step where the analysis of the top four is performed no new observations was done due to that the total changes was small. But as can be seen in the table there are changes in the top four within the three new cases. This is of course connected to the small difference in score between these technologies. The result of this is that fermentation (2nd) and pyrolysis are still the best, except for one case these two technologies are on first or second place. Bark is the largest loser of the weighting system and LPS is the winner. Bark is winner in two criteria’s (DS Technology and Independence) but these two criteria’s has the lowest weighting factor and this affect the weighted score a lot. LPS have scores in the middle but extra high score (11) in the residue criteria which have a high weighting factor.
9 Discussion

This work shows that there is a huge potential for bio based products in the future. There are probably not possibilities to replace all fossil products, due to both resource availability and technical hurdles, therefore it is important to focus on some of them. As mentioned before gas has a broad market and there are strong intensives to get out more and “new” gases on the market. The gas market increase every year and EU-27 is at the moment depending on Russian natural gas which can give political problems in the future. Although technologies producing gaseous products do not have any residues that are of interest for this Master’s thesis.

The top four technologies in the ranking are not unexpected due to their products and residues but there are some statements to do about the result. First anaerobic digestion is a promising technology for the future and will maybe earn a higher rank than place seven. This technology has here lost credits because of the non suitable residue, which is one of the main focuses in this Master thesis. Also the feedstock to the anaerobic digester can be based on all organic materials not only wooden feedstock and an investment in wooden based biogas is probably not a sustainable idea when the wood can be used for other purposes. If different gaseous products are the most interesting, a solution with gasification is the most appropriate. Another technology that need comments is hydrolysis, this technology is in the top four and have quite high credits due to good residues. Here it is necessary to keep in mind that hydrolysis in this case is a technology to separate lignin and cellulose. This means that the produced target products are intermediates or improvements of the feedstock.

The lignin technologies that can be found in the middle of the ranking often give good residues, however large facilities (pulp mills and separation facilities) are necessary. The result of this is that older mills can be retrofitted and new mills can be built in this way, but it is not the most interesting option for the energy or chemical companies to invest in paper mills. In the last two places different liquefaction technologies are found, there can be opportunities for these technologies in the future, but in this ranking with these categories the score is lower than for the other technologies.

There are both positive and negative aspects with different solutions of technologies combinations, but the target of this Master thesis is to get valuable products and residues that can decrease the usage of coal and crude oil. One superior option is to produce ethylene from the bioethanol and then replace the fossil based ethylene. As described in the report ethylene can be produced from ethanol (fermentation) or from syngas (gasification) and because of the residue the recommendation will be to focus on fermentation.

Connected to the chapter 7.4 about combination of technologies there are also important questions about logistics. The heat from the coal power plant should not go longer than necessary, this to prohibit loses of heat. Transportation regarding heat, products and residues all has to be considered to be as efficiently as possible. With a close location the transportation of biorefinery residues can be minimized and can in the best case be transported on conveyor to the power plant. Transports of products or intermediate products which have a higher economic value can be done with truck or train and in some case, depending on the site location, with barges. The texts about a modern paper mill and the Stenungsund industry cluster shows the future potential to combined a biorefinery, a coal fired power plant and a chemical plant. As shown in the report a couple of different technologies can be connected to produce different
products. The most important is not to produce many products, it is to produce large quantities of some of them to be an important market player.

In the work with the carbon cycle and the energy balance many different assumptions where made. The results from both analyses are very dependent on these assumptions and results could be illuminating to fit different purpose. To make this report as transparent as possible follows here a discussion on factors that has an effect on the results.

For all type of biomass the yield varies a lot with both geographical location and spices cultivated. This report does take these fluctuations into account by doing a sensitivity analysis. Conventional farming has increased its yields by 1.5 to 2 % each year the last 50 years, this will also be the case for energy crop cultivation, but such optimisation is not taken into account in this report. If this would be taken into consideration it is likely to believe that energy crop yield will increase in the future.

Also conversion efficiency varies a lot with different technologies and end product. For both crude oil use and biomass use the raw material is converted into several valuable products along the value chain, and for most technologies the residues are taken care of and used. This makes the assumptions around conversion efficiencies complex.

The comparison of needed cultivated area with available farmland can be done with a number of different reference values. In this project two different values were chosen, available arable land and total farmland. Often the land use is affected by many factors which makes it complex question. To mention a few parameters the availability of land is affected by competition of interest, national restrictions, shape of land and location. All these parameters can prohibit utilisation due to technical, social and political hurdles. In EU-27 there are also many restrictions regarding nature preservation which limits further cultivation.

When starting discussion with chemical industry for a possible cooperation insight and arguments around biomass economy is valuable. To get well built up arguments more complex calculation on biomass economy has to be performed. Although it is surprising to see how much the crude oil price has to increase in order to reach biomass levels. It would be interesting to see more detailed calculations to see if this picture is changing, with for examples political subsidies.

We have through this project learned many important lessons both regarding methodology and target image. If we would do this once again we would try to have a clearer picture of what we wanted to achieve at the end. This would have given us the opportunity to go deeper in to some areas that we found more interesting, for example the ranking system. We did try to find an appropriate methodology instead of looking forward to what we wanted to learn. Regarding methodology it is important to keep a broad mindset, even if the goal stands clear. This was the case with the carbon cycle and the energy balance, here we learned well how to see a problem out of different angels to avoid a lock in. One big downbeat in this study is that there have only been theoretical studies of different technologies, this due to the fact that there are very few biorefineries existing, producing chemical products. In the beginning of the project we had hoped to see facilities and implementations live, but unfortunately this was not possible.
10 Conclusions and recommendations

This Master thesis concludes that there are good technical potentials for cooperation between the chemical industry and the energy sector. The fact that there are very few commercial biorefinery producing chemicals indicates that this branch is far away from replacing crude oil. If energy industry wants to benefit from development towards green products now is the right time to start cooperation, before any new power plants are build or even planned.

There are five solid residues from different biorefinery technologies that can be co-combustion with coal. These are char coal, lignin, AHR, SSFR and bark. Out of these five, lignin, AHR and SSFR are the most suitable both to chemical properties and technology used.

In chapter 7.4 different combinations of biorefinery technologies are presented. Due to the market potential analysis bioethanol will gain large market shares in the future. This makes the combination with fermentation (2\textsuperscript{nd}) and hydrolysis to the most prominent solution, based on the ranking. Hydrolysis has a high score in the ranking but needs combination with other technologies to give a target product. This technology combination needs further development on wooden based ethanol production before commercialisation of is possible. Other technologies with high ranking scores are pyrolysis and gasification, the problem with these technologies are that the residue is hard to extract from the plant.

Another important question around the change from crude oil based to biomass based chemistry is the question about land use. For the performed energy balance the expected demand for cultivated land area is higher than for the expectations for the carbon cycle. This pin points the complexity on how to determine the need for energy crops to cover the crude oil used today in chemical industry. Results depend heavily on conversion rate and this figure is changing with better technology and different end products. Conversion rate used here for crude oil and biomass into useable product is a simplification and this should be kept in mind when studying the results.

Results from carbon cycle and energy balance give an indication on how much biomass that will be needed in the future and calculations shows that the demand is far lower than the available land in EU-27. The complexity of problem definition makes it hard to determine if performed calculations are “right” and depending on input data, results can be manipulated to fit the purpose of stakeholder. Calculations in this Master thesis shows that there is enough theoretical land areal to cultivate biomass to replace the use of crude oil in chemical industry, but before this can happen many social and economical hurdles has to been torn down. Although it should be mentioned that the use of crude oil in EU-27 is 460.6 million toe and chemical industry uses only 69.1 million toe. Further investigation on how much of theoretical land area in EU-27 that actually can be used for cultivation of food or energy crops is needed. Calculations in this Master’s thesis on biomass economy showed that crude oil still do have a great economical advantage over biomass as raw material.

Investigations connected to logistics and process integration between chemical sites and power plants has to be performed deeper. It is necessary that the chemical industry think about if and how they are ready to implement these new green chemicals and if they fulfil the quality requirements.
The chemical industry is known to be a very traditional industry which has low exchangeability on raw material and is loyal to their supplier. They rather stay with one supplier known to be good, than change to a cheaper or more environmentally friendly one which they do not have the same trust for. As for today there are no clear incentives, such as tax reduction or subsidy, for the chemical industry to exchange crude oil to renewable feedstock and crude oil is still cheaper and easier to convert into valuable products. To speed up this process political incentive has to take place, but also cooperation with other branches is believed to encourage the exchange. Although the chemical industry is very complex, it is becoming greener and to still be competitive enterprises has to follow this trend.
11 References


Vattenfall AB. (2010). *Six sources of energy - one energy system (Coal power)*. Stockholm: Vattenfall AB.


Appendix A

The list here below shows people we have contacted during the work with our Master thesis. In the list you may find name, title, contact information and date of conversation by email, telephone or personally.

<table>
<thead>
<tr>
<th>Name</th>
<th>Title / Contact information</th>
<th>Date of contact</th>
</tr>
</thead>
</table>
| Stefan Wirsenius | PhD Physical Resource Theory  
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Appendix B

Table 13: Total score and ranking for all categories, $S=$Score and $W=$Weighting.

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Weighted</th>
<th>Total</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Score</td>
<td>Rank</td>
<td>Score</td>
</tr>
<tr>
<td><strong>Technology</strong></td>
<td>$S$</td>
<td>$W$</td>
<td>$S$</td>
</tr>
<tr>
<td>Anaerobic Digestion</td>
<td>12</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>Barking</td>
<td>5</td>
<td>1.25</td>
<td>3</td>
</tr>
<tr>
<td>Fermentation (1st)</td>
<td>11</td>
<td>2.75</td>
<td>10</td>
</tr>
<tr>
<td>Fermentation (2nd)</td>
<td>14</td>
<td>3.5</td>
<td>14</td>
</tr>
<tr>
<td>Gasification</td>
<td>13</td>
<td>3.25</td>
<td>13</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>6</td>
<td>1.5</td>
<td>6</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>7</td>
<td>1.75</td>
<td>4</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>2</td>
<td>0.5</td>
<td>5</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>1</td>
<td>0.25</td>
<td>9</td>
</tr>
<tr>
<td>LPS</td>
<td>8</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>3</td>
<td>0.75</td>
<td>1</td>
</tr>
<tr>
<td>Organosolv</td>
<td>4</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>9</td>
<td>2.25</td>
<td>11</td>
</tr>
<tr>
<td>Transesterfication</td>
<td>10</td>
<td>2.5</td>
<td>7</td>
</tr>
<tr>
<td><strong>Summary</strong></td>
<td>105</td>
<td>26.25</td>
<td>105</td>
</tr>
</tbody>
</table>

3 TP = Target product  
4 DS = Development status  
5 CPP = Combination with power plant  
6 MP = Market potential
<table>
<thead>
<tr>
<th>Technology</th>
<th>Target product</th>
<th>Potential TP</th>
<th>DS Technology</th>
<th>Independence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaerobic Digestion</td>
<td>Biogas that can replace NG</td>
<td>Can replace NG &amp; NG has a growing market with good infrastructure</td>
<td>Existing world wide</td>
<td>Can be used on its own &amp; is often connected to farms or waste disposal</td>
</tr>
<tr>
<td>Barking</td>
<td>Wood material or bark free tree feedstock</td>
<td>There is already a lot of bark on the world market</td>
<td>A well known technology used for decades</td>
<td>Don't need anything more than a tree buyer</td>
</tr>
<tr>
<td>Fermentation (1st)</td>
<td>Ethanol</td>
<td>Growing market as fuel &amp; chemical</td>
<td>Existing especially in Brazil &amp; USA</td>
<td>Give ethanol and can stand independent</td>
</tr>
<tr>
<td>Fermentation (2nd)</td>
<td>Ethanol</td>
<td>Growing market as fuel &amp; chemical</td>
<td>Demonstration and existing</td>
<td>Due to the lignocelluloses feedstock some pretreatment is necessary</td>
</tr>
<tr>
<td>Gasification</td>
<td>Synthetic gas</td>
<td>Huge potential as NG replacer but also for chemical extraction</td>
<td>Demonstration and existing</td>
<td>Work alone</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>Different sugars from starch such as cellulose</td>
<td>Extracted cellulose can i.e. be used for fermentation (2nd)</td>
<td>Old existing technology</td>
<td>Give product such as sugar but need connection to work as biorefinery.</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>Pulp/cellulose</td>
<td>Paper market increase, but not significant. Cellulose can be interesting</td>
<td>Pilot plant i.e. Bäckhammar</td>
<td>Need paper mill and extraction technology</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>Liquids such as alcohols</td>
<td>Good for bio based chemicals</td>
<td>Pilot plant</td>
<td>Quite independent, give product on site</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>Mixture of gas &amp; liquids</td>
<td>Good for bio based chemicals and gas</td>
<td>Developed for coal work with fermentation</td>
<td>Need for separation</td>
</tr>
<tr>
<td>LPS</td>
<td>Pulp/cellulose</td>
<td>Hard to forecast, but good quality</td>
<td>Existing in France/India</td>
<td>Need a paper mill</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>Pulp/cellulose</td>
<td>Hard paper market</td>
<td>Demonstration and existing</td>
<td>Need a paper mill</td>
</tr>
<tr>
<td>Organosolv</td>
<td>Pulp/cellulose</td>
<td>Low pulp quality, not good</td>
<td>Existing</td>
<td>Need a paper mill</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>Bio-liquids (oil) &amp; Syngas</td>
<td>Interesting potential for bioprod.</td>
<td>Demonstration and existing</td>
<td>Give product on site</td>
</tr>
<tr>
<td>Transesterification</td>
<td>FAME</td>
<td>Biofuel for transport but with decreasing subsidies</td>
<td>Existing</td>
<td>Give product on site</td>
</tr>
</tbody>
</table>
Table 15: Background of judgement for score table part two.

<table>
<thead>
<tr>
<th>Technology</th>
<th>Residue</th>
<th>CPP</th>
<th>Residue MP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anaerobic Digestion</td>
<td>A wet residue that maybe can be used as fertilizer</td>
<td>This is no good options for co-combustion with coal</td>
<td>There are other possible use such as fertilizer</td>
</tr>
<tr>
<td>Barking</td>
<td>Bark</td>
<td>Fulfil a lot of requirements for co-combustion. Question about chemical composition and water content</td>
<td>Can be used for energy services but also for i.e. leather treatment</td>
</tr>
<tr>
<td>Fermentation (1st)</td>
<td>Not so energy rich but can be sent to the digester</td>
<td>Low possibilities</td>
<td>Energy services with digester</td>
</tr>
<tr>
<td>Fermentation (2nd)</td>
<td>DS, AHR, SSFR lignin rich residues</td>
<td>Spec. AHR and SSFR can be used for co-combustion with coal</td>
<td>There is no other big markets, maybe as fertilizer</td>
</tr>
<tr>
<td>Gasification</td>
<td>Char coal</td>
<td>Mainly used directly in process and not feasible to extract</td>
<td>Energy services but most direct in process</td>
</tr>
<tr>
<td>Hydrolysis</td>
<td>Lignin rich residue as AHR &amp; SSFR</td>
<td>Very good possibilities due to the lignin rich residue</td>
<td>Very good possibilities for energy use, similar to fermentation (2nd)</td>
</tr>
<tr>
<td>LignoBoost</td>
<td>Lignin</td>
<td>Shows good properties for co-combustion</td>
<td>Research on chemical extraction otherwise energy service</td>
</tr>
<tr>
<td>Liquefaction (direct)</td>
<td>Not well known, but wet</td>
<td>No good known options</td>
<td>Possibilities for digester</td>
</tr>
<tr>
<td>Liquefaction (indirect)</td>
<td>Not well known, but not so wet as direct</td>
<td>No good known options</td>
<td>Can be digester options</td>
</tr>
<tr>
<td>LPS</td>
<td>High quality lignin</td>
<td>Good for co-combustion</td>
<td>Other chemical products</td>
</tr>
<tr>
<td>Near-Natural</td>
<td>Lignin &amp; Acetic acid</td>
<td>Good for co-combustion</td>
<td>Possibilities for other use</td>
</tr>
<tr>
<td>Organosolv</td>
<td>Clean lignin (high separation)</td>
<td>Good lignin, good combustion</td>
<td>Clean lignin can be used for chemicals</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>Char coal</td>
<td>Good co-combustion possibilities</td>
<td>Optimize Char coal and extract</td>
</tr>
<tr>
<td>Transesterfication</td>
<td>Glycerol</td>
<td>Possibilities with drying</td>
<td>Use in chemical industry</td>
</tr>
</tbody>
</table>
Appendix C

Carbon cycle and energy balance

Table 16: Figures for calculation of carbon content in energy crop mix.

<table>
<thead>
<tr>
<th></th>
<th>Unit</th>
<th>Willow</th>
<th>Hemp</th>
<th>Reed Canary Grass</th>
<th>Poplar</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water content at harvest</td>
<td>%</td>
<td>50</td>
<td>10.0-15.0</td>
<td>10.0-15.0</td>
<td>50.0-55.0</td>
</tr>
<tr>
<td>Dry mass yield</td>
<td>t&lt;sub&gt;DM&lt;/sub&gt;/ha/year</td>
<td>6</td>
<td>5</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td>Max yield</td>
<td></td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Net calorific value (dry)</td>
<td>MJ/kg DM</td>
<td>18.62</td>
<td>17.48</td>
<td>18</td>
<td>18.1</td>
</tr>
<tr>
<td>Net calorific mean value</td>
<td>MJ/kg DM</td>
<td>18.05</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hydrogen</td>
<td>kg H/kg DM%</td>
<td>6.26%</td>
<td>6.00%</td>
<td>6.10%</td>
<td>7.70%</td>
</tr>
<tr>
<td>Carbon</td>
<td>kg C/kg DM</td>
<td>49.80%</td>
<td>47.30%</td>
<td>48.60%</td>
<td>39.70%</td>
</tr>
<tr>
<td>Carbon content</td>
<td>gC/MJ</td>
<td>26.75</td>
<td>27.06</td>
<td>27.00</td>
<td>21.93</td>
</tr>
<tr>
<td>Carbon content mean value</td>
<td>gC/MJ</td>
<td>25.68</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 17: Basis for diagram showing the result of carbon cycle and energy balance.

<table>
<thead>
<tr>
<th>Area cultivated to cover carbon coming from crude oil</th>
<th>Yield 20 t&lt;sub&gt;DM&lt;/sub&gt;/ha</th>
<th>7.0 Mha/year</th>
<th>Percent of total farmland</th>
<th>3.9%</th>
<th>Percent of arable land</th>
<th>6.3%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Area cultivated to cover energy coming from crude oil</td>
<td>Yield 20 t&lt;sub&gt;DM&lt;/sub&gt;/ha</td>
<td>7.8 Mha/year</td>
<td>Percent of total farmland</td>
<td>4.3%</td>
<td>Percent of arable land</td>
<td>7.0%</td>
</tr>
<tr>
<td></td>
<td>Yield 10 t&lt;sub&gt;DM&lt;/sub&gt;/ha</td>
<td>15.6 Mha/year</td>
<td>Percent of total farmland</td>
<td>8.7%</td>
<td>Percent of arable land</td>
<td>14.1%</td>
</tr>
<tr>
<td></td>
<td>Yield 5 t&lt;sub&gt;DM&lt;/sub&gt;/ha</td>
<td>31.2 Mha/year</td>
<td>Percent of total farmland</td>
<td>17.3%</td>
<td>Percent of arable land</td>
<td>28.1%</td>
</tr>
</tbody>
</table>

Following three equations shows calculations performed to calculate the energy balance. Here the calculations are done with a yield of 10 t<sub>DM</sub>/ha/year.

Equation 13: Calculations on total energy coming from crude oil within chemical industry.

\[
\text{Amount of crude oil used in chemical industry} \times \frac{\text{Energy content}}{\text{Oil equivalent}} \times \text{conversion efficiency crude oil to useful product}
\]

\[
= 69.1 \text{ M toe} \times 42 \frac{\text{GJ}}{\text{toe}} \times 97 \% = 2815 \text{ PJ}
\]

Equation 14: Calculations on the amount of biomass DM that is needed to cover energy demand.

\[
\text{Amount of total energy coming from crude oil} \div \text{Energy content in energy crops mean value DM} / 1000 \ (\text{to get tonne})
\]

\[
= \frac{2815 \text{ PJ}}{18.05 \text{ MJ/kg<sub>DM</sub>}} /1000 = 155963102 \ t_{<sub>DM</sub>}
\]
Equation 15: Calculations on the cultivated area needed to produce the necessary biomass.

\[
\frac{155963102 \, t_{DM}}{10 \, \text{ha/year}} \times \frac{1}{1000000} = 14.0 \, \text{Mha/year}
\]

Calculations for biomass economy; for chapter 7.5.1.

Table 18: Basic figures for biomass economy calculations.

<table>
<thead>
<tr>
<th>Basic facts</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Crude oil</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Energy content oil</td>
<td>37</td>
<td>MJ/L</td>
</tr>
<tr>
<td>Density of crude oil</td>
<td>840</td>
<td>kg/m³</td>
</tr>
<tr>
<td>Crude oil price</td>
<td>112</td>
<td>$/barrel</td>
</tr>
<tr>
<td></td>
<td>78</td>
<td>€/barrel</td>
</tr>
<tr>
<td>Tonne of oil equivalent</td>
<td>42000</td>
<td>MJ/toe</td>
</tr>
<tr>
<td>Crude oil use in chemical industry</td>
<td>69.1</td>
<td>M toe</td>
</tr>
<tr>
<td>1 barrel</td>
<td>158.9873</td>
<td>L/barrel</td>
</tr>
<tr>
<td><strong>Biomass</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Energy content energy crops (mean value DM)</td>
<td>18.05</td>
<td>MJ/kg</td>
</tr>
<tr>
<td>Price biomass</td>
<td>200</td>
<td>€/MWh</td>
</tr>
<tr>
<td>1 MJ</td>
<td>0.000278</td>
<td>MWh/MJ</td>
</tr>
<tr>
<td>Fraction sugar</td>
<td>0.67</td>
<td></td>
</tr>
<tr>
<td>Fraction lignin</td>
<td>0.33</td>
<td></td>
</tr>
<tr>
<td>Need for biomass (dry)</td>
<td>139591449288</td>
<td>kg</td>
</tr>
<tr>
<td><strong>Hard coal</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hard coal price average</td>
<td>80.95</td>
<td>€/tce</td>
</tr>
<tr>
<td>1 tce = energy from one tonne of coal</td>
<td>29.3076</td>
<td>GJ/tce</td>
</tr>
<tr>
<td><strong>Other variables</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Euro = 1.4313 US dollar</td>
<td>1.4313</td>
<td>$/€</td>
</tr>
<tr>
<td>1 US dollar = 0.698909701 Euro</td>
<td>0.6989</td>
<td>€/$</td>
</tr>
<tr>
<td>Energy consumption EU-27</td>
<td>75558</td>
<td>PJ</td>
</tr>
<tr>
<td>Emission Rights Spot price</td>
<td>13.33</td>
<td>€/t CO₂</td>
</tr>
<tr>
<td>CO₂-emission</td>
<td>94.6</td>
<td>g CO₂/MJ</td>
</tr>
</tbody>
</table>
Equation 16: Calculations on total cost for crude oil.

\[
\left( \frac{\text{Crude oil price}}{\text{Liters per barrel}} \right) \times \frac{\text{Energy content oil}}{\text{Crude oil use in chemical industry}} \times \text{Currency rate } \frac{€}{$} = \left( \frac{112 \frac{\text{€}}{\text{barrel}}}{158.99 \frac{L}{\text{barrel}}} \right) \times 69.1 \text{M} \text{toe} \times 4200 \frac{\text{MJ}}{\text{toe}} \times 0.6989 \frac{€}{\text{MJ}} = 38619 \text{ million } €
\]

Equation 17: Calculations on total cost for biomass.

\[
\text{Need for biomass} \times \text{Energy content in energy crops} \times \text{Price of biomass} = 139591449288 \text{ kg} \times 18.05 \frac{\text{MJ}}{\text{kg}} \times 200 \frac{€}{\text{MWh}} \times 0.000278 \frac{\text{MWh}}{\text{MJ}} \times 10^6 = 139979 \text{ million } €
\]

Scenario 1

Equation 18: Calculations on future crude oil price.

\[
\frac{\text{Total cost for biomass} \times \text{Energy content in oil}}{\text{Crude oil use in chemical industry}} = \frac{139979 \text{ million } € \times 37 \frac{\text{MJ}}{L} \times 158.99 \text{L/barrel}}{42000 \frac{\text{MJ}}{\text{toe}} \times 69.1 \text{M} \text{toe}} = 284 \text{ €/barrel}
\]

Equation 19: Increase in percentage.

\[
\frac{284 \text{ €/barrel}}{78 \text{ €/barrel}} = 362\%
\]

Scenario 2

Equation 20: Calculations on revenue from sold lignin.

\[
\frac{\text{Need for biomass} \times \text{Energy content energy crops} \times \text{Fraction of lignin} \times \text{Hard coal average price}}{\text{Tonne of coal equivalent}} = \frac{139591449288 \text{ kg} \times 18.05 \frac{\text{MJ}}{\text{kg}} \times 33\%}{10^4} \times \frac{80.95 \frac{€}{tce}}{29.31 \frac{€}{tce}} = 2320 \text{ million } €
\]

Equation 21: Calculations on future crude oil price if lignin is sold.

\[
\frac{(\text{Total cost for biomass} - \text{Revenue from sold lignin}) \times \text{Energy content oil}}{\text{Crude oil use in chemical industry}}
\]
Scenario 3

Equation 22: Calculations on avoided gCO₂.

\[
\text{Energy coming from crude oil} \times \text{Fraction of cellulose in biomass} \times \text{CO}_2\text{emissions} =
\]

\[
2815.13 \text{ PJ} \times 67\% \times 94.6 \frac{\text{gCO}_2}{\text{MJ}} \times 10^3 = 178428823 \text{ tCO}_2
\]

Equation 23: Calculations on avoided emissions rights.

\[
\frac{\text{Avoided emissions} \times \text{Emission rights spot price}}{178428823 \text{ tCO}_2 \times 13.33 \text{ €/tCO}_2 \times 10^6} = 2378 \text{ million €}
\]

Equation 24: Calculations on future crude oil price if lignin is sold and emission permits are avoided.

\[
\frac{\text{Total cost for biomass} \times \text{Energy content oil}}{\text{Crude oil use in chemical industry}} =
\]

\[
\frac{(139979 - 2320 \text{ million €}) \times 37 \frac{\text{MJ}}{L} \times 158.99\text{L/barrel}}{42000 \frac{\text{MJ}}{\text{toe}} \times 69.1\text{Mtoe}} = 274 \text{ €/barrel}
\]