Deliverable D3.1

Report on identified biomass conversion chains, their general socio-economic impacts and the translation of criteria into practically applicable indicators and thresholds

WP 3

July 2011
Global-Bio-Pact website: www.globalbiopact.eu
INTRODUCTION TO DELIVERABLE D3.1

1.1 Introduction

Work Package 3 of the Global-Bio-Pact project covers socio-economic impacts of biofuel/bioproduct conversion chains. The objective of this WP is to assess the sustainability of existing and new biofuel/bioproduct conversion chains with respect to its economic and social impacts. Thereby, emphasis will be put on industrial conversion chains in comparison with small-scale use of biomass. Close cooperation with WP2 will be maintained in order to assess the economic and social impacts of the whole life cycle of biofuels/bioproducts.

Contrary to other renewable energy sources like wind, solar and hydro-energy, biomass can be used for multiple applications like food, feed, biofuels and bio-products in addition to heat and power generation. Many first generation biofuels like palm oil, soy, sunflower and rapeseed for diesel generation and sugar cane and beet for ethanol production require considerable quantities of agricultural land. The last decade much attention has been paid to the environmental impacts of these crops, which has among others resulted in European sustainability criteria as described in the Renewable Energy Directive (RED) (2009/28/EC 2009). Within the Global-Bio-Pact project we are interested in the future outlook of our increasingly bio-based economy. Will future bioenergy conversion chains, like ‘biorefineries’ or ‘second generation biofuel production’ imply more sustainable practises? How can we guarantee the sustainability of these biofuels, especially socio-economic sustainability issues that are not covered by the RED. Furthermore it is observed that while current sustainability schemes pay much attention to biofuel production, its conversion seems to attract less attention.

In the Description of Work Deliverable D3.1 is described as a report on (1) identified biomass conversion chains, (2) their general socio-economic impacts and (3) the translation of criteria into practically applicable indicators and thresholds.

For practical reasons, these three topics have been addressed in two separate reports and one common instruction document developed in cooperation with WP2 covering socio-economic impacts of biomass production:

- Report Task 3.1 Identification of current and future industrial and small-scale conversion chains
- Report Task 3.2 Identification of socio-economic impacts of conversion chains
- Template for Global-Bio-Pact case studies with detailed instructions.

The report Task 3.1 and 3.2 are described below and attached as Annex 1 and Annex 2.
1.2 Report task 3.1 Identification of current and future industrial and small-scale conversion chains

This report provides an excellent overview of current and future biomass conversion technologies, with special focus on the technologies described in the case study i.e. sugar cane, jatropha, palm oil, and soy based biofuels, as well as on the different types of biorefineries. Figure 1 shows a dazzling number of possible biorefinery options.

![Figure 1 Schematic overview flow chart: biomass to products (Kamm, Gruber et al. 2006)](image)

For lignocellulosic biomass several pathways are described, using a systematic approach distinguishing between biochemical, thermochemical and hybrid pathways.

The following main routes exist for the processing of lignocellulosic biomass:

- A *biochemical pathway*, in which enzymes and other micro-organisms are used to convert cellulose and hemicellulose components of the feedstocks into sugars prior to their fermentation to produce ethanol and/or other products. In addition the lignin fraction of the biomass can be recovered and used for biochemical production.

- A *thermochemical pathway*, where pyrolysis/gasification technologies produce a synthetic gas from which a wide range of fuels and chemicals can be produced.

- A *hybrid pathway* when a combination of thermochemical and biochemical treatments is applied to one fraction.

- With elements of both main routes another pathway can be described, the ‘two platform biorefinery’, in which the cellulose and hemicellulose fractions are biochemically upgraded (the sugar platform), and the lignin fraction upgrades in a thermochemical way (syngas platform).

Also the status of the different pathways is discussed. For instance, Figure 2 shows that a large part of the small plants are operational, while most of the medium and large scale plants are planned but not yet operational. This clearly shows that most lignocellulosic biorefineries are still in the pilot and demonstration scale.
From the described technologies, the BTG-pyrolysis oil and the Lignol process have been selected for further analysis in the case study on the socio-economic impacts of large scale 2\textsuperscript{nd} generation conversion technologies.

1.3 Report Task 3.2 Identification of socio-economic impacts of conversion chains

Each activity that takes place in a biomass conversion chain as well as each input and output has impacts. Raw materials, labour and capital are the classic ingredients needed for a factory to operate. Technology could be added as a fourth factor that is materialised in capital goods (equipment/hardware, information technology, etc.) and embedded in humans (technical and organisational skills, etc.). The activities in the conversion chain result in various outputs such as final products, jobs, salaries, profits, but also emissions, waste, transport movements, etc. See Figure 3.

The biomass conversion chain (its inputs, outputs and activities) will have various impacts such as socio-economic, fiscal, environmental, and traffic impacts. The impacts can take place at various levels:

- Production unit level
- Community level
- Regional level
- National level
- International level.
Various methods have been developed to assess and quantify the impacts of planned interventions (policies, programs, plans, projects), such as:

- Socio-economic impact assessment (SEIA)
- Environmental impact assessment (EIA)
- Strategic environmental assessment (SEA)
- Social impact assessment (SIA)
- Development impact assessment / sustainable development
- Fiscal impact analysis
- Traffic impact analysis

Within the Global-Bio-Pact project, the socio-economic impacts of biofuel/bioproduct conversion chains are the main topic of interest. Therefore, the socio-economic impact assessment (SEIA) is expected to be the most relevant assessment method and described in further detail in the Task 3.2 report. Furthermore, a general overview of possible impacts is provided, as well as other lists of impacts that could be used as checklists when preparing a selection of relevant impacts for a particular project.

1.4 The translation of criteria into practically applicable indicators and thresholds

The translation of criteria into practically applicable indicators and thresholds is one of the main topics of the Global-Bio-Pact project as a whole. While biomass certification systems are generally used for biomass production, feasibility studies, environmental and socio-economic impact assessments are often used to check the sustainability of biomass conversion plants. There are some particular differences between the two approaches that have to be taken into account.

Biomass certification schemes measure whether the normative sustainability criteria are met by the use of indicators for compliance. The selection of the accurate and precise indicators is very relevant. Project operators are responsible for data collection on indicators of compliance, which should be therefore not be a too complicated or time consuming process.
Impact assessments are systematic processes to identify, predict and evaluate the effects of proposed actions and projects. These assessments are applied prior to major decisions and commitments being made, while biomass certification generally takes place after project implementation. Impact assessments contain in-depth analyses and are performed by specialists. The expected impacts should be described in a transparent and understandable way to the project developer. The project developer - and possibly government bodies and other involved stakeholders - subsequently decide whether mitigation measures are needed or, in the worst case, whether the project can proceed at all.

<table>
<thead>
<tr>
<th>Issue</th>
<th>Certification systems</th>
<th>Impact assessments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timing</td>
<td>Carried out after project implementation</td>
<td>Carried out before project implementation</td>
</tr>
<tr>
<td>Depth</td>
<td>Project operators must provide proof that criteria are met.</td>
<td>Qualified experts perform in-depth assessments</td>
</tr>
<tr>
<td>Appreciation</td>
<td>Project must meet pre-defined criteria that are determined in advance and integrated in the certification system.</td>
<td>The assessments result in an overview of possible impacts. The project developer and stakeholders must decide whether these impacts are acceptable.</td>
</tr>
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</table>

The described differences between the two instruments might be a bit exaggerated. Impact assessments will be implicitly normative and certification systems might contain indicators that are descriptive rather than normative. Certification schemes and impact assessments can also complement each other. An interesting example is the certification scheme of the Roundtable on Sustainable Biofuels (see www.rsb.org). It requests participating operators to perform a screening exercise to determine whether assessments like an Environmental and Social Impact Assessment are required. A special RSB Screening Tool (RSB 2011) is developed for this purpose. In case biofuel operations will have significant impacts, as measured during the screening exercise, a social impact assessment process shall be carried out. RSB provides further guidance on how to carry out these impact assessments. This could be an interesting way to address the relevant socio-economic issues in more depth while using a biomass certification scheme.

In the case studies of the Global-Bio-Pact project, data will be collected on the various socio-economic impacts. The data collection is not based on a limited pre-defined set of indicators and thresholds, in contrary, the case study partners are encouraged to identify and describe any relevant issue. In order to structure the process, the WP2 and WP3 leaders and the coordinator have developed a Template for Global-Bio-Pact case studies with detailed instructions on the collection of the main impacts, like economics, employment generation, working conditions, health issues, food issues, land use competition and conflicts, and gender issues. Where appropriate, reference is made to the task 3.2 report on identification of socio-economic impacts of conversion chains. Since this template for case studies is developed for internal use by project participants, it is not attached to this D3.1 report.
REFERENCES


ANNEX
A. REPORT TASK 3.1 IDENTIFICATION OF CURRENT AND FUTURE INDUSTRIAL AND SMALL-SCALE CONVERSION CHAINS
Identification of current and future industrial and small scale conversion chains

WP 3 – Task 3.1

July 2011
The Global-Bio-Pact project (Global Assessment of Biomass and Bioproduct Impacts on Socio-economics and Sustainability) is supported by the European Commission in the 7th Framework Programme for Research and Technological Development (2007-2013). The sole responsibility for the content of this report lies with the authors. It does not represent the opinion of the Community. The European Commission is not responsible for any use that may be made of the information contained therein. The Global-Bio-Pact project duration is February 2010 to January 2013 (Contract Number: 245085).

Global-Bio-Pact website: www.globalbiopact.eu
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1 INTRODUCTION

In this report current and future industrial and small-scale conversion chains are identified for a number of biomass feedstock. The report corresponds to task 3.1 and as part of deliverable D 3.1 of the Global-Bio-Pact project (www.globalbiopact.eu).

The following approach is used:

- Biomass can be used for the production of: (1) food and feed, (2) bioproducts, (3) biofuels, and/or (4) energy (heat and electricity). This paper covers conversion chains for the production of biofuels and/or bioproducts, possibly in combination with energy, food and feed production. However, conversion chains solely directed to the production of food and feed fall outside the scope of this report.
- Depending on the structure and composition of the biomass (oil, starch, sugar, lignocellulose) different conversion routes can be used to produce energy (heat, electricity), biofuels, and/or bioproducts. Therefore, a distinction is made between conversion chains based on (1) oil containing biomass, (2) sugar and starch biomass, and (3) lignocellulosic biomass.
- Current biomass conversion technologies are described in chapter 2, covering mainly commercially available technologies. For each conversion technology the (expected) commercial scale of production is described.
- The distinction between small and industrial scale has been made according the following working definition (based on BTG’s experience with biomass conversion chains)\(^1\):
  - micro (household) scale: < 1 MW\(_{\text{th input}}\), < 0.25 MW\(_{\text{e output}}\) and/or < 500 tonnes product
  - small scale: 1-20 MW\(_{\text{th input}}\), 1-5 MW\(_{\text{e output}}\) and/or < 0.5-10 ktonnes product
  - medium industrial scale: 20-200 MW\(_{\text{th input}}\), 5-50 MW\(_{\text{e output}}\), and/or 10-100 kton product
  - large industrial scale: >200 MW\(_{\text{th input}}\), >50 MW\(_{\text{e output}}\), and/or >100 kton product
- Future conversion chains are described using the concept of biorefineries, thereby covering a whole range of integrated biofuel, bioproduct and bioenergy production installations. Different types of biorefineries are described in chapter 3.
- In chapter 4 relevant conversion chains are selected for further analysis in the case studies and in task 3.2 of the Global-Bio-Pact project.

The report has the following purposes:

- Providing the reader with an overview of current and future conversion technologies and their scale of application.
- Offer background information for the selection of the case study on ‘Socio-economic impacts of large-scale second generation conversion technologies in EU/North America’.

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\(^1\) These capacities are indicative and roughly comparable; for energy applications the electric or thermal capacity can be used; for liquid and solid fuels the capacity can be expressed in kilo tonnes per year. Example: 10,000 tonnes biomass with 15 GJ/ton corresponds to 20 MW\(_{\text{th input}}\) or 5 MW\(_{\text{e output}}\) at an electric efficiency of 25% and 7,500 production hours per year.
• Select current and future conversion chains for further analysis in task 3.2 of the Global-Bio-Pact project.
2 CURRENT BIOMASS CONVERSION CHAINS

2.1 Classification of biomass resources

Biomass consists of a large variety of wastes, residues and crops that can be used for the production of solid, liquid and gaseous energy carriers and products.

Biomass can be defined as ‘the biodegradable fraction of products, waste and residues from agriculture (including vegetal and animal substances), forestry and related industries, as well as the biodegradable fraction of industrial and municipal waste’ (2001/77/EC 2001).

Biomass can be classified according to its sector of origin, like forestry, agricultural and waste sectors; or to the way it is released: main crops, primary harvest residues, secondary processing residues and tertiary residues after end use. In this paper, the following classification based on the composition of the biomass will be used:

- Oil containing biomass
- Sugar and starch biomass
- Lignocellulosic biomass
  - woody biomass
  - grassy biomass

**Oil containing biomass**

Oil containing biomass includes common agricultural crops like rape, sunflower, and soy, as well as Jatropha, an oil-containing non-food crop that can survive on poor soils under dry conditions. Furthermore, used frying oil, animal oils and fats and fat containing algae can be added to the list. The oils have a high energy content of 32-40 GJ/ton. Traditionally vegetable oils are mainly used for food applications. Oils can also be used as input material for biodiesel production, biogas production and direct combustion.

**Sugar and starch biomass**

Sugar and starch biomass includes agricultural commodities like sugar beet, sugar cane, corn, wheat, potatoes, etc. Sugar and starch crops have an energy content of 17-18 GJ/ton. Their main application is in food, but technical applications like glue, adhesive, pharmaceutical applications etc. are common as well. Sugar and starch are also commonly used for ethanol production and derived chemicals.

**Lignocellulosic biomass**

Lignocellulosic biomass refers to plant biomass that is composed of cellulose, hemicellulose, and lignin. Lignocellulosic biomass can be grouped into four main categories:

- wood residues (including sawmill and paper mill discards),
- agricultural residues (including corn stover and sugarcane bagasse),
- dedicated energy crops (including wood plantations and grasses), and
• end-of-life biomass waste (including demolition wood and the organic fraction of municipal waste).

Lignocellulosic biomass is expected to play a key role as cheap input for new conversion technologies for the production of liquid biofuels and bioproducts. It has three primary components: cellulose, hemicellulose and lignin.
• Cellulose has a strong molecular structure made from long chains of glucose molecules.
• Hemicellulose is a relatively amorphous component that is easier to break down with chemicals and/or heat than cellulose.
• Lignin is essentially the glue that provides the overall rigidity to the structure of plants and trees (trees typically have more lignin, which makes them able to grow taller than grasses).

For each main type of biomass the main conversion chains are described for the production of energy, fuels and chemicals. A distinction will be made between small, medium and large scale conversion technologies.

2.2 Biomass conversion chains for oil crops

Main biomass conversion chains for oil biomass are biodiesel and pure plant oil production. They are described in more detail in the following two chapters.

2.2.1 Biodiesel production and application

Figure 1 shows the main inputs and outputs of the biodiesel production process, based on the most common biodiesel production technology called "trans-esterification". In this process, oils and fats are combined with mono-alcohol in the presence of base catalyst to form fatty acid esters (biodiesel).
Animal and plant fats and oils are typically made of triglycerides which are esters containing three free fatty acids and the trihydric alcohol, glycerol. In the transesterification process, the alcohol is deprotonated with a base to make it a stronger nucleophile. Commonly, ethanol or methanol is used. The reaction has no other inputs than the triglyceride and the alcohol. Normally, without external input, this reaction will be either very slow or will not happen at all. Heat, as well as an acid or base are used to help the reaction proceed more quickly. It is important to note that the acid or base are not consumed by the transesterification reaction, thus they are not reactants but catalysts. Biodiesel has to be filtered and purified to meet fuel grade specifications required for use in diesel engines.

More detailed descriptions of the biodiesel production process can be found in various publications, like (Mittelbach and Remschmidt 2006), (Schober and Mittelbach 2009), (Knothe, Gerpen et al. 2005), and on websites like www.top-biofuel.org. It is possible to replace fossil diesel by 100% pure biodiesel. However, biodiesel is mainly used for blending with ordinary diesel to meet targets as set in the frame renewable energy policies in many countries.

Biodiesel factories form medium to large scale industrial conversion chains. The capacities can vary between 10-500 ktonnes/year, most of them larger than 30 ktonnes/year. The average plant capacity of biodiesel plants in Europe is 131 ktonnes/year.

2 Based on FACTBOX-Biodiesel plants across Europe (situation June 2007)
http://uk.reuters.com/article/idUKL3037791620070621
Practically all biodiesel is used for transport purposes; most of it is blended with fossil diesel, part of it is sold as 100% biodiesel.

### 2.2.2 Example: biodiesel from soy

Soy is a leguminosae plant native to East Asia, widely grown for its edible bean which has numerous uses. Soybeans contain about 18% oil and 43-49% protein. Fat-free soybean meal is a primary, low-cost, source of protein for human consumption and animal feeds. Soy vegetable oil is widely used as cooking oil and increasingly used for biodiesel production. The United States, Brazil, and Argentina are the world’s main soy producers. In 2010-2011, roughly 50 million tons of soy were produced in Argentina, which is about 20% of the world production. In 2010, about 2.5 million tons of soy biodiesel were produced in Argentine, while in 2005 biodiesel production was still marginal. This illustrates that although soy biodiesel is not Argentina’s main soy product, its importance has increased significantly. Soy oil conversion takes place in two main steps, (1) the production of crude soy oil and (2) further refining steps. Both steps are described below.

#### Production of crude soy oil

Soy oil is present in the soy seed. Harvesting is done mechanically by a combine, it cuts the soy plant with the soy beans and the combine separates the soy seeds from the soy beans. The soy seeds are (mechanically) pressed, and the resulting oil is filtered. Typically, an oil yield of 15-20% is possible. This crude soy oil needs some further purification and the product is refined soy oil. The refined soy oil is further processed to biodiesel and the press cake contains proteins mainly which are used as cattle feed. See Figure 2.

Steps in the crude soy oil production process:

1. The soy crushers buy soy beans from suppliers
2. A pre-treatment such as cleaning, drying and dehulling is necessary prior to oil extraction.
3. The soy beans are crushed and pressed and the crude soy oil is extracted.
4. The press cake is used as cattle feed.
Refining of crude soy oil

The crude soy oil still contains many oil-insoluble and oil-soluble impurities that need to be removed. The oil-insoluble materials are removed with filtration and the soluble materials are removed in five steps.

1. Degumming: the phosphatides are removed by adding water or acid.
2. Deacidification: in this step the development of rancid flavors of free fatty acids (FFA) are prevented. The content of these compounds in unrefined soy oil is between 0.6-6%.
3. Bleaching: in this step colorants are removed from the oil and this will extend the storage life of the oil. Usually, an absorbing substance such as bleaching earth, silica gel or activated carbon is added to remove the colorants.
4. Deodorization: in this step odorous substances are removed by steam distillation.
5. Dehydration: in the final step water impurities are removed. Traces of water may decrease the conversion of soy oil into biodiesel.

Biodiesel production

Although refined soy oil can be used as automotive fuels in their pure form, their use does require some serious engine modifications. Alternatively, the oil can be used for biodiesel production. Biodiesel can be used directly in most (modern) diesel engines, either in its pure form or blended with fossil diesel. It is produced through the process of transesterification, which is a reaction between refined soy oil (90%) and with methanol (10%). The reaction products are biodiesel (90%) and glycerine (10%).

Biodiesel can be produced from a range of vegetable and animal oils and fats. However, the quality of the biodiesel largely depends on the attributes of the raw material. Fuel stability, ignition behaviour, smoothness of combustion and (in colder climates) winter

3 Some (plastic) parts in the engine fuel system can be degraded by biodiesel over time. However, in response to the upcoming biodiesel market, today most manufacturers are using resistant materials.
operability are important fuel characteristics that are dependent on the composition of the oil. There is no single oil that scores perfectly on all attributes, but soy oil comes close to an ideal raw material (Körbitz 2008).

2.2.3 Pure plant oil production and application

Although the name „pure plant oil“ (PPO) refers to a vegetable origin like rapeseed, soybean etc., also oils from other resources, e.g. waste oil and animal fat, can be included in this category. Other publication also use the more generic term „straight vegetable oil“ (SVO). Like in biodiesel production the oil needs to be extracted followed by refining in order to remove undesirable substances, such as phosphatides, free fatty acids, waxes, tocopherols and colorants. In case of PPO no transesterification process takes place, which makes the PPO production process simpler than the biodiesel production process, and easier to apply on a smaller scale. For an extensive description of the PPO (and biodiesel) production process, please refer to (Rutz and Janssen 2007).

Already in 1900 PPO was used as a fuel to run diesel engines; however, its current application as transport fuel is more limited than biodiesel. Properties of pure plant oil (PPO) largely differ when compared to fossil diesel. For example the viscosity of PPO is much higher, especially at cooler temperatures. This property leads to technical challenges in winter running and when cold starting in conventional engines. Since PPO tends to gum up at colder temperatures, it has been difficult to blend it with conventional diesel fuel. On the positive site, the flashpoint of pure plant oil is significantly higher than that of normal diesel and is therefore particularly safe in storage and transport and easy to handle. Additionally, PPO is biodegradable in a short time in soil and waters and e.g. in Germany, it is not classified in any water hazard class (Rutz and Janssen 2007).
2.2.4 Example 1: Palm oil conversion chains

Palm oil conversion takes place in two main steps, (1) the production of crude palm oil and (2) further refining steps. Both steps are described below.

Production of crude palm oil
In palm oil mills fresh fruit bunches (FFB) are processed into crude palm oil (CPO), palm kernels and - in some of the plants – into palm kernel oil (PKO). Empty bunch ash (soda), fibres, shells and palm oil mill effluent (POME) are the side products of the process. See Figure 3 and the following description based on (Yew 2007). The steps in the process are numbered; the treatments of main by-products after each separation process are indicated with the number followed by “a” (e.g. 7a).

Steps in the crude palm oil production process:
1. The palm oil mill buys fresh fruit bunches (FFB) from their suppliers or harvests the FFBs from own plantations.
2. Filled lorries with FFB are weighed at the weigh-bridge section to ensure that the weight of FFB are match with the document.
3. After weighing, the FFB is graded at the grading section for quality control and also to determine the price of the FFB.
4. FFB are then unloaded into RAMP section.
5. The RAMP unloads FFB into cages that are charged into a sterilizer to cook the FFB. The overall sterilization takes around 85 minutes.

Figure 3 Crude oil milling process
Source: Plots Investment Group http://plotspalmoil.com/plantations.htm
6. The sterilized fruit bunches (SFB) are dropped into the thresher hopper and then into the feeder, where the feeder controls the feeding of SFB into the thresher. When the SFB enter the thresher, the thresher separates loose fruits from bunches.

7. Sterilized fruits from the thresher go through a sterilized fruit conveyor and elevator to the digesters. Digesters are filled with ¾ full of sterilized fruits, to soften the fruits before the press.

7a. Empty bunches and un-stripped bunches from the threshing go into a hard bunch crasher. At the hard bunch crasher, un-stripped bunches fall into the cages again for second round sterilization, while the empty fruit bunches (EFB) go into the incinerator for burning into ash; fibre from empty bunches can be used for fertilizer and for fuel at the boiler section.

8. Softened sterilized fruits are pressed to obtain crude oil and press cake.

8a. The press cakes consist of the mixed fibre and nuts and enter into a depericarper drum, where fibres are sucked up to fibre cyclone by cyclone fan, at the same time. Fibres are used as a fuel at the boiler section. Nuts from the depericarper drum go to the polishing drum by elevator to remove fibres from nuts. After the polishing drum, the de-stoner fan sucks fibres from nuts to the fibre cyclone, and nuts go into the nut silos and then into a nut cracker (ripple mill) to get cracked mixtures. These cracked mixtures will go through two winnowing systems (LTDS) for separation. Shells and kernels (cracked mixture) which are not 100% separated at the LTDS, conveyed to clay bath for further separation by using vibrating screen. Wet kernels go to kernel silos for drying, and these dried kernels will then store in kernel bunkers for dispatch, while shells to the boiler as fuel.

9. Crude oil flows into a crude oil gutter and then into a vibrating screen to separate dirt and sludge from crude oil. Dirt and sludge flow into conveyor back to digesters for oil recovery, and crude oil flows into crude oil tank then pumped to the oil room. In the crude oil tank, there is a skimmer to skim the pure oil at the upper layer of the crude oil and straight pump to the pure oil tank. Oil and sludge will underflow at the crude oil tank and pumped to clarifier 1 in the oil room. In the oil room, crude oil from the press is pumped into clarifier 1. Basically, the crude oil will form into 2 layers of oil, the pure oil at the upper layer while the oil plus sludge at the bottom layer. The coil in the clarifier heats up the crude oil to enable the pure oil to merge and float up easily to the top. Pure oil will overflow into a skimmer in the clarifier to the pure oil tank, and oil plus sludge and dirt will underflow to clarifier 2. The same concept applied in clarifier 2, where pure oil will overflow into a skimmer to reclaim oil tank and back to clarifier 1.

9a. The remaining oil, sludge and dirt will underflow to sludge tank then into the sludge siever to remove dirt and sands with bigger particle. Later, sludge and oil are pumped up to the decanter to remove sludge plus dirt, which are also called the decanter cake. Decanter cake used to make fertilizer. The oil and sludge from decanter are then flow into a decanter tank then pumped up to a baffle tank which located on top of the clarifier 2. From baffle tank to sludge separators, hot water added into sludge separators to separate the pure oil from the oil plus sludge. The sludge separators contain hundreds of bowls which piled up and function like a centrifuge. Sludge at the bottom is drained out and pure oil at the top is pumped into a fat oil tank then back into clarifier 1. Sludge and dirt at the bottom of the pure oil tank flows into the reclaim oil tank for recovery by going back into clarifier 1 again.
10. Pure oil from pure oil tank flows into a vacuum dryer with nozzles. These nozzles spray the hot oil into droplets with moisture in a vapour form, the vacuum sucks out the moisture and left the oil in a liquid form. Pure oil, commonly known as crude palm oil or CPO, which is produced from the mill is pumped into storage tanks for dispatch.

**Refining of crude palm oil**

Crude palm oil (CPO) is further processed by either physical or chemical refining to produce either refined, bleached and deodorised palm oil (RBDPO) or neutralised, bleached and deodorised palm oil (NBDPO). These are subjected to fractionation to obtain the respective liquid olein fraction and the solid stearin fraction. See Figure 4 and the description below based on (Hai 2002). Of the two processes, physical refining is the predominant approach adopted by the refineries as it is simpler, less capital intensive, more efficient and produces a lower effluent load.

*Physical or steam refining* begins with degumming when the CPO is treated with food grade phosphoric acid or citric acid to remove natural gums in the form of phosphatides, followed by bleaching with activated earth (Fuller’s Earth) under vacuum to remove colouring matters as well as to adsorb any metal ions. The treated oil is then heated to 240 - 260 °C under 2-6 mm Hg (MEOMA, 2002) for simultaneous de-acidification and deodorisation. The free fatty acids are stripped off by live steam and is recovered together with the entrained oil as palm fatty acid distillate. The steam distillation process also removes odours and off-flavors from the CPO (‘Deodorisation’). The oil is then cooled to 55°C before polishing. Refined Bleached Deodorised Palm Oil (RBDPO) is the basic oil product sold on the world's commodity markets.

In the *chemical refining process*, the free fatty acids present in CPO are removed by neutralisation with caustic soda (sodium hydroxide), the concentration of the latter being dependent on the quality of the CPO feedstock. This chemical reaction produces neutralised CPO and a soap stock; the latter is separated from the oil by a high-speed separator. The neutralised oil is subjected to earth bleaching to remove colour pigments and metal ions followed by deodorisation - steam distillation under vacuum to remove odoriferous matters such as aldehydes and ketones.
The refined oil contains triglycerides of various compositions and melting points, the main fractions being palm olein and palm stearin. These fractions can be separated by dry fractionation, detergent fractionation and solvent fractionation. Dry fractionation is commonly used whereby the refined oil is allowed to crystallise under controlled temperature and the resultant slurry is pumped through a membrane filter press to obtain the liquid olein fraction and the solid stearin portion. The olein could also be fractionated for a second time (‘double fractionation’) to produce a ‘super olein’ and a solid palm mid-fraction (PMF) which is the feedstock for production of specialty fats and other products.

After refining, palm oil possesses all the characteristics required for many uses without hydrogenation. By fractionating palm oil, its uses are further increased. Hydrogenation and inter-esterification increase its uses even further, especially in making specialised and sophisticated edible products. Its versatility makes it a suitable raw material for many applications both in edible and non-edible products. There are several classes of foodstuffs made wholly or in large part from oils and fats. Some of the products include:frying/cooking oils and fats, shortening, Vanaspati (vegetable ghee), margarine
and spreads, and imitation dairy products. Non-food applications of palm oil include soaps, oleochemicals like fatty acids, methyl esters, fatty alcohols, glycerine and epoxidised palm oil.

Table 1 Main applications of palm oil derivates. Source (Salmiah 2000)

| Fatty acids               | - Medium chain triglycerides for use in the flavour and fragrance industries  |
|                         | - Processing aids for rubber products                                      |
|                         | - Production of candles                                                    |
|                         | - Manufacture of cosmetic products from myristic, palmitic and stearic acids |
|                         | - Production of soaps via a neutralisation process                          |
|                         | - Production of non-metallic or non-sodium soaps                            |
| Fatty esters            | - Production of pure soap                                                  |
|                         | - Alpha-sulphonated methyl esters as active ingredients for washing and     |
|                         |   cleaning products (anionic surfactants)                                   |
|                         | - Palm-based methyl esters as a substitute for diesel fuel for vehicles     |
|                         |   and engines                                                               |
| Fatty alcohols          | - Fatty alcohol sulphates (anionic surfactants)                            |
|                         | - Fatty alcohol ethoxylates (nonionic surfactants)                         |
|                         | - Fatty alcohol ether sulphates (anionic surfactants)                      |
| Fatty nitrogen          | - Imidazolines with good surface active properties (rust prevention)        |
| compounds              | - Esterquats as softeners                                                   |
| Glycerol (Monoglycerides | - Wide range of applications such as a solvent for pharmaceutical products, |
| and Diglycerides)      |   humectants in cosmetics and tobacco, stabilisers, lubricants, antifreeze, |
|                         |   etc.                                                                      |

Biodiesel from palm oil

Like other vegetable oils, palm oil can be used for the production of biodiesel through transesterification into methyl esters. Depending on the design, palm oil based biodiesel plants use either accept crude palm oil (CPO) or refined, bleached and deodorised palm oil (RBDPO). Direct use of CPO requires an additional esterification step while RBDPO can undergo direct transesterification. See Figure 5. CPO’s treatment into a good quality biodiesel can take many shapes (using pure chemically-based materials, biocatalysts such as potassium-oxide and sodium-oxide or even microorganisms such as Pseudomonas fluorescens); subsequent transesterification usually involves the use of a sodium methoxide solution, or alternatively a high purity sodium hydroxide solid catalyst (Toh and Koh 2008).

Compared to other oil crops, palm oil is characterized by high amounts of medium-chain saturated and monounsaturated fatty acids. Therefore, the resulting palm biodiesel has a high filter plugging point and cloud point, and is referred to as “summer grade biodiesel”. A “winter grade biodiesel” can be produced by either a partial crystallization or a fractional distillation step to remove the C₁₆ fraction; the C₁₆ fraction can be further hydrogenated to produce a feedstock suitable for the manufacture of MES, a key ingredient in the detergent industry. Both summer and winter palm biodiesel grades are commercially available.
In recent years, high palm oil prices have stimulated the search for alternative biodiesel feedstock. Palm fatty acid distillate (PFAD) is a potentially valuable low-cost feedstock for biodiesel (Cheah, Toh et al. 2010). PFAD is generated in the fatty acid stripping and deodorization stages of the CPO refining process. Although the basic process for the conversion of high-acid oil feedstock to biodiesel is well known, it has been carried out mainly in small-scale batch-type processes. A breakthrough came in October 2009 with the successful operation of the world’s first continuous large-scale 200 MT/day PFAD biodiesel plant (in Sumatra, Indonesia). In this plant, owned by a large Asian based multinational palm oil group, fresh PFAD from the refineries is sent directly to the PFAD biodiesel plant for conversion to biodiesel. The biodiesel yield from this plant approaches 100%, and it fully meets EN (European Standards) specifications. After distillation, the PFAD biodiesel also passes the ASTM Cold Soak Filtration Test, introduced in 2008.
2.2.5 Example 2: Jatropha oil conversion chains

Jatropha oil is present in the jatropha seed kernel. After harvesting, the hulls of the seeds are removed in a de-hulling step. The kernels are (mechanically) pressed, and the resulting oil is filtered. Typically, 75-85% of the available oil is extracted. Both the oil and the by-products (press cake and hull) can be used for energy production or further processing. See Figure 6.

![Diagram of jatropha oil conversion chains](image)

Figure 6 Application of jatropha seeds Source: (Vis, Reumerman et al. 2009)

According to FACT Foundation (the Netherlands) the potential energy production from a jatropha plantation is made up of approx. 35% of energy from oil, 9% of energy in fruit shells and 35% of energy in press cake and oil residues. Thus it can be seen that adding energy use of fruit shells, press cake and oil residues can more than double the energy output, while improving the compost output as well. This makes addition of biogas into a pure jatropha oil model a very interesting new component. Biogas also has the advantage of returning the maximum of organic matter back to the land.

**Biodiesel production**

Although vegetable oils can be used as automotive fuels in their pure form, their use does require some serious engine modifications. Alternatively, the oil can be used for biodiesel production. Biodiesel can be used directly in most (modern) diesel engines\(^4\), either in its pure form or blended with fossil diesel. It is produced through the process of transesterification, which is a reaction between oil (90%) with methanol (10%). The reaction products are biodiesel (90%) and glycerine (10%).

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\(^4\) Some (plastic) parts in the engine fuel system can be degraded by biodiesel over time. However, in response to the upcoming biodiesel market, today most manufacturers are using resistant materials.
Biodiesel can be produced from a range of vegetable and animal oils and fats. However, the quality of the biodiesel largely depends on the attributes of the raw material. Fuel stability, ignition behaviour, smoothness of combustion and (in colder climates) winter operability are important fuel characteristics that are dependent on the composition of the oil. There is no single oil that scores perfectly on all attributes, but jatropha comes close to an ideal raw material (Körbitz 2008).

Box 2: Jatropha oil attributes

Jatropha oil is a vegetable oil that is, due to its toxicity, unfit for human or animal consumption. It is mainly composed of palmitic, stearic, oleic and linoleic fatty acids (Schultz 2008). Specific physical attributes:

- Lower heating value: 39.0 MJ/kg
- Density: 0.91 kg/litre
- Energy content of 1 litre jatropha oil: 35.49 MJ/litre
- Energy content of 1 litre jatropha oil: 9.86 kWh/litre
- Viscosity (at 25 °C): 50 cS
- Saponification Number: 192
- Iodine Value: 97

Electricity production

Like many vegetable oils, jatropha oil can be used directly for the generation of electricity in (modified) diesel engine – generator sets.

- Since the early 2000s, the use of vegetable oils for the production of combined heat and power (CHP) has increased steadily in Europe. Particularly in the last 2-3 years, many engine plants running on rapeseed and palm oil have been commissioned in for instance Germany, Italy and Belgium. Plant sizes range from below 100 kWe to several dozens of MW. Also, the use of jatropha is anticipated: a 9 MWe CHP plant on jatropha oil is now under construction in Belgium.

- In developing countries, the interest in using vegetable oils (particularly locally produced jatropha) for the production of electricity and mechanical power is growing, for example for rural applications. MFC Nyetaa (Mali-Folkecenter Nyetaa) is currently carrying out a variety of projects in 12 villages which will use Jatropha oil as fuel. 3 new projects combine jatropha oil with solar or biogas to have renewable energy hybrid systems. Another recent example is FACT Fuels, which has installed several generator sets in combination with their jatropha cultivation scheme in Mali. A further example is Winrock India, which installed several electricity generator sets running on jatropha oil for electrifying remote villages in Chhattisgarh.

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5 See: www.jatropha.de/p-o-engines/index.html
6 See http://www.winrockindia.org/act_proj_ene_prom_bio_1.htm,
• In a wider international context, application of vegetable oils in UNDP’s Multifunctional Platform (MFP) Programme is a possibility.

*Use as a cooking fuel*
Experiments with the use of vegetable oils as cooking fuels have led to the development of several types of cook stoves. An example of a modern, efficient stove is the Protos Plant Oil Cooker of B/S/H (Shiroff 2008). This cooker was developed in recent years for application in developing countries. At present, it is being marketed in a number of countries in Asia.

*Use for small-scale soap production*
Soap is a proven market for jatropha oils in rural Africa. Jatropha oil has good properties for making soap and soap production can be accomplished with minimal investment of supplies and equipment. The jatropha soap is often marketed as a medical soap and can be sold for a good price.

*Use of residues*
Finding applications for the valorisation of by-products of jatropha production is seen as one of the major challenges to improve the economics of jatropha growing.

• The jatropha press cake that is left after oil pressing has so far been seen as a fertiliser to be returned to the soil. It contains mainly proteins but due to its toxicity it cannot be used as animal feed. Experiments with detoxification, and breeding of non-toxic varieties, are on-going.

• Recent experiments with press cake indicate that it can be used for the production of biogas, through anaerobic digestion. The biogas can be used for energy production; the digester effluent, which still contains all the nutrients in the press cake, can be applied as a fertiliser on the land. MFC Nyetaa is currently carrying out a project in this area in Garalo, southern Mali. First studies by FACT show good economic results.

• Jatropha seed hulls constitute about one third of the seed weight. Experimental work in the Netherlands show that hull can be used as raw material for fibreboard production, wood-plastics composites or as a feedstock for the production of bioenergy of biofuels (Heeres 2008).

2.2.6 *Anaerobic digestion*
Oils and fats can be used as co-substrates for co-digestion with manure. For more information on anaerobic digestion please refer to section 2.4.6.

2.3 *Biomass conversion chains for sugar and starch biomass*
The main conversion chain for sugar and starch crops is ethanol production which includes a number of steps including fermentation. Sugar and starch crops can be used for anaerobic digestion as well.
2.3.1 Ethanol production from sugar and starch biomass

Ethanol is generally produced either synthetically from petrochemical feedstock (petroleum) or by microbial fermentation which is applicable to bioethanol production. The process for production of fuel bioethanol from biomass can be broken down to the following steps.

- Feedstock production: harvesting, reception, storage
- Physical pre-treatment: milling
- Saccarification: conversion of starch and cellulose into sugar
- Chemical treatment: dilution of the sugars with water and addition of yeast or other organisms
- Fermentation: production of ethanol in solution with water along with waste and by-products
- Distillation: separation of ethanol
- Dehydration: Removal of the remaining water by molecular sieves (anhydrous ethanol)
- Co-product preparation: Drying of the alcohol free stillage (mash) for high-value animal feed.

The steps in the feedstock-to-ethanol conversion process largely depend on the type of feedstock. Table 2 shows an overview of the ethanol production process for main feedstock types.
<table>
<thead>
<tr>
<th>Feedstock type</th>
<th>Feedstock</th>
<th>Harvest technique</th>
<th>Feedstock conversion to sugar</th>
<th>Process heat</th>
<th>Sugar conversion to Alcohol</th>
<th>Co-products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sugar crops</td>
<td>cane</td>
<td>Cane stalk cut, mostly taken from field</td>
<td>Sugars extracted through bagasse crushing, soaking, chemical treatment</td>
<td>Primarily from crushed cane (bagasse)</td>
<td>Fermentation and distillation of alcohol</td>
<td>Heat, electricity, molasses</td>
</tr>
<tr>
<td></td>
<td>sugar beet</td>
<td>Beets harvested, foliage left on the field</td>
<td>Sugar extraction</td>
<td>Typically from fossil fuel</td>
<td>Fermentation and distillation of alcohol</td>
<td>Animal feed, fertilizer</td>
</tr>
<tr>
<td>Starch crops</td>
<td>wheat</td>
<td>Starchy parts of plants harvested; stalks mostly left in the field</td>
<td>Starch separation, milling, conversion to sugars via enzyme application</td>
<td>Typically from fossil fuel</td>
<td>Fermentation and distillation of alcohol</td>
<td>Animal feed (e.g. distillers dried grains)</td>
</tr>
<tr>
<td></td>
<td>corn</td>
<td>Starchy parts of plants harvested; stalks mostly left in the field</td>
<td>Starch separation, milling, conversion to sugars via enzyme application</td>
<td>Typically from fossil fuel</td>
<td>Fermentation and distillation of alcohol</td>
<td>Animal feed (e.g. distillers dried grains), sweetener</td>
</tr>
<tr>
<td></td>
<td>potatoes</td>
<td>Potatoes harvested, foliage left on the field</td>
<td>Washing, mashing, cooking, starch separation, conversion to sugars via enzyme application</td>
<td>Typically from fossil fuel</td>
<td>Fermentation and distillation of alcohol</td>
<td>Animal feed, industrial use</td>
</tr>
<tr>
<td>Cellulosic crops</td>
<td>trees</td>
<td>Full plant harvested (above ground)</td>
<td>Cellulose conversion to sugar via saccharification (enzymatic hydrolysis)</td>
<td>Lignin and excess cellulose</td>
<td>Fermentation and distillation of alcohol</td>
<td>Heat, electricity, animal feed, bioplastics, etc.</td>
</tr>
<tr>
<td></td>
<td>grasses</td>
<td>Grasses cut with regrowth</td>
<td>Cellulose conversion to sugar via saccharification (enzymatic hydrolysis)</td>
<td>Lignin and excess cellulose</td>
<td>Fermentation and distillation of alcohol</td>
<td>Heat, electricity, animal feed, bioplastics, etc.</td>
</tr>
<tr>
<td>Waste biomass</td>
<td>Crop residues, forestry waste, municipal waste, mill waste</td>
<td>Collected, separated, cleaned to extract material high in cellulose</td>
<td>Cellulose conversion to sugar via saccharification (enzymatic hydrolysis)</td>
<td>Lignin and excess cellulose</td>
<td>Fermentation and distillation of alcohol</td>
<td>Heat, electricity, animal feed, bioplastics, etc.</td>
</tr>
</tbody>
</table>

Source: (Rutz and Janssen 2007)
Sugar-to-Ethanol Process
The simplest way to produce ethanol is the sugar-to-ethanol production. Sugar crops like sugar beets or cane contain substantial amounts of (six-carbon)\textsuperscript{7} sugars which can be fermented directly to ethanol. Although fungi, bacteria, and yeast microorganisms can be used for fermentation, the specific yeast Saccharomyces cerevisiae (Bakers’ yeast) is frequently used to ferment glucose to ethanol. Theoretically, 100 grams of glucose will produce 51.4 g of ethanol and 48.8 g of carbon dioxide.

In Brazil and in most tropical countries which produce ethanol, sugar cane is the most common feedstock for ethanol production. In these countries the costs of ethanol production from sugar cane are among the lowest of any biofuels.

Starch-to-Ethanol Process
A large portion of starch based bioethanol is produced from the starch component of grain crops, primarily corn and wheat in the US and wheat and barley in Europe. Starch molecules are made up of long chains of glucose molecules which have to be broken into simple glucose molecules (saccharification). Therefore starchy materials require a reaction of starch with water (hydrolysis). Typically hydrolysis is performed by mixing the starch with water to form slurry which is then stirred and heated to rupture the cell walls. During the heating cycle, specific enzymes are added, which break the chemical bonds. Organisms and enzymes for starch conversion and glucose fermentation on a commercial scale are readily available. The starch-to-ethanol process is shown in Figure 8. In conventional starch-to-ethanol processes, only the starchy part of the crop plant is used. Section 3.4.3 describes a biorefinery process using the whole starch crop.

\textsuperscript{7} Six-carbon sugars are much easier to convert to ethanol than five carbon sugars. See also chapter 3.5.1 on ethanol production from lignocellulosic biomass.
Ethanol - industrial versus small scale conversion chains

Although plant capacities vary widely ranging from 1.5 - 500 mln. litres/year, ethanol production is generally speaking a large scale business. The average capacity of European wheat/starch ethanol installations is 152 mln. litres/year; the average capacity of sugar/juice based ethanol production units is 132 mln litres/year. In the USA most of the ethanol production plants are corn based, their average capacity is 192 mln litres/year. The small scale initiatives (1.5 - 10 mln litres/year) mainly concern ethanol based on less common feedstock such as cheese whey, C-starch, seed corn, milo, brewery and beverage waste (as well as pulp and other lignocellulosic feedstock).

8 Based on 2010 data from ePURE http://www.epure.org/statistics.php?id=5
9 Based on 2010 data from EPAC Ethanol Producers And Consumers http://www.ethanolmt.org/plants.html
### 2.3.2 Example: ethanol production in Brazil and Costa Rica

In Brazil and Costa Rica, sugar cane is used as feedstock for ethanol. As of late 2007, sugarcane fields occupied about 7.8 million hectares in Brazil, or about 2% of all arable lands available in the country. The yearly yield of sugar cane in season 2008/09 was 569 mln tonnes of sugar cane, resulting in 31 mln tonnes of sugar, 18 mln m$^3$ of hydrous ethanol and 9.3 mln m$^3$ of anhydrous ethanol (UNICA 2011). This makes Brazil the number one producer of sugarcane in the world, followed by India, Thailand and Australia. Costa Rica has a much smaller yield of 4.1 mln tonnes of sugar cane in 2009 (FAOSTAT 2011). Main production regions are South-Central Brazil, where close to 90% of overall production is concentrated, and the Northeast, which accounts for the remainder. There are two harvests per year, which allows Brazil to produce sugar and ethanol year round for both the internal market and for export.

Sugar cane processing units can be divided into mills that produce only sugar, mills with distillery plants that produce sugar and ethanol, and independent distillery plants that produce ethanol only. If used in ethanol production only, each ton of ground sugarcane would result in 89 litres of hydrated or 85 litres of anhydrous ethanol today; if used exclusively in sugar production, each ton would yield 118 kg of sugar and 10 litres of ethanol from the molasses. It should be noticed, however, that under normal market operation the Brazilian average yield is 71 kg of sugar and 42 litres of alcohol per tonne of crushed sugarcane (Kaltner, Azevedo et al. 2005). Below the process scheme to produce ethanol (without sugar production) is illustrated.

The sugar cane is harvested by cutting the cane stalk mechanically or by hand; subsequently the stalks are taken from the field and transported to the processing facility. It takes two main steps from feedstock to ethanol. First, the sugars are extracted from the feedstock and second, the sugars are converted by fermentation and distillation into ethanol. See Figure 9 for all necessary steps.

Steps from feedstock to ethanol fuel:
1. The ethanol production facility receives sugar cane from the production fields.
2. It is first cleaned of dirt and any other extraneous material that may be present.
3. It is then chopped or shredded to change the harvested material to a fibrous material ready for sugar extraction.
4. During the milling process, the sugars (in form of juice) are squeezed out and the leftovers (fibrous materials) are called bagasse which is used as fuel for the processing facility.
5. The juice is then pasteurized, partially evaporated and centrifuged into syrup, or molasses.
6. The easiest method to produce ethanol from sugar is the sugar-to-ethanol production. Yeast is added to the syrup (sugar) and these organisms transform the six-carbon sugars (mainly glucose) to glucose. The formed products are ethanol and carbon dioxide.
7. The mixture of ethanol and yeast is centrifuged (distilled), the yeast is extracted from the ethanol and the end product is the biofuel ethanol.
Uses of residues

- Bagasse is the fibrous material that is leftover after the milling process. It contains a lot of energy which is burned in boilers for steam production and used for power and heat generation.
- Molasses: sugar cane molasses is a common ingredient in baking, often used in baked goods such as gingerbread cookies. Furthermore it can be used for ethanol production as well.

2.3.3 Anaerobic digestion

Sugar and starch biomass such as corn can be used as co-substrates for co-digestion with manure. For more information on anaerobic digestion please refer to section 2.4.6.

2.4 Biomass conversion chains for lignocellulosic biomass

A large number of technologies exist for the conversion of lignocellulosic biomass. Lignocellulosic biomass can be directly converted into energy by combustion. Other thermochemical processes such as gasification, pyrolysis, or carbonisation allow the production of intermediary products such as syngas, pyrolysis oil and charcoal that could be combusted for energy or used for the production of materials, fuels and or chemicals. Anaerobic digestion results in a methane rich biogas that can be regarded as an intermediate product as well. Ethanol production by fermentation of lignocellulosic biomass is challenging; a key characteristic of lignocellulosic biomass is that it is
naturally resistant to being broken down into their constituent parts, particularly in comparison to first-generation biomass feedstock such as sugar starch or vegetable oil.

The above mentioned conversion chains are described in the sections below, i.e. combustion, followed by the main thermochemical processes that produce intermediate products (gasification, pyrolysis and carbonisation) as well as the biochemical process of anaerobic digestion.

2.4.1 Combustion

Biomass combustion is the most common biomass conversion technology, applied on household and industrial levels since ancient times. Over the last decades, modern biomass combustion technologies have emerged like fully automated pellet boilers, co-firing, and efficient combined heat and power production from a large variety of biomass resources. Biomass combustion is used to generate heat, electricity or mechanical energy. After combustion only flue gas is generated, and - besides ash - no by products are produced. The following common combustion technologies can be distinguished on small (household), medium (larger buildings and industry) and large (power sector) scale:

- Biomass combustion on household level (heat only)
  - Charcoal stoves (cooking in developing countries)
  - Barbeque stoves (leisure use)
  - Fireplaces for wood blocks (low efficient, usually only for leisure use)
  - Wood block stoves and boilers
  - Wood pellet stoves and boilers.

- Biomass combustion for (non) residential buildings and industrial sector (heat and/or electricity)
  - Wood pellet boilers for houses and block heating
  - Wood chips boilers for block heating
  - Wood chips and straw boilers for district heating (and electricity production)
  - Boilers for industrial heat (and electricity) production using sawdust, wood chips, bark, black liquor, straw, etc.
  - Boilers for electricity production using wood chips, straw, etc.
  - Gas engines for the production of electricity and heat from biogas (from anaerobic (co)digestion of manure, waste water and industrial organic residues)
  - Gas engines for the production of electricity from landfill gas.

- Biomass combustion in the traditional power sector
  - Co-firing of biomass pellets (wood, straw, other) and torrefied biomass (wood) in coal fired power plants
  - Co-firing of liquid fuels (palm oil, fats) in power plants.

After the right pre-treatment (especially drying and sizing) practically all biomass can be combusted for energy generation. The most common biomass types for heat and electricity production are chipped, pelletised or carbonised wood. Compared to many other solid biomass types such as straw and other agricultural residues, wood has a
relatively low content of elements like chlorine and alkalies that can harm the conversion equipment during combustion. Nevertheless combustion of agricultural residues such as straw, bagasse, rice husks is also common practice.

For more information on combustion and co-combustion see for instance (Loo and Koppejan 2008); Also IEA task 32 on Combustion and Cofiring (http://www.ieabcc.nl/) is worth visiting. For more information on pellets see for instance the Pellet Handbook of (Obenberger and Thek 2010).

2.4.2 Gasification

Biomass gasification is an endothermic thermal conversion technology where a solid biomass fuel is converted into a combustible gas. A limited supply of oxygen, air, steam or a combination serves as the oxidising agent. The product gas consists of carbon monoxide, carbon dioxide, hydrogen, methane, trace amounts of higher hydrocarbons (ethane), water, nitrogen (with air as oxidant) and various contaminants, such as small char particles, ash, tars, higher hydrocarbons, alkalies, ammonia, acids, and the like. For firing the product gas in a boiler simple de-dusting of the product gas in a cyclone is sufficient, while for other applications such as gas engines, turbines and upgrading for biofuel/chemical production, various degrees of tar removal and extensive gas cleaning is needed.

Different types of gasifier designs can be distinguished:
- Fixed bed: downdraft, updraft, crossdraft
- Fluid bed: bubbling, circulating (CFB), twin-bed
- Entrained flow
- Stage gasification with physical separation of pyrolysis, oxidation and/or pyrolysis zones.

The following main applications can be distinguished:
- Gasification for heat production
- Fixed bed gasifiers for power or CHP production from biomass
- Co-firing gas from biomass in existing power plants
- Fluid bed gasification for power production
- Syntheses gas production for liquid (transportation) fuels. Synthesis gas differs from producer gas as it contains only CO and H₂ and is usually produced with oxygen as gasification agent.

**Industrial versus small scale applications**

Biomass gasification can be applied at very small scale (fixed bed at household level) up to extremely large scale as applied in refineries (entrained flow). See Figure 10.
At small scale, gasification is usually implemented for heat, or combined heat and power production (e.g. in rural electrification projects) up to a few MW. Small scale gasifiers combined with a gas turbine or engine can reach significantly higher electric efficiencies than small scale steam cycle combustion systems. At the medium to large scale, the market segment may change to co-firing, IGCC and/or syngas production for a wider variety of products. An entrained flow gasifier does not encounter large size restrictions and its capacity can easily reach several hundreds of MW, which could be combined with Fischer-Tropsch liquid production. Many gasification concepts are currently under demonstration and evaluation. An excellent source for more information is the Handbook Biomass Gasification (Knoef 2005); furthermore it is worthwhile to check IEA Task 33 on thermal gasification of biomass\(^\text{10}\).

\(^{10}\) http://www.gastechnology.org/webroot/app/xn/xd.aspx?it=enweb&xd=iea/homepage.xml
2.4.3 Pyrolysis

Fast pyrolysis is a process in which organic materials are rapidly heated to 450 - 600 °C in absence of air. Under these conditions, organic vapours, permanent gases and charcoal are produced. The vapours are condensed to pyrolysis oil. Typically, 50 - 75 wt.% of the feedstock is converted into pyrolysis oil.

Pyrolysis transforms difficult-to-handle biomass of different nature into a clean and uniform liquid, called pyrolysis oil. Not only woody biomass, but also agricultural residues such as empty fruit bunches can be converted into pyrolysis oil. Pyrolysis oil can be used for the production of renewable/sustainable energy and as a building block for the production of chemicals. Its energy density is four to five times higher than the original solid material, which offers important logistic advantages.

![Pyrolysis process diagram](image)

**Figure 11 BTG’s fast pyrolysis process. Source: BTG**

Pyrolysis oil production is in its demonstration phase. BTG works on the development of a 5 ton/hour input pyrolysis unit that can produce 26,000 ton/year pyrolysis oil in Hengelo, the Netherlands. In 2005, a 2 ton/hour plant was delivered to a client in Malaysia. Generally speaking pyrolysis oil production is viable at production rates of 20,000 - 50,000 pyrolysis oil per year, which requires an input of 30,000 - 75,000 tonnes of dry biomass, which is comparable with medium sized biomass plants. Pyrolysis oil knows various applications, from heat boilers for block, district and industrial heat to large scale biorefineries and co-refining in existing fossil refineries.

For more information one could refer to (Bridgwater, Czernik et al. 1999), IEA Task 34 on Biomass Pyrolysis ([http://www.pyne.co.uk](http://www.pyne.co.uk)) and company websites such as [www.btgworld.com](http://www.btgworld.com) and [www.btg-btl.com](http://www.btg-btl.com).
2.4.4 Carbonisation

Charcoal production from wood is the most common carbonisation technology, but also agro residues like cotton stalks can be carbonised and further upgraded to household fuels. The main applications for carbonised wood are cooking (in developing countries) and barbeque use. In some countries like Brazil, carbonised wood is used in steel making. Also specialities such as activated carbon for filters and medicine are made of charcoal.

![Charcoal production with Beehive kilns in Brazil. Source: BTG](image)

The production of charcoal can be performed in earth or metal kilns, or in more advanced installations that allow the full use of the energy rich vapours that are released. The scale of production ranges from a few m$^3$ per batch up till installations that can produce 1,500 tonnes/unit per year. Large wood plantations and charcoal companies can use multiple charcoal installations in parallel, but their total production capacities are usually well below 100,000 tonnes/year. The technologies are mature and found all over the world. The World Bank and FAO report regularly on the impacts of charcoal production for household use. For a charcoal handbook see for instance (Krishnan and Robinson 1999).

2.4.5 Torrefaction

Torrefaction is a partial carbonisation process at temperatures of 200-400°C, making the biomass ‘crispy’, comparable with roasting of coffee beans. The torrefied biomass is suitable for co-firing in coal-fired power plants. Compared to carbonised biomass, a higher percentage of the initial energy content of the biomass stays in the product. Torrefaction is a relatively novel concept; the first demonstration plants have been built in the Netherlands; the capacity of facilities is medium sized (60,000 ton/year) and seen the application of the torrified products in power plants, up-scaling to large production plants >100 kton could be anticipated. For more information see for instance
http://www.dutchtorrefactionassociation.eu/ and company websites such as http://www.topellenergy.com. IEA task 32 ‘biomass combustion and cofiring (http://www.ieabcc.nl/) also covers torrefaction.

2.4.6 Anaerobic digestion

Anaerobic digestion is the production of a methane rich biogas from wet biomass sources like manure, kitchen and garden waste, wastewater, etc., or from dedicated energy crops such as from corn silage. The biogas can be used for heat and power generation using gas engines, or upgraded for use in the natural gas grid.

Main anaerobic digestion approaches include:

- Manure digestion
- Manure co-digestion with additional materials (co-substrates)
- Digestion with dedicated energy crops
- Digestion of kitchen and garden waste
- Landfill gas extraction
- Anaerobic waste water treatment
- Micro digestion units for cooking (developing countries)

All approaches are mature and commercially available. In most cases the biogas is used for electricity production with a gas engine and generator set, resulting in a considerable electric efficiency of 30-40%. In case of manure digestion, the heat of the engine is used to keep the digester in an optimal temperature range and excess heat can be used for heating buildings, or drying. If no nearby heat users are available, as often the case with landfill gas extraction, the excess heat is lost. The biogas can be used for heat only applications as well, including cooking on biogas. Upgrading of biogas to ‘green gas’ for use in the natural gas grid is in its early commercial phase; it can also be used to fuel cars.

Figure 13 Small scale anaerobic digestion installation. Source BTG
Manure co-digestion allows the use of a wide range of co-substrates like sugar/starch biomass (corn, food waste), oil containing biomass (e.g. used frying oil), and lignocellulosic biomass such as grasses. Woody biomass is not suitable as a co-substrate for anaerobic digestion. The scale of most digestion technologies is in the range of 10 kW_e up till 5 MW_e, which is relatively small. Sometimes small biogas grids are used to collect biogas from multiple manure (co)digesters.

More information can be found in the German Biogas Praxis (Eder and Schulz 2006), which is known by experts. English speaking readers could for instance consult the Complete Biogas Handbook (House 2006) as well as websites such as http://www.biogas-info.co.uk or the Biogas Handbook of the BiG-East project (www.big-east.eu) (Al Seadi 2008).

2.5 Conclusion

The overview of current industrial and small-scale biomass conversion chains can be summarised in a Biomass Technology Combination Matrix as presented in Table 3. In principle, all biomass feedstock could be combusted for energy generation, but lignocellulosic biomass is preferred from an economic point of view.

<table>
<thead>
<tr>
<th>Combination Matrix</th>
<th>Oil containing biomass</th>
<th>Sugar and starch containing biomass</th>
<th>Lignocellulosic biomass</th>
</tr>
</thead>
<tbody>
<tr>
<td>Combustion</td>
<td>++</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Gasification, pyrolysis</td>
<td>-</td>
<td>-</td>
<td>++</td>
</tr>
<tr>
<td>Carbonisation, torrefaction</td>
<td>-</td>
<td>-</td>
<td>++</td>
</tr>
<tr>
<td>Anaerobic digestion</td>
<td>++</td>
<td>++</td>
<td>-</td>
</tr>
<tr>
<td>Fermentation – ethanol</td>
<td>-</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Transesterification - biodiesel</td>
<td>++</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Note: ++ = common combination; + = possible combination - = not common combination

Gasification, pyrolysis, carbonisation and torrefaction are strongly linked to solid lignocellulosic biomass sources. Anaerobic digesters can use a range of feedstock, except woody biomass. Obviously, transesterification and production of pure plant oil is linked to oil containing biomass; and fermentation to sugar and starch crops. Ethanol production through fermentation of lignocellulosic crops is challenging but could potentially result in high yield of ethanol from a wide range of feedstock.

The scale of production of the current conversion chains is summarised in Table 4. Basic physical properties of the feedstock and final products influence the common scale of production. Table 4 shows that ethanol production from sugar and starch crops and biodiesel production from oil have the largest scale of production, while conversion technologies for lignocellulosic feedstock such as combustion, gasification, pyrolysis, carbonisation and torrefaction are often applied in the range of the small and medium scale applications. It has to be kept in mind that raw biomass feedstock such as wood and agricultural residues have a higher bulk density and are available in smaller quantities than oil seeds and sugar/starch crops from large scale agriculture. This could affect the
expected scale and feedstock of future lignocellulosic feedstock refineries. Nevertheless the wood based paper and pulp industry operates at a large scale. The future possible size of lignocellulosic ethanol production could therefore be estimated by studying the paper industry.

<table>
<thead>
<tr>
<th>Table 4 Scale of production of current biomass conversion technologies¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electric output</td>
</tr>
<tr>
<td>Thermal input</td>
</tr>
<tr>
<td>Production capacity</td>
</tr>
<tr>
<td>&lt;10k ton/year</td>
</tr>
<tr>
<td>Combustion</td>
</tr>
<tr>
<td>Gasification</td>
</tr>
<tr>
<td>Pyrolysis</td>
</tr>
<tr>
<td>Carbonisation</td>
</tr>
<tr>
<td>Torrefaction</td>
</tr>
<tr>
<td>Anaerobic digestion</td>
</tr>
<tr>
<td>Fermentation (sugar/starch)</td>
</tr>
<tr>
<td>Transesterification</td>
</tr>
</tbody>
</table>

Note: ++ = common capacity range; + = possible capacity range but less common; - = not common capacity range

¹ The capacities are indicative; for energy applications the electric or thermal capacity can be used; for liquid and solid fuels the capacity can be expressed in kilotonnes per year. Example: 10,000 ton biomass with 15 GJ/ton corresponds to 20 MW\text{thermal input}, or 5 MW\text{electric output} at an electric efficiency of 25% and 7500 production hours per year.

For several reasons anaerobic digesters are usually small scale applications. First of all, only part of the incoming biomass can be converted into biogas which means that, compared to other conversion chains, relatively large input quantities of biomass are needed. Consequently, this leads to higher transport costs and a lower radius from which the (wet) biomass can be collected. Secondly, the voluminous biogas cannot be stored and needs to be used almost directly; and last but not least the electric efficiency of biogas combustion in gas engines is already high, which reduces the need to scale up.

All mentioned existing conversion chains can play a role in future conversion chains, mainly as part of different biorefinery concepts, which are presented in the next chapter.
3 BIOREFINERIES

In this chapter various conversion routes are presented that might have an important future role as 2nd generation biofuel and biomaterial production processes. These conversion routes are usually presented as ‘biorefineries’. In this chapter, definitions and classifications of biorefineries are presented, followed by a selection of biorefineries relevant for oil containing biomass, sugar/starch biomass and especially lignocellulosic biomass based conversion routes.

3.1 Definition of biorefineries

The term biorefinery is a popular concept nowadays and, depending on the type of activity and stakeholders involved, different definitions for biorefinery are currently used. (Ree and Annevelink 2007) present the following examples of definitions of biorefineries.

- A biorefinery is a facility that integrates biomass conversion processes and equipment to produce fuels, power, and value-added chemicals from biomass. The biorefinery concept is analogous to today’s petroleum refinery, which produces multiple fuels and products from petroleum (NREL 2007).
- A biorefinery is a cluster of bio-based industries producing chemicals, fuels, power, products, and materials.
- A biorefinery is an overall concept of a promising plant where biomass feed stocks are converted and extracted into a spectrum of valuable products (DOE 2007).
- Biorefinery is the separation of biomass into distinct components which can be individually brought to the market either directly after separation, or after further (biological, thermo chemical/chemical) treatment(s) (Elbersen 2003).
- Biorefining is the transfer of the efficiency and logic of fossil-based chemistry and substantial converting industry as well as the production of energy onto the biomass industry (Kamm, Gruber et al. 2006).
- Biorefineries are integrated bio-based industries, using a variety of technologies to produce chemicals, biofuels, food and feed ingredients, biomaterials (including fibres) and power from biomass raw materials (EU Eurofinery Euroview 2007).
- Addition of pure plant oil into traditional oil refineries (Shell 2007).
- Biorefinery is efficient use of the entire potential of raw materials and by-streams of the forest-based sector towards a broad range of high added-value products (by co-operation in between chains) ((Biorefinery Taskforce FTP 2007).
- A biorefinery is an integrated cluster of bio-industries, using a variety of different technologies to produce chemicals, biofuels, food ingredients, and power from biomass raw materials (Europabio, 2007).

Following (Ree and Annevelink 2007), the following definition as provided by the IEA Bioenergy Task 42 on Biorefineries, will be used in this report.

| Biorefinery is the sustainable processing of biomass into a spectrum of marketable products and energy. |

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3.2 Types of biorefineries

Just as with crude oil refineries, the feedstock - biomass - is separated and refined to produce various fuels and products in a biorefinery. Biorefineries are not new. An existing example is a modern sugar factory that can fine-tune the plant to produce either more bio-ethanol or more sugar. Another example is the starch hydrolysis plant, for the production of among others glucose, which can be used for the production of many chemicals and products such as ethanol, acetic acid, etc.

Biorefineries come in all shapes and sizes and there are many classifications possible. Among others, classifications can be based on:

- Raw material input (ligno-cellulosic biomass, aquatic biomass, etc.),
- Type of technology (thermo-chemical treatment, microbial degradation, etc.),
- Status of technology (conventional vs. advanced, 1st and 2nd generation), and
- Main (intermediate) product (syngas, sugar and lignin platform).

The following general classification, presented by (Kamm, Gruber et al. 2006) is often used nowadays:

- **Conventional biorefineries** - many existing industries make use of biorefinery concepts, like for instance in the sugar, starch, vegetable oils, feed, food, pulp and paper industries and the traditional biofuel industry.

- **The lignocellulosic feedstock biorefinery (LCF)** - where “nature-dry” biomass, such as wood and other cellulose-containing biomass is first separated into cellulose, hemicellulose and lignin, after which further processing can take place.

- **Thermo-chemical biorefineries** - where thermo-chemical technologies are used like pyrolysis or gasification to produce an intermediary product that can be refined thermo-chemically into a portfolio of value added products. A specific type of thermo-chemical biorefineries makes use of the existing fossil oil based petrochemical infrastructure.
Two platform biorefinery - Here two platforms are distinguished; the sugar platform and the syngas platform. The sugar platform is based on biochemical processes, and the syngas platform is based on thermo-chemical conversion.

Whole crop biorefinery - this concept uses raw materials such as cereals or maize. The straw may be utilised as in an LCF biorefinery, the seed may be converted to starch, or grinded to meal, followed by further processing.

Green biorefineries – where “nature wet” biomass such as green grass are converted to various products using microbial degradation.

Marine biorefinery - in which aquatic biomass (algae) are treated by cell disruption, product extraction and separation into, for instance, lipids and “algomass”.

Relevance of biorefinery types for different biomass types
In section 2.1 three main types of biomass were identified:
- oil containing biomass,
- sugar and starch biomass and
- lignocellulosic biomass (wood/straw and grass).

In the next sections biorefinery types relevant for these types of biomass are highlighted.

3.3 Biorefineries for oil containing biomass - the oleochemical industry
Vegetable and animal oils and fats are used in food, feed and other applications including cosmetics, shampoos, soaps, detergents, pharmaceutical products, paint, inks, industrial lubricants, plastics, waxes, candles, etc. (see for instance http://www.apag.org/oleo/index.htm). The oleochemical industry can be regarded as ‘conventional biorefineries’ responsible for the production of the above mentioned non-food non-feed applications.

The production of biodiesel from plant oils can also be regarded as part of the oleochemical industry. However (part of) the traditional oleochemical industry opposes the subsided use of oils and fats for transport fuels since it has led to considerable market distortions. In particular the drop in glycerine price induced by the increased biodiesel and subsequent glycerine production has hit the European oleochemical industry.

On-going research is focused on the use of oleochemical building blocks in order to increase the percentage of non-petroleum based components in various applications (Kamm, Gruber et al. 2006) (Vol II, p 295-300). It is however questionable how far these developments should be regarded as new biorefinery activities separate from the existing oleochemical industry.

For further information on the oleochemical sector see for instance the handbook of (Gunstone and Hamilton 2001) and (Kamm, Gruber et al. 2006).
3.4 Biorefineries for sugar and starch biomass

Upgrading of sugar and starch biomass is mainly performed in ‘conventional biorefineries’, but also new developments can be observed. In the next sections sugar-based and starch-based biorefineries are presented as well as the -starch crop based-whole crop biorefinery concept.

3.4.1 Sugar based biorefinery

Sugar mills operating in Brazil and other countries are large units functioning as autonomous industrial complexes. Their main products are sugar, fuel ethanol and bagasse. These systems can be regarded as ‘conventional biorefineries’ and have been described in section 2.3.1. The combined sugar and ethanol production, which is common in Brazil, could be called sugar-based biorefinery.

One interesting application is the production of poly-3-hydroxybutyrate (PHB), an environmentally degradable material belonging to the polyhydroxyalkanoates (PHA) family. PHB is a bioplastic that can be processed as a conventional thermoplastic in most industrial transformation processes including extrusion, injection, and thermopressing. It is fully biodegradable under the conditions of industrial waste composting sites (Kamm, Gruber et al. 2006). PHB is slightly more biodegradable than polylactic acid (PLA) - the most widely used bioplastic, and has a wider range of applications ranging from stiff packaging to highly elastic materials for coatings (Evans 2010).

PHB is produced in an aerobic fermentation process in which the sugar carbon source is converted into a biopolymer by means of the microorganism Ralstonia eutropha.

Figure 15 Sugar cane processing to sucrose, ethanol, by-products and new products. Source (Kamm, Gruber et al. 2006)
Recovery of PHB in biomass from fermentation involves several complex steps, for example microorganism cell breakdown, removal of impurities, and purification of the final product. For more information see (Kamm, Gruber et al. 2006) (Vol I, p 212-223).

Production of PHB can be integrated in a sugar and ethanol mill; the milling season lasts 180 days whereas PHB facilities could work all year round, meaning that the sugar input (medium quality standard sugar) needs to be stored. The required thermal energy and power could be supplied by the combustion of bagasse. The investment costs of a 10 kton PHB plant is estimated at 38 mln USD. The estimated production costs of integrating a PHB plant with an existing sugar mill are lower than those of a standalone PHB plant (Kamm, Gruber et al. 2006) (p 223).

No commercial sugar cane based PHB production plant was found yet. However, Telles, a joint venture of Metabolix and Archer Daniels Midland (ADM) has completed the initial phase of construction of a 50,000 ton/year corn based PHA manufacturing plant in Clinton, Iowa. This first ever large-scale PHA manufacturing facility will allow a range of PHA polymers to be produced and commercialized in large volume markets (Evans 2010).

3.4.2 Starch based biorefinery

Starch is a carbohydrate consisting of a large number of glucose units joined together by glycosidic bonds. It is the most important carbohydrate in the human diet and is contained in such staple foods as potatoes, wheat, maize (corn), rice, and tapioca (cassava).

**Potato starch**

Potato is the most important tuber used for industrial isolation of starch. Potato has the advantage that the extraction process is simpler than processes used for cereals, because it is not necessary to swell or mill them or prepare a dough. The disadvantages are the lower starch content of tubers and the large quantity of fruit water produced. Examples of (potato) starch based products are paper (mainly paper sizing), building additives (such as starch ethers for the dry mortar industry), adhesives (such as wallpaper, paper sack, envelop, billposting adhesives), textile (warp yarn sizing agents), etc. (see for instance www.avebe.com).

**Corn starch**

In the processing of corn based starch, a general distinction is made between traditional dry milling and wet milling.

**Dry milling** ethanol production plants work by grinding the grain to flour. The flour is then fermented to produce ethanol. Ethanol is separated off, and the residue (the “sillage”) is then dried to get DDGS “dried distillers grains with solubles”, which is used as cattle fodder. The CO₂ released during fermentation is captured and sold separately (see drawing).
In the wet milling process, grain is first soaked in water and sulphurous acid and 'steeped' for 1 to 2 days. It is then separated into its four basic components: starch, germ, fibre and protein. A metric tonne of corn grain yields, on average, 684 kg starch, 237 kg corn gluten feed, 45 kg gluten meal (60% protein) and 34 kg corn oil in a typical wet milling operation (Kamm, Gruber et al. 2006). Each component can be further processed to yield products. The greatest portion of the wet milled corn starch is converted to ethanol or syrup. The extent of hydrolysis determines whether a corn syrup (20-70 dextrose equivalent\(^{11}\) (DE)) or dextrose syrup (94-98 DE) is produced. Converting 95 or higher DE glucose syrup provides the refiner with three options. It can be refined as dextrose or 95 DE corn syrup, fermented to ethanol, or isomerised to high fructose corn sweetener (HFCS). In the USA the most widely used varieties of high-fructose corn syrup are: HFCS 55 (mostly used in soft drinks) and HFCS 42 (used in many foods and baked goods). In the USA, HFCS is among the sweeteners that have primarily replaced sucrose (table sugar).

The dry milling process is commonly used for the production of ethanol. It is cost effective, and capital expenditures are limited. Wet milling requires more investments, but the plant is more flexible, and a bigger portion of the grain is used to produce a variety of products. Although both processes can be regarded as ‘conventional biorefineries’, the development of the wet milling process with its flexible outputs is seen as exemplary for future biorefineries.

Furthermore, starch can be upgraded by:
- plasticization (co- and mix-polymerization, compounding with other polymers)
- chemical modification
  - etherification into carboxymethyl starch
  - esterification and re-esterification into fatty acid esters via acetic starch
  - splitting reductive amination into ethylenediamine, etc.,
  - hydrogenative splitting into sorbitol, ethylene glycol, propylene glycol, and glycerin
- bio-technological conversion into poly-3-hydroxybutyric acid (PHB).

Like in case of the oleochemical industry, the development of these new products, will be based on and developed largely by the existing starch processing industry.

\(^{11}\) The Dextrose Equivalent (DE) is a measure of the total dry reducing sugars in the syrup calculated as dextrose and expressed as a percentage of the total dry substance of the solution.
3.4.3 Whole crop milling biorefinery

When the upgrading of starch is combined with the upgrading of lignocellulosic feedstock (straw) at the same site, it is called ‘whole crop biorefinery’. The whole crop milling biorefinery is based on dry or wet milling of starch biomass such as rye, wheat, and maize, in which also the lignocellulosic fraction (straw) is applied in a lignocellulosic biorefinery:

- The grain will deliver starch, and derived products.
- The straw (a mixture of chaff, nodes, ears and leaves), represents a lignocellulosic feedstock, and may be further be processed in a lignocellulosic feedstock biorefinery, following biochemical and/or thermochemical pathways (see section 3.5).

The first step is the mechanical separation into a grain and straw fraction; where the portion of grain is approximately 20 wt% and the portion of straw is 80 wt%. Both streams will be further processed separately. Both fractions can be processed further to result in a portfolio of end products. See Figure 16.

![Figure 16 Potential products from a whole crop biorefinery (Kamm, Gruber et al. 2006)](image)

Many dry and wet milling plants can be regarded as ‘conventional biorefineries’, in which various starch based products are produced. It is expected that this existing large industry will develop toward the whole crop biorefinery concept. The starch part of the feedstock competes with food.
3.5 Biorefineries for wood/straw based lignocellulosic biomass

The following main routes exist for the processing of lignocellulosic biomass:

- A **biochemical pathway**, in which enzymes and other micro-organisms are used to convert cellulose and hemicellulose components of the feedstocks into sugars prior to their fermentation to produce ethanol and/or other products (see section 3.5.1). In addition the lignin fraction of the biomass can be recovered and used for biochemical production (see section 3.5.2).

- A **thermochemical pathway**, where pyrolysis/gasification technologies produce a synthetic gas from which a wide range of fuels and chemicals can be produced (see section 3.5.3).

- A **hybrid pathway** when a combination of thermochemical and biochemical treatments is applied to one fraction (section 3.5.4).

- With elements of both main routes another pathway can be described, the ‘two platform biorefinery’, in which the cellulose and hemicellulose fractions are biochemically upgraded (the sugar platform), and the lignin fraction upgraded in a thermochemical way (syngas platform) (see section 3.5.5).

**Status and scale of production**

An excellent overview of current biochemical, thermochemical and hybrid lignocellulosic biorefinery projects and their status can be found on (Bacovsky, Dallos et al. 2010), and also accessible through [http://biofuels.abc-energy.at/demoplants/projects/mapindex](http://biofuels.abc-energy.at/demoplants/projects/mapindex).

![Figure 17 Number of biorefinery project in certain capacity ranges sorted by technology. Source (Bacovsky, Dallos et al. 2010)](image-url)
(Bacovsky, Dallos et al. 2010) collected information on 66 projects, of which 37 were classified to use a biochemical pathway, 23 a thermochemical pathway and 6 a hybrid pathway. Figure 17 shows that most of these projects (45 out of 66) are small scale, 12 are medium scale and 9 large scale projects. Figure 18 shows that a large part of the small plants are operational, while most of the medium and large scale plants are planned but not yet operational. This clearly shows that the lignocellulosic biorefineries are in the pilot and demonstration scale.

Figure 18 Planned, under construction and operational lignocellulosic biorefinery projects. Source: (Bacovsky, Dallos et al. 2010)

3.5.1 Biochemical biorefinery - lignocellulosic ethanol production

Lignocellulosic feedstock biorefineries are based on the fractionation of lignocellulosic-rich biomass sources into the intermediate output streams cellulose, hemicellulose and lignin, which can then be further processed into a portfolio of bio-based end-products, materials, chemicals, fuels and power and/or heat. See Figure 19.
Although various possibilities exist, most research is focused on the production of ethanol from both the cellulose and hemicellulose fractions (this section) and the utilization of the lignin fraction (section 3.5.2).

**Ethanol production from cellulose and hemicellulose**
Lignocellulosic biomass consists of three main components: cellulose, hemicellulose and lignin. Producing ethanol from lignocellulosic biomass depends on converting the complex cellulosic and hemicellulosic carbohydrates into simple sugars which are fermented to ethanol by a variety of microorganisms. The cellulose contains hexose (C-6) sugars; the hemicellulose contains hexose (C-6) and pentose (C-5) sugars. Table 5 shows the carbohydrate composition of different lignocellulosic feedstock.

**Table 5 Carbohydrate composition of different lignocellulosic feedstock**

<table>
<thead>
<tr>
<th>Plant name</th>
<th>Glucan (%)</th>
<th>Mannan (%)</th>
<th>Galactan (%)</th>
<th>Xylan (%)</th>
<th>Arabinan (%)</th>
<th>Lignin (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>White wood</td>
<td>46.5</td>
<td>11.6</td>
<td>1.2</td>
<td>6.8</td>
<td>1.6</td>
<td>26.7</td>
</tr>
<tr>
<td>Douglas fir</td>
<td>43.46</td>
<td>10.76</td>
<td>4.66</td>
<td>2.77</td>
<td>2.67</td>
<td>31.3</td>
</tr>
<tr>
<td>Oak</td>
<td>40.63</td>
<td>1.97</td>
<td>1.22</td>
<td>19.19</td>
<td>0.36</td>
<td>23.91</td>
</tr>
<tr>
<td>Aspen</td>
<td>45.97</td>
<td>2.10</td>
<td>7.9</td>
<td>17.74</td>
<td>1.23</td>
<td>20.30</td>
</tr>
<tr>
<td>Maize cob</td>
<td>34.0</td>
<td>0.5</td>
<td>1.0</td>
<td>14.0</td>
<td>1.7</td>
<td>13.1</td>
</tr>
<tr>
<td>Wheat straw</td>
<td>37.0</td>
<td>0.3</td>
<td>6.5</td>
<td>18.9</td>
<td>5.6</td>
<td>13.6</td>
</tr>
<tr>
<td>Resulting</td>
<td>D-glucose</td>
<td>D-mannose</td>
<td>D-galactos</td>
<td>D-Xylose</td>
<td>L-arabinose</td>
<td></td>
</tr>
<tr>
<td>sugar type</td>
<td>(Hexose)</td>
<td>(Hexose)</td>
<td>(Hexose)</td>
<td>(Hexose)</td>
<td>(Pentose)</td>
<td></td>
</tr>
</tbody>
</table>

Source: based on (Kamm, Gruber et al. 2006), vol 2, p 109
Depending on the feedstock, next to the conversion of C-6 sugars, the conversion of C-5 sugars of the hemicellulose is essential to achieve high ethanol conversion rates. The biochemical conversion of lignocellulosic biomass into ethanol consists of three main steps:

1. pre-treatment: sizing, removal dirt, releasing cellulose from hemicellulose and lignin
2. (hemi) cellulase-mediated hydrolysis of (hemi)cellulose into sugars (C-5 and C-6)
3. fermentation of C-5 and C-6 sugars into ethanol.

These steps can be carried out with varying degrees of consolidation and are described below.

**Pretreatment**

Due to the strong bonds of the ligno-cellulose structure, pretreatment processes are generally severe and represent a significant cost element of the whole pathway. The aim of the pretreatment is to open up the cellular structure of the biomass by removal of the lignin fraction, so that the cellulose and hemicellulose can be exposed to enzymatic hydrolysis, and to avoid the release of other components toxic for the hydrolysis process. Woody biomass pretreatments that are under investigation include (1) one stage steam explosion, (2) two stage steam explosion in the presence of SO$_2$ or sulphuric acid, (3) ammonia fibre explosion (AFEX), (4) dilute mineral acid, and (5) organosolv delignification. Steam explosion under mildly acidic conditions is currently the state-of-the-art pretreatment technology. Laboratory testing has shown that the recovery of C-5 and C-6 sugars and lignin from softwood after steam-explosion pretreatment can be significantly lower than for agricultural residues or hardwoods such as poplar. This problem is one of the primary drivers for assessing alternative pretreatment methods for softwoods. More information on different pretreatment methods is summarised in (Sims and Taylor 2008) (p 46-50).

**Enzymatic hydrolysis**

Hydrolysis is a chemical process in which a certain molecule is split into two parts by the addition of a molecule of water, in this case the production of sugars from cellulose by either acid hydrolysis or enzymatic hydrolysis:

- **Acid hydrolysis** can be applied to convert cellulose to sugars. Using sulphuric acid to hydrolyse lignocellulosic fibres into sugars has not been very economical. Processes that hydrolyse fibres with dilute acid and pressure have tended to degrade too much of the hemicellulose sugars before they can be fermented into ethanol, causing low yields. The other option, using concentrated acid at low pressures, requires purchasing and then recycling expensive quantities of acid.

- **Enzymatic cellulose** hydrolysis processes are of interest because enzymes exclusively catalyse specific reactions. Therefore, different from acidic hydrolysis, there are no side reactions or by-productions and the hydrolysis can potentially be performed with yields approaching 100%. Enzymatic hydrolysis of conventional starchy feedstocks has already replaced acid hydrolysis in modern ethanol facilities.
Cellulase is a group of enzymes that cellulose converts to glucose. Efficient enzymatic hydrolysis of cellulose requires the synergistic activity of three groups of enzymes (Sims and Taylor 2008):

- Endo-β-1,4-glucanases are enzymes that hydrolyse accessible regions on cellulose chains and split the chains to provide sites for other enzymes to interact.
- Cellobiohydrolases act on these sites and begin to remove successive cellobiose units (i.e. two joined glucose units).
- β-glucosidase is a specialised enzyme that hydrolyses cellobiose to glucose.

Commercial cellulase preparations are typically deficient in β-glucosidase so require supplementing in order to relieve product inhibition caused by accumulation of cellobiose and other soluble sugars. This adds to the already substantial cost for enzymes. Substantial efforts are made to reduce the total enzyme costs, including enzyme production costs, pretreatment methods that reduce enzyme use and enzyme recycling.

**Fermentation**

Fermentation of glucose (C-6) to ethanol is not difficult provided there is an absence of inhibitory substances such as furfural, hydroxyl methyl furfural, or natural wood derived inhibitors such as resin acids. For more than 20 years research activities have been directed towards the development of improved micro-organisms for the fermentation of the pentose (C-5) sugars. For cost effective processing, such organisms must be able to co-ferment both glucose and pentose sugars together. Significant progress has been made in engineering such micro-organisms, and pentose fermentation has been achieved on ideal substrates. Their sensitivity to inhibitors and the production of unwanted by-products remain serious problems yet to be overcome if these systems are to become commercially viable. Most of the current lignocellulosic ethanol production units convert only the hexose sugars into ethanol.

**Status and scale of production**

Ethanol production from lignocellulosic biomass is in its demonstration phase. Whilst the costs for lignocellulosic feedstocks themselves are expected to be lower than for oil, sugar and starch crops, the difficulty of converting these feedstocks to liquid fuels means that the conversion technologies are prone to being more expensive than current conversion technologies. Several initiatives can be found in Europe and especially the USA. Because of the demonstration character, the plants are generally smaller than sugar/starch ethanol plants.

(Bacovsky, Dallos et al. 2010) provide a recent overview of (planned) demonstration units which have capacities of 50-500,000 tonnes/year. The information can also be accessed through a map on [http://biofuels.abc-energy.at/demoplants/projects/mapindex](http://biofuels.abc-energy.at/demoplants/projects/mapindex). For more information regarding the process please refer to (Kamm, Gruber et al. 2006). Also (Sims and Taylor 2008) provide an excellent overview of the status of a.o. lignocellulosic ethanol production.
3.5.2 Biochemical biorefinery – lignin utilization

Lignin represents a considerable proportion of the structural components of plants. It accounts for approximately 10-12% of some short annual plants and up to 30% and more for some coniferous trees. Most lignin in the chemical pulping industry is burned in chemical recovery boilers. Nevertheless, most experience with lignin recovery is build up by the chemical pulping industry. Significant commercial markets, totalling over one million tonnes per year, already exist for lignins that have been recovered from chemical pulp mills. The value of lignin in these markets is generally an order of magnitude higher than its fuel value. Furthermore lignin from some types of biorefineries will most probably have better performance characteristics and be of greater commercial value for its chemical properties than lignins from existing chemical pulping operations (Kamm, Gruber et al. 2006) (Vol 2, p 165).

Traditional lignin products from paper mills
In paper mills the cellulose is separated from the remaining hemicellulose and lignin fraction, forming black liquor in the kraft process or red/brown liquor in sulphite pulping, from which lignin fractions can be obtained. The main commercial lignin types are lignosulfonates from sulphite pulp mills, and - to a lesser extent - thiolignins from Kraft pulping processes.

Lignosulfonates are recovered from the spent pulping liquids (red or brown liquor) from sulfite pulping. The most widely used industrial process is the Howard process, in which 90–95% yields of calcium lignosulfonates are precipitated by adding of excess calcium hydroxide. Lignosulfonate can be used for road dust suppression, feed pellet binders and animal feed additives. More advanced applications of lignosulfonates, promoted by the larger mills, are its use as dispersants, emulsifiers and surfactants. Borregaard LignoTech produces 400,000 tons of lignosulfonates per year while the total market is below 800,000 tonnes/year. Because of the steadily declining number of sulphite pulp mills, the amount of available lignosulfonates has also declined (Kamm, Gruber et al. 2006) (Vol 2, p 173).

Today vast quantities of lignin are processed in the kraft pulping industry, probably more than 70 million tonnes per year, but of this amount more than 99% is burned in chemical recovery furnaces. The recovery of kraft lignin from black liquors is neither simple nor inexpensive. MeadWestvaco is the main producer of kraft lignins. The total production of kraft lignin is estimated to be below 100,000 tonnes/year (Kamm, Gruber et al. 2006) (Vol 2, p 170). Kraft lignin finds applications as rubber reinforcers, activated carbon, carbon black substitues, phenolic resin components, and as raw materials for the production of methylsulfonates (competing with lignosulfonates).

Lignin from biorefineries
The options for lignin recovery from biochemical biorefineries depends mainly on the type of pretreatment. The various fibre explosion processes open up the tight structure of the fibre to expose cellulose more for enzymatic attack. In the subsequent enzymatic hydrolysis, the presence of lignin reduces the efficiency of the process as the cellulase tends to bind with the lignin. Therefore, after the initial explosion stage, lignin extraction from the pre-treated material is often included, usually with hot ethanol or hot alkali.
The organosolv process combines pretreatment and (hemicellulose and) lignin removal in one step. It involves treatment of the raw lignocellulosic material with an aqueous organic solvent (frequently ethanol) at temperatures in the range of 180-200 °C, in which most of the hemicellulose and lignin is hydrolysed and dissolved in the liquor. Following washing of the remaining solid fibre, the residual lignin is a minor part of the finely disrupted material. The lignin is separated from the “black” liquor and other components can be extracted as well in further steps as shown in Figure 20.

Recent interest in environmentally friendly chemical pulping has encouraged the investigation and development of organosolv pulping. Various organosolv processes have been proposed and investigated over the last twenty-five years or so. These include the Alcell process, using ethanol; Acetosolv, using acetic acid; Formacell, using formic acid; pulping with phenol; and the Organocell process using methanol.

The Alcell process, developed by Repap Enterprises Inc, during 1987–1997, is perhaps the most commercially advanced of the ethanol-based organosolv pulping processes. The Alcell pulping process was operated in a pre-commercial demonstration plant at Repap’s Miramichi pulp and paper mill in New Brunswick, Canada. This plant, with a design capacity of 30 tonnes of pulp per day, operated intermittently from 1989 to 1996 and produced more than 3,700 tonnes of organosolv lignin. This lignin was in the form of a dry powder and was sold in supersacks containing approximately 800 kg each. All of the lignin produced at the plant was sold commercially during the time the plant operated, except for some minor portion that was used for internal study and development purposes. Unfortunately, with the financial collapse and consequent breakup of Repap in 1997, this plant was closed, but the technology has since been acquired by Lignol Innovations Corporation of Vancouver, Canada, and is now being commercialized as a biorefinery technology with the cellulose fraction being used for ethanol production instead of pulp (Kamm, Gruber et al. 2006) (Vol 2, p 179).
Lignin recovery is also addressed by other biorefinery process developers, including hybrid options, like lignin recovery from pyrolysis oil.

**Applications of lignin from biorefineries**

One of the most important applications of lignin is the replacement of phenol-formaldehyde (PF) resin applications. PF is produced by reacting phenol and formaldehyde in the presence of an acid or alkaline catalyst. Since phenol is now produced almost exclusively from benzene instead of from coking operations, increases in crude oil prices cause a significant rise in the cost of raw materials and a consequent increase in the prices for phenolic resins. The manner of substituting lignin for PF-resins can vary from a simple blending of dry powder lignin with dry powder phenolic resin to the use of organosolv lignin as a primary phenolic component during the manufacture of the resin. Table 6 shows the main present applications of PF in which lignin could play a role as a substitute. The degree of substitution varies and depends on the specific application and R&D efforts. Already in 1995, when Repap was preparing to construct a 450 tonne per day Alcell pulp mill at Atholville, New Brunswick, several studies were commissioned to identify marketing strategies for the 56,000 tonnes of organosolv lignin output from this mill. These studies clearly showed that several hundred thousand tonnes of lignin per year could be easily sold at that time for an average price close to the price of PF resins. Lignosulfonates and thiolignins have been marketed for incorporation into phenolic resins for some time, but they do not appear to have met with significant acceptance by the resin industry.

<table>
<thead>
<tr>
<th>PF- Application</th>
<th>Current market size (tonnes/year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Panel board adhesives</td>
<td>USA 550,000</td>
</tr>
<tr>
<td></td>
<td>World 1.2 mln</td>
</tr>
<tr>
<td>Thermoset resins for moulded products</td>
<td>USA 55,000</td>
</tr>
<tr>
<td></td>
<td>World 200,000</td>
</tr>
<tr>
<td>Friction materials</td>
<td>USA 27,000</td>
</tr>
<tr>
<td></td>
<td>World &gt; 120,000</td>
</tr>
<tr>
<td>Foundry resins</td>
<td>USA 55,000-60,000</td>
</tr>
<tr>
<td></td>
<td>World 150,000</td>
</tr>
<tr>
<td>Insulation materials</td>
<td>USA 110,000</td>
</tr>
<tr>
<td></td>
<td>World -</td>
</tr>
<tr>
<td>Decorative laminates</td>
<td>USA 100,000</td>
</tr>
<tr>
<td></td>
<td>World 300,000</td>
</tr>
</tbody>
</table>

\(^a\) North America

Lignin from biorefineries can also serve as an anti-oxidant, for instance in animal feed supplements, rubber products and lubricant industry. It could also be used in markets currently served by lignosulfonates, i.e. in concrete admixtures, dye dispersants, asphalt emulsifiers, agricultural applications, and as dispersants for herbicides, pesticides and fungicides. See (Kamm, Gruber et al. 2006) (Vol 2 p 181-199).
3.5.3 Thermochemical biorefinery

Most thermal biorefineries are based on a combination of gasification, gas cleaning and upgrading and fuel synthesis.

Gasification
In most thermochemical biorefineries dry lignocellulosic feedstock is subjected to a severe heat treatment in the presence of a controlled amount of air (or oxygen) so that gasification takes place. Compared to ‘ordinary’ gasification, the product gas or syngas is used for fuel/chemical production instead of conventional heat and/or power production. In some applications pyrolysis is used as a pretreatment before gasification. Gasification techniques applied are quite equally split between fluidised bed gasifiers and entrained flow gasifiers (Bacovsky, Dallos et al. 2010). Generally the fluidised bed gasifiers are built smaller than entrained flow gasifiers.

Gas cleaning and upgrading
The gas needs to be cleaned for the prevention of corrosion, erosion and deposits in the process lines as well as prevention of poisoning of catalysts. Typical components that need to be removed are dust, alkali metal compounds, tar, sulphur compounds, nitrogen and chloride compounds. After that the gas can be upgraded, for instance by a water-gas-shift reaction, gas reforming and removal of inert gas fractions (mainly CO₂).

Fuel synthesis
Starting from the syngas several fuel processing pathways are possible, such as Fischer-Tropsch synthesis, methanol synthesis, and production of synthetic natural gas (SNG). Ethanol production can be produced from producer gas following either a thermochemical or a microbiological pathway.

Fischer-Tropsch synthesis
Fischer-Tropsch synthesis is a catalysed chemical reaction in which carbon monoxide and hydrogen are produced from the gasification of a solid feedstock such as coal or woody biomass into a range of liquid hydrocarbons. Typical transition metal catalysts are based on iron and cobalt.

Useful reactions give alkanes \((2n+1)\ H_2 + nCO \rightarrow C_nH_{(2n+2)} + nH_2O\)

The process was developed in 1923 by Prof Franz Fischer and Dr. Hans Tropsch. In the 1940s 600,000 t/yr of liquid fuels were made from coal in Germany from coals. South Africa built three coal-to-liquid SASOL plants in the 1980s using the FT process to convert coal to gasoline and diesel. Shell has operated a commercial FT-diesel plant in Malaysia since 1993. It uses natural gas as a feedstock to produce primarily low sulphur diesel fuels and food-grade wax (Sims and Taylor 2008).

Similar conversion technologies can be applied to biomass-derived syngas. However, before catalysis the raw syngas must be cleaned in order to remove inhibitory substances that would inactivate the catalyst. Volatile tars as well as sulphur, nitrogen and chlorine...
compounds should be removed. The ratio of hydrogen to carbon monoxide in the syngas may need to be adjusted and the CO$_2$ by-product may need to be removed.

**Synthetic natural gas production**
Methanisation is a highly exothermal reaction in which carbon monoxide, carbon dioxide and hydrogen are converted into methane and water. The reaction is catalysed typically by nickel oxide catalysts under pressure of 20-30 bar. The formation of carbon (coking) is a possible undesired side reaction of this process. Cleaning and upgrading of the obtained gas is required for increasing the methane content and reaching natural gas quality, which consists of 98% methane. For this upgrading primarily the drying, desulphurisation and the separation of methane and carbon dioxide is necessary.

**Alcohol (Methanol, ethanol, and higher alcohols) production**
Higher alcohol synthesis (HAS) is performed by the use of catalysts. The catalytic HAS transformation of the syngas to alcohols is typically performed in fixed bed reactors. The optimization of heat removal is of particular interest because the process is highly exothermic. The gas cleaning needs to be adjusted to the requirements of the catalysts. This way of ethanol production may be seen as alternative to lignocellulosic ethanol production.

**Status and scale of production**
Of the 28 thermochemical biorefineries presented by the overview of IEA task 39$^{12}$ ten are operational, of which five pilot plants, four demonstration plants and one commercial plant (Range fuels, Georgia, United States). This shows that commercial operation of these technologies is challenging. A basic technical challenge is that most biomass based syngas is considerably more heterogeneous than natural gas based syngas leading to resulting variations in quality, while the input gases must be relatively clean in order for the synthesis processes (e.g. FT, SNG) to function in a commercially viable way. Most economic studies have indicated that deployment of large scale commercial plants is required to gain the necessary cost reductions from both economies of scale and learning experience for these processes.

### 3.5.4 Hybrid biorefineries
Hybrid biorefineries make use of a combination of thermochemical and biochemical approaches. Examples are
- Gasification followed by fermentation of the resulting gaseous mixture of carbon monoxide, hydrogen and carbon dioxide.
- Fast pyrolysis followed by hydrolysis and/or fermentation of anhydrosugars found in the resulting pyrolysis oil.

**Gasification and fermentation**
The microbiological production of alcohols is a fermentative process based on the utilisation of hydrogen, carbon monoxide and carbon dioxide. Besides alcohols such as ethanol and butanol other chemicals such as organic acids and methane can be obtained.

$^{12}$ http://biofuels.abc-energy.at/demoplants/projects/mapindex
The main advantage of the microbiological process is the mild process conditions compared to thermochemical ethanol synthesis. Due to the lower sensitivity of the used microorganisms towards sulphur, the gas cleaning costs may be significantly reduced. Moreover, there is a higher reaction specificity for a product than in case of inorganic catalysts which is combined with the high tolerance towards the CO/H\textsubscript{2} ratio (no gas shift reaction is needed). The main disadvantage is the limited gas to liquid mass transfer rate requiring specific reactor designs (Bacovsky, Dallos et al. 2010).

**Biorefineries based on fast pyrolysis**

Fast pyrolysis is a process in which organic materials are rapidly heated to 450 - 600 °C in absence of air. Under these conditions, organic vapours, permanent gases and charcoal are produced. The vapours are condensed to pyrolysis oil. Typically, 50 - 75 wt.% of the feedstock is converted into pyrolysis oil. The pyrolysis oil is a mixture of many compounds, of which the main components are:

- acids, aldehydes, sugars and furans from the carbohydrate fraction (cellulose and hemicellulose)
- phenolic compounds, aromatic acids and aldehydes derived from the lignin fraction.

There are several possibilities for acquiring upgraded products from pyrolysis oil, as shown in Figure 21. A strong point of pyrolysis oil is that it can be produced relatively near the biomass source and transported to a larger biorefinery for further upgrading.

One promising upgrading option is hydrogenation, in which the pyrolysis oil is prepared for the use in conventional refineries. The upgrading process primarily involves oxygen removal by hydro-treatment followed by hydrocracking after which co-processing in conventional refineries is possible. The market for conventional fuels and chemicals is large; even a few percent of co-processing in a conventional refinery would involve large volumes of upgraded pyrolysis oil.
3.5.5 Two platform concept biorefinery

The two platform concept biorefinery is based on fractionation of biomass into mainly a sugar (cellulose and hemicellulose) and a lignin fraction (Figure 12):

- The sugar fraction (cellulose and hemicellulose) will be biochemically converted using a so-called “sugar platform” into a portfolio of potential bio-products, such as: materials, chemicals, and fuels.

- The lignin fraction (and the residues from the biochemical process) will be thermo-chemically converted using a so-called “syngas platform” into a syngas for the potential production of a spectrum of bio-based products, including power and/or heat, to meet the internal process power and heat requirements.

![Figure 22 Two platform concept biorefinery (Kamm, Gruber et al. 2006)](image)

3.6 Biorefineries for grassy lignocellulosic biomass - green biorefineries

Like other lignocellulosic biomass green plants, such as grass, alfalfa, immature grain etc. contain cellulose, hemicellulose and lignin fractions. However, in comparison with other vegetable biomass green plants are characterised by a high content of aqueous cell juice with carbohydrates of low molecular weight, a large amount of enzymes (proteins) for photosynthesis and a relatively low content of lignin in the cell walls.

Green biorefineries are based on traditional technologies of green forage preservation, leaf-protein extraction, chlorophyll (pigment) production, and modern biotechnological and chemical conversion methods. All technological concepts of green biorefineries include the separation of the cell juice from the plant framework. Both fractions are subjected to different biotechnological and physiochemical conversion methods. See also Figure 17.

The green biorefinery concept differs from the others because fresh biomass is processed. This means that specific points of interest have to be taken into account, e.g. rapid
primary processing or use of preservation methods (i.e. silage or decomposition) is necessary to prevent degradation of the harvested materials. Three main primary fractionation methods can be distinguished:

- direct wet fractionation
- silage production followed by wet fractionation
- (enzymatic, hydrolytic, or thermal) decomposition followed by fractionation.

Subsequently, various treatments like heat, treatment with organic and inorganic acids, acid anaerobic fermentation, centrifugation, and gel filtration into a leaf nutrient concentrate and brown juice can be used to obtain the different components, such as proteins, lipids, glycoproteins, lectins, sugars, free amino acids, dyes (carotenones), enzymes, lactic acids, minerals and other materials can be obtained. The main markets are in protein recovery for animal feed, preparation of amino acid mixtures for nutrition, and use of lactic acid for bioplastics, chemicals etc.

The press-cake (fibre) fraction can be pressed into pellets for fodder or biofuels or used for biogas production. Furthermore, upgrading to fibre products is possible for applications such as insulation materials, fibre boards, pulp and paper. Finally, applications known from woody lignocellulosic biorefineries could be applied such as hydrolysis and fermentation for ethanol production, lignin utilisation, etc.

**Status and scale of production**

Green biorefineries are still mainly in the pilot plant phase. The pilot plants in Utzenaich (Austria) and Brandenburg (Germany) focus on lactic acid recovery. In Switzerland a demonstration plant provides a grass fibre insulation product on a commercial basis. In the Netherlands several initiatives are under development, amongst others directed to
protein recovery and fibre utilisation. Given the logistics of this (low) density feedstock, it is expected that future commercial green biorefineries will have a medium scale of 10,000 - 100,000 tonnes per year.

3.7 Conclusion

Figure 24 gives a schematic overview of the main steps from raw biomass to products, showing the different platforms.

Table 6 gives an overview of the biorefinery types that were investigated in this chapter together with the feedstock, products, predominant technology, phase of development and the (expected) commercial scale. Small scale means an annual output below 10 ktonnes; medium scale between 10-100 ktonnes; and large scale means > 100 ktonnes. The two platform concept and whole crop biorefinery were not presented since they are based on the other concepts.
<table>
<thead>
<tr>
<th>Type of biorefinery</th>
<th>Feedstock</th>
<th>Products</th>
<th>Predominant technology</th>
<th>Main phase of development</th>
<th>(Expected ) scale&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oleochemical industry</td>
<td>Vegetable and animal oils and fats</td>
<td>Cosmetics, shampoos, soaps, detergents, paint, inks, lubricants, etc.</td>
<td>Fat splitting, hydrogenation, concentration, distillation, refinement, etc.</td>
<td>Commercial</td>
<td>Medium to large</td>
</tr>
<tr>
<td>Sugar based</td>
<td>Sugar cane, beets</td>
<td>New products by sugar or ethanol chemistry, like bioplastics (PHB)</td>
<td>Microbiological polymerisation</td>
<td>Pilot, demo</td>
<td>Small to medium</td>
</tr>
<tr>
<td>Starch based</td>
<td>Potato</td>
<td>Paper (sizing), building additives, adhesives, etc.</td>
<td>Chemical modification: etherification, esterification, splitting</td>
<td>Commercial</td>
<td>Large</td>
</tr>
<tr>
<td></td>
<td>Corn</td>
<td>Corn or dextrose syrup, sweeteners (HFCS)</td>
<td>Wet milling, hydrolysis, isomerisation.</td>
<td>Commercial</td>
<td>Medium to large</td>
</tr>
<tr>
<td></td>
<td>Starch biomass</td>
<td>Bioplastics, PHB, modified products (PHA)</td>
<td>Plasticization, chemical modification</td>
<td>Pilot, demo</td>
<td>Medium</td>
</tr>
<tr>
<td>Biochemical – ethanol production</td>
<td>lignocellulosic biomass: e.g. straw, miscanthus, wood</td>
<td>Ethanol</td>
<td>Pre-treatment, chemical or enzymatic hydrolysis, fermentation, separation</td>
<td>Demo</td>
<td>Medium to large</td>
</tr>
<tr>
<td>Biochemical – lignin utilisation</td>
<td>Spent sulphite pulping liquids (red or brown liquor)</td>
<td>Lignosulfonates (feed pellet binders, dust surpression, dispersants, emulsifiers, surfactants)</td>
<td>Lignin precipitation. In Howard process by addition of excess calcium hydroxide</td>
<td>Commercial (declining, because of decline in sulphite pulping mills)</td>
<td>Medium</td>
</tr>
<tr>
<td></td>
<td>Lignin fraction of lignocellulosic biomass</td>
<td>Replacement phenol formaldehyde resin applications</td>
<td>Organosolv delignification, lignin precipitation</td>
<td>Pilot, demo</td>
<td>Medium</td>
</tr>
<tr>
<td>Thermochemical</td>
<td>lignocellulosic biomass</td>
<td>Diesel, petrol, synthetic natural gas, alcohols</td>
<td>Gasification, gas cleaning and upgrading, catalytic synthesis (SNG, FT, HAS)</td>
<td>Pilot, demo</td>
<td>Large</td>
</tr>
<tr>
<td>Hybrid</td>
<td>lignocellulosic biomass</td>
<td>Ethanol, alcohols, acids, methane</td>
<td>Gasification/pyrolysis, fermentation</td>
<td>Pilot</td>
<td>Medium</td>
</tr>
<tr>
<td>Green biorefineries</td>
<td>wet biomass: green grasses and green crops</td>
<td>Lactic acid, bioplastics</td>
<td>Fractionation in juicy and fibre fraction, various microbiological and biochemical treatments to obtain the components. Fibre upgrading.</td>
<td>Pilot</td>
<td>Small to medium</td>
</tr>
</tbody>
</table>

<sup>a</sup> Expected annual output at commercial scale: small: <10 ktonnes/year; medium: 10-100 ktonnes/year; large: >100 ktonnes/year
4 SELECTION OF CHAINS FOR FURTHER ANALYSIS

The following selection criteria for selecting conversion chains for further analysis of socio-economic impacts are used:

- The input biomass is widely available;
- The input biomass of biorefineries is not competing with food/feed crops;
- There is a realistic prospect that the biorefinery technology will become commercially available between, say, now and 2025.
- The market for the end products is promising; and the total market volume large.

Based on these selection criteria, a preliminary assessment of the presented current and future industrial and small-scale conversion chains has been made. Table 8 provides a summary matrix. BTG has not the intention to judge on the presented technologies. However, the approach makes a justified choice of the main conversion technologies for further analysis possible.

Table 8 Summary of conventional conversion technologies and future biorefineries and their main phase of development

<table>
<thead>
<tr>
<th>Concept</th>
<th>Availability feedstock</th>
<th>No competition with food/feed</th>
<th>State of development</th>
<th>Market perspective</th>
<th>Selected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biodiesel production</td>
<td>+</td>
<td>--</td>
<td>++</td>
<td>++</td>
<td>√</td>
</tr>
<tr>
<td>Pure plant oil production</td>
<td>+</td>
<td>--</td>
<td>++</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Anaerobic digestion</td>
<td>+/-</td>
<td>+/-</td>
<td>++</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Ethanol from sugar/starch crops</td>
<td>+</td>
<td>--</td>
<td>++</td>
<td>+</td>
<td>√</td>
</tr>
<tr>
<td>Combustion lignocellulosic biomass</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>Gasification lignocellulosic biomass</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Pyrolysis woody lignocellulosic biomass</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Carbonisation/torrefaction lignocellulosic</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>biomass</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oleochemical industry</td>
<td>+</td>
<td>-</td>
<td>++</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>Sugar based biorefinery</td>
<td>+</td>
<td>--</td>
<td>+/-</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Starch based biorefinery</td>
<td>+</td>
<td>--</td>
<td>++</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>Biochemical lignocellulosic biorefinery</td>
<td>++</td>
<td>++</td>
<td>+/-</td>
<td>+</td>
<td>√</td>
</tr>
<tr>
<td>(bioethanol &amp; lignin utilisation)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thermochemical lignocellulosic biorefinery</td>
<td>++</td>
<td>++</td>
<td>+/-</td>
<td>+</td>
<td>√</td>
</tr>
<tr>
<td>Hybrid lignocellulosic biorefinery</td>
<td>++</td>
<td>++</td>
<td>-</td>
<td>+</td>
<td>√</td>
</tr>
<tr>
<td>Green biorefineries</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td></td>
</tr>
</tbody>
</table>

Conventional biodiesel from oil crops and ethanol production from sugar and starch crops have been selected because of their relevance and impact in current production systems including competition (for land) with food and feed production. Combustion has been selected as it is the most important current biomass conversion technology for heat and power production. Much is known about the socio-economic impacts as well. For the future conversion technologies, the lignocellulosic biorefinery concepts have been selected. They are seen as solutions to the current bioliquid production chains, having
lower claims on agricultural land and having higher energy yields at lower carbon and energy inputs, compared to oil, sugar and starch based biorefineries.
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Salmiah, A. (2000). Non-food Uses of Palm Oil and Palm Kernel Oil... MPOPC Palm Oil Information Series, Kuala Lumpur.


B. REPORT TASK 3.2 IDENTIFICATION OF SOCIO-ECONOMIC IMPACTS OF CONVERSION CHAINS
Task 3.2 Identification of socio-economic impacts of conversion chains

Introduction to socio-economic impact analysis

WP 3 – Task 3.2

May 2012
The Global-Bio-Pact project (Global Assessment of Biomass and Bioproduct Impacts on Socio-economics and Sustainability) is supported by the European Commission in the 7th Framework Programme for Research and Technological Development (2007-2013). The sole responsibility for the content of this report lies with the authors. It does not represent the opinion of the Community. The European Commission is not responsible for any use that may be made of the information contained therein. The Global-Bio-Pact project duration is February 2010 to January 2013 (Contract Number: 245085).

Global-Bio-Pact website: www.globalbiopact.eu
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1 INTRODUCTION TO IMPACT ASSESSMENTS

1.1 Introduction

Each activity that takes place in a biomass conversion chain as well as each input and output has impacts. Raw materials, labour and capital are the classic ingredients needed for a factory to operate. Technology could be added as a fourth factor that is materialised in capital goods (equipment/hardware, information technology, etc.) and embedded in humans (technical and organisational skills, etc.). The activities in the conversion chain result in various outputs such as final products, jobs, salaries, profits, but also emissions, waste, transport movements, etc. See Figure 1.

The biomass conversion chain (its inputs, outputs and activities) will have various impacts such as socio-economic, fiscal, environmental, and traffic impacts. The impacts can take place at various levels:

- Production unit level
- Community level
- Regional level
- National level
- International level.

Moreover, several types of impacts can be distinguished:

Direct impacts

Direct impacts are the direct consequences of a proposed project’s location, construction or operation on the socio-economic environment. The direct socio-economic impacts of a large-scale development are often manifested as changes in socio-economic structures (e.g. increased employment opportunities, increased income levels, new or expanded social services, etc.).
**Indirect impacts**
Indirect impacts are the secondary consequences of direct impacts (e.g. altered consumption patterns, increased business opportunities and/or an increased need for particular services). The types of indirect impacts that the proposed development may cause, depend largely on an individual and/or community’s priorities, and their ability to manage change.

**Cumulative impacts**
Cumulative impacts are repeated impacts on a valued component. The accumulation of insignificant impacts happening over time can cause one significant impact. An example of a cumulative impact is the effect on housing availability and the cost of living in a community that is experiencing an extended period of in-migration of people employed by several consecutive developments in one region.

### 1.2 Types of impact assessments
Various methods have been developed to assess and quantify the impacts of planned interventions (policies, programs, plans, projects), such as:
- Socio-economic impact assessment (SEIA)
- Environmental impact assessment (EIA)
- Strategic environmental assessment (SEA)
- Social impact assessment (SIA)
- Development impact assessment / sustainable development
- Fiscal impact analysis
- Traffic impact analysis

Within the GlobalBiopact project, the socio-economic impacts of biofuel/bioproduct conversion chains are the main topic of interest. Therefore, the *socio-economic impact assessment (SEIA)* is expected to be the most relevant assessment method. Nevertheless, it is worthwhile to be acquainted with the other impact assessment types. In the next section the SEIA is described and related to the other impact assessment methods.

### 1.2.1 Socio-economic impact assessment
The socio-economic impacts assessment (SEIA) can be defined as follows:
- *Socio-economic impact assessment (SEIA)* is the systematic analysis (used during EIA) to identify and evaluate the potential socio-economic and cultural impacts of a proposed development on the lives and circumstances of people, their families and their communities (Mackenzie 2007).
- *Socio-economic impact assessment (SEIA)* examines how a proposed development will change the lives of current and future residents of a community (Edwards 2011)

The goals of SEIA may vary from simply reducing the negative effects of these actions on people to maximizing their positive benefits and to contribute to sustainable development.
The concepts used in SEIA are derived from a number of social disciplines, including economics, sociology, geography, anthropology and political science. The key issue and challenge in SEIA is understanding the nature of social or economic impacts. An impact is a change in conditions caused by a development, such as a road or a mine. Generally, socio-economic impacts are changes in the human condition. They are changes in the economic and social conditions of local communities, vulnerable groups (such as women, children or poor), businesses and employees, districts, provinces or even the nation. Generally, health and cultural impacts (e.g., language loss) are also the subject of SEIA, but are not always covered in depth, as they may need special study. Social and economic impacts may each require specific study and analysis using various techniques.

Various other assessment methods have been developed in order to determine the impacts of projects, policies, programs and plans. Below a selection of these assessment methods are defined and related to SEIA.

1.2.2 Environmental impact assessment and strategic environmental assessment

Environmental impact assessment and strategic environmental assessment are generally defined as follows:

- **Environmental Impact Assessment (EIA)** is a systematic process to identify, predict and evaluate the environmental effects of proposed actions and projects. This process is applied prior to major decisions and commitments being made. A broad definition of environment is adopted. Whenever necessary, social, cultural and health effects are considered as an integral part of EIA (UNEP 2002).

- **Strategic Environmental Assessment (SEA)** refers to a formal process of systematic analysis of the environmental effects of development policies, plans, programmes and other proposed strategic actions. This process extends the aims and principles of EIA upstream in the decision-making process, beyond the project level and when major alternatives are still open (UNEP 2002).

Socio-economic impact assessments (SEIA) are often seen as additional to environmental impact assessments (EIA). (Mackenzie 2007) states: “In the past EIA focused on direct and indirect biophysical impacts of proposed developments (i.e. impacts of development activities on water, air, land, flora and fauna). In recent years the impacts of industrial development on society, culture and different forms economic activity have gained equal importance in EIA.” Especially when the social impacts are high, for instance when a big dam is planned, it is obvious that carrying out a SEIA in addition to an EIA is essential. EIA procedures and frameworks have been used as a base to develop SEIA.

1.2.3 Social impact assessment

Social impact assessment can be defined as follows:

- **Social impact assessment (SIA)** includes the process of analysing, monitoring and managing the intended and unintended social consequences, both positive and negative of planned interventions (policies, programs, plans, projects) and any social change processes invoked by those interventions. Its primary purpose is to
bring about a more sustainable and equitable biophysical and human environment (IAIA 2003).

- Social impact Assessment (SIA) identifies the consequences to people of any proposed action that changes the way they live, work, relate to one another, organise themselves and function as individuals and members of society, with particular attention to the mitigation of adverse or unintended aspects. This definition includes social-psychological changes, for example to people’s values, attitudes and perceptions of themselves and their community and environment (based on (UNEP 2002), topic 13).

The main types of social impact that occur as a result of these project-related changes can be grouped into five overlapping categories (UNEP 2002) (topic 13):

- lifestyle impacts – on the way people behave and relate to family, friends and cohorts on a day-to-day basis;
- cultural impacts – on shared customs, obligations, values, language, religious belief and other elements which make a social or ethnic group distinct;
- community impacts – on infrastructure, services, voluntary organisations, activity networks and cohesion;
- amenity/quality of life impacts – on sense of place, aesthetics and heritage, perception of belonging, security and liveability, and aspirations for the future;
- Health impacts – on mental, physical and social well-being, although these aspects are also the subject of health impact assessment.

The definitions of Social Impact Assessment (SIA) are very comparable to those of Socio-economic impact analysis (SEIA). These assessment types are sometimes mixed. However, it is clear that in a proper SEIA both social and economic impacts are studied.

1.2.4 Development impact assessment / sustainable development

The classic definition of sustainable development is “development that meets the needs of the present without compromising the ability of future generations to meet their own needs”

1. The United Nations 2005 World Summit Outcome Document refers to the "interdependent and mutually reinforcing pillars" of sustainable development as economic development, social development, and environmental protection. See Figure 2. By combining environmental impact assessment and socio-economic impact assessments the sustainable development impact can be assessed.

---

(Sustainable) Development impact assessment is defined as follows:

- **Development impact assessment** involves a process to comprehensively evaluate the consequences of development on a community. The assessment process should be an integral part of the planning process as it provides extensive documentation of the anticipated economic, fiscal, environmental, social and transportation-related impacts of a particular development on a community (Edwards 2011).

- **Sustainable development assessment (SDA)** is an overarching methodology (with many components), which is used in evaluating investment projects (as well as programs and policies), to ensure balanced analysis of both development and sustainability concerns. The ‘economic’ component of SDA is based on conventional economic and financial analysis (including cost benefit analysis). The other two key components are environmental and social assessment (EA and SA). However, many other more specialized types of assessments may be included within an integrated SDA.

### 1.2.5 Other impact assessments analyses

Some other impact assessment analyses were found:

- **Fiscal impact analysis** estimates the impact of a development or a land use change on the costs and revenues of governmental units serving the development. (The analysis enables local governments to estimate the difference between the costs of providing services to a new development and the revenues—taxes and user fees, for example—that will be generated by the development.) (Edwards 2011)

- **Traffic impact analysis** is a study which assesses the effects that a particular development’s traffic will have on the transportation network in the community. Traffic impact studies should accompany developments which have the potential to impact the transportation network (Edwards 2011).

Fiscal impact analysis could be part of an economic impact assessment. A traffic impact analysis could typically be included in an environmental impact assessment.
2 SOCIO-ECONOMIC IMPACT ANALYSIS

2.1 Introduction
In the last decade, broad guidelines for the practice of socio-economic impact assessment (SEIA) have been developed at the practitioner level. For example, principles for SEIA have been developed by the International Association for Impact Assessment (IAIA) (IAIA 2003); Mackenzie has published socio-economic impact assessment guidelines (MVEIRB 2007); and UNEP has published an Environmental Impact Assessment Training Resource Manual, that includes a chapter on social impact assessment (UNEP 2002). Based on this documentation it is possible to determine an approach for assessing the socio-economic impacts for the case studies in the Global Biopact project.

2.2 Steps of the SEIA process
The following main steps are followed in the SEIA process (Mackenzie 2007):

1. Scoping and issues identification: The proposed project must be well-defined. Social and economic issues must be identified as well as the geographic and temporal study boundaries.

2. Determining the social and economic baseline: There must be a good understanding of the impacted community or communities and the general socio-economic conditions in the project area.

3. Predicting and analysing impacts: The assessment must be able to project what the social and economic impacts may be, including the effect of potential interactions between factors and over the lifetime of the development.

4. Determining significance: There must be an assessment of the importance of the social and economic impacts of the project.

5. Mitigation, management and monitoring: Once impacts and their significance are understood, decisions must be made about whether the project should proceed. If so, measures must be identified to avoid or lessen negative impacts and maximize positive impacts (mitigation). Management of the mitigation needs to occur and on-going monitoring of the projects effects must be carried out to ensure thresholds are not crossed.

These steps are further worked out in the next sections. More information can also be found in (Mackenzie 2007).
2.3 Scoping and issues determination - appropriate level of SEIA

Before starting an SEIA it is important to determine its scope consisting of:

- The scope of development
- The scope of issues
- The scope of assessment

Scope of development
The scope of development includes a description of the project to be studied in the SEIA, including the needed human resources, skills, goods and services and changes to the physical infrastructure.

Scope of issues
Potentially relevant impacts need to be identified. An initial selection can be made with help of existing long lists of possible impacts. Initially or later in the process, also community members need to be involved to ensure that relevant impacts are included.

The Global Biopact Task 2.1 report (van Dam, Faaij et al. 2010) provides a list of socio-economic impacts relevant for biomass production, classified under the following themes:

- Working conditions and rights
- Economic aspects
- Competition and availability of natural resources
- Social aspects and welfare
- Health impacts
- Food security
- Smallholder aspects
- Policy and governance aspects
- Land tenure and rights
- Participatory aspects

Each theme consists of a number of potential impacts. The theme working conditions and rights includes for instance:

- Freedom of association and collective bargaining
- Forced labour
- Elimination of child labour and protection of children and young persons
- Equality of opportunity and treatment
- Minimum wages
- Working time
- Health and safety
- Social security
- Unemployment benefit
- Social security for migrant workers
- Maternity protection
- Migrant workers

Most of these themes and their underlying impacts potential impacts could be relevant for biomass conversion projects too.
If needed, other lists of potential impacts can be used to support the process of impact identification. An initial list of issues - mainly relevant on community level - is provided below (Mackenzie 2007):

- Housing (access, appropriateness, affordability)
- Family/household stability
- In-migration and out-migration
- Maintenance of cultural values such as language
- Access to land for traditional uses
- Traditional economy-harvesting success
- Income and levels of disposable income
- Cost of living and inflation
- Employment levels
- Community expectations
- Business opportunities
- Gender equity
- Inter-generational equity
- Access to education/training and their perceived value
- Human health concerns including access to services
- Pressure on infrastructure (roads, buildings)
- Public safety concerns
- Level and accessibility of social services provided
- Lifestyle choices
- Boom and bust economic cycles
- Archaeological/heritage resources

Another extended list of potential impacts from (UNEP 2002) is provided in Annex A.

During the SEIA process some issues initially included might be found less relevant, and some new issues might be added to the selection.

The scope of assessment
Depending on the type of the listed potential impacts, the spatial boundaries of the SEIA can be defined. It is likely that many social impacts take place on company and community level; some impacts such as the contribution to the GDP can be assessed on a national level. Furthermore, it should be defined which stages of the project are included in the SEIA. The following stages can be distinguished: planning, construction, operation, decommissioning, and post closure stage.

Level of detail of the SEIA
SEIA can be carried out at different levels of detail. It is reasonable to link this level to the size of the project and the expected level of concern related impacts. (MVEIRB 2007) distinguishes basic, moderate and comprehensive SEIAs and developed a test to determine which level is appropriate.

Basic SEIA
In a basic SEIA the following information should be included:
1. A record and description of efforts to consult potentially affected communities and other parties.
2. A development description, including the following socio-economic data:
   - Total estimated capital costs of the proposed development, including annual operating costs
   - Approximate number of workers including the developer’s employees and contractors, and number of person days/years of work for the proposed development, including subcontracting
   - Identified archaeological resources within the footprint of the proposed development
   - A list of any extra regional infrastructure required for the proposed development to proceed
3. Any identified potential impacts on the socio-economic environment, and suggestions for mitigating these impacts

**Moderate SEIA**

In moderate SEIA a distinction is made between the construction, operating, maintenance and decommissioning phase of the proposed project. Additional information requirements are described in (MVEIRB 2007) (chapter 3, table 7).

**Comprehensive SEIA**

Complex large-scale and long-term developments such as large mines, oil and gas operations, pipelines, large new highways, hydroelectric dams, etc. demand for a comprehensive SEIA. The SEIA needs to be carried out well in advance of the proposed development. See (MVEIRB 2007) (chapter 3, page 28-29)

**2.4 Determination of baseline**

The developer should describe the current socio-economic and cultural environment and the context of the proposed project. It can be difficult to determine whether an impact is caused by the proposed project. The socio-economic environment will continue to evolve whether the project occurs or not. The occurrence of two simultaneous projects/developments can make it hard to attribute the impacts between the projects. Even the issue whether an impact is adverse or beneficial, depends on an individuals’ personal choice. For example, increased disposable income can create stronger families, brighter futures for children and greater health; or it can fuel anti-social behaviour (MVEIRB 2007).
The choice of methods and tools for characterising and predicting social and cultural impacts is essential.

2.5 Methods for predicting and determining economic impacts

Economic issues are given substantial emphasis in SEIA. Possible economic impact assessment tools include fiscal analysis, cost benefit analysis and input/output analysis.

2.5.1 Fiscal analysis

Fiscal impact analysis estimates the impact of a project or development on the costs and revenues of governmental units serving the project or development. It focuses on the inter-relationship between project viability and government costs and revenues. Government obtains revenues from a project through a variety of taxes, fees, and royalties. Government may also impose conditions on the developer that will raise the costs of government institutions managing and monitoring the environmental and socio-economic standards of a project. If the net cost of all of these elements is too high, the project will not proceed. A balance is required. Fiscal analysis also concerns intergovernmental relationships with respect to project revenues and costs (Mackenzie 2007).

2.5.2 Cost-benefit analysis

Cost-benefit analysis is a technique used to compare the various costs associated with an (investment) project with the benefits that it proposes to return. Most feasibility studies use cost-benefit analysis to determine the feasibility of a project. Typical indicators used are Net Present Value (NPV), Internal Rate of Return (IRR), Simple Payback Period and figures showing yearly cash flows. In order to make this calculation, traditionally, the main inputs and outputs of the project need to be identified, including direct labour costs, use of intermediary products, quantities of waste etc. as far as they have a direct financial impact on the proposed project. In addition, it is possible to quantify the costs and benefits of environmental impacts, cost effectiveness of mitigation and, where possible, environmental and social costs of intangibles (for example, value of country food or costs of pollution) in monetary units (e.g. dollars, euros). In some cases, the environmental and social cost/benefit estimates provided in the SEIA are then used to perform an overall
economic analysis of the project. An overall economic analysis evaluates the total economic value of a project (e.g. will the project provide society, overall, with overall positive benefits or not?).

An example data format for cost-benefit analysis for biomass production is provided in Annex B.

### 2.5.3 Input/output analysis

Input-output (I/O) analysis studies the interrelationships within and between economic sectors of a country and can be used to determine the impacts of an economic activity on the whole economy. The I/O method is based on a country’s I/O table, which is available from national statistical bureaus and which generally concerns the country’s economy for a time period of one year. There are two options by which a new industry can be introduced to the economy. The first method is based on creating a new final demand vector, while the second method is based on including the new industry in the technology matrix. Despite the first method’s popularity, the second method has the advantage that it accounts for the impacts of the introduction of a new sector in a more complete manner. That is to say, the second way not only accounts for the inputs being bought by the new sector from the existing sectors, but can also account for its outputs being consumed by the existing sectors (Wicke 2006). The construction of an input-output table requires a large amount of data on inter-industry flows and other variables. Governments are often the only organizations with adequate resources for designing these models, and collecting and analysing the required data. Other agencies wishing to use input/output models must often rely on existing models developed by government (Mackenzie 2007). For more information and examples of input/output analysis applied to a biomass conversion chain see (Wicke 2006), (van den Broek, van den Burg et al. 2000), (Wicke, Faaij et al. 2006).

### 2.6 Methods for predicting and determining social impacts

Many consultative techniques are used in SEIA to identify issues, predict impacts and plan for mitigation. These include surveys, public meetings, workshops, focus groups, networks, and checklists. See Table 1 for an overview of commonly used tools.
**Table 1 Tools for determination of social impacts (Mackenzie 2007)**

<table>
<thead>
<tr>
<th>Social analysis techniques</th>
<th>Description</th>
<th>Issues</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Surveys / Questionnaires</strong></td>
<td>• Continuous or one-time</td>
<td>• If a carefully designed survey keeps turning up a particular answer, causality is suggested</td>
</tr>
<tr>
<td></td>
<td>• Targeted at impacted individuals (e.g. those employed during project, workers spouses, etc.)</td>
<td>• Poor design can yield inadequate</td>
</tr>
<tr>
<td></td>
<td>• If a carefully designed survey keeps turning up a particular answer, causality is suggested</td>
<td>• Responses</td>
</tr>
<tr>
<td><strong>Focus Groups / Workshops</strong></td>
<td>• Held in groups of 6 or less (the smaller the group, the more productive the session) of individuals well informed on a particular topic</td>
<td>• A well-conducted focus group/workshop can yield a great deal of very useful information and insight.</td>
</tr>
<tr>
<td></td>
<td>• Collaborate to move towards consensus on key issues.</td>
<td>• Moderate disagreement would normally suggest that there should be no attribution until more evidence of causality has been obtained</td>
</tr>
<tr>
<td><strong>Community Meetings</strong></td>
<td>• Held in public to identify community based concerns</td>
<td>• Effective when identifying broad issues regarding impacts (e.g. do you think what is happening is good or bad?)</td>
</tr>
<tr>
<td></td>
<td>• Provides opportunity for open dialogue</td>
<td>• Good indicator of public support / unhappiness</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• A poorly organized public hearing can be counterproductive, leading to polarization of views; to unfounded fears about the socio-economic impacts of the project; or to unfounded confidence in the project.</td>
</tr>
<tr>
<td><strong>Networks / Technical Advisory Committees</strong></td>
<td>• Experts on particular issues relevant to the assessment process who lend advice on an ongoing basis (community leaders/ policy analysts)</td>
<td>• Difficult to establish. Development can take time and energy</td>
</tr>
<tr>
<td><strong>Checklists</strong></td>
<td>• Matrices are useful in ensuring relevant impacts are identified. Design requires giving consideration to key component impacts of a project.</td>
<td>• Useful in making inter-community comparisons – identifying how various communities may see things differently.</td>
</tr>
<tr>
<td><strong>Ethnographic / Ethnohistoric Studies</strong></td>
<td>• Focused study of the impacts of development on indigenous communities on social organization</td>
<td>• Difficult to carry out in the timeframe of an SEIA. Alternative is the Rapid Ethnographic Assessment Procedures (REAP) of cultural mapping, in-depth interviewing, focus groups supplemented with limited survey research.</td>
</tr>
</tbody>
</table>

These techniques are effective for identifying present vulnerability and identifying development futures, and involve stakeholders in the identification of issues and concerns. Once issues and concerns are identified, the social analyst normally consults case studies of similar projects to compare impacts. If time permits, focused ethnographic research may be carried out. Otherwise rapid cultural appraisal techniques can be used.
In the Global BioPact project ‘rapid appraisal’ techniques can be applied. An example of the design of a rapid appraisal method is given below. The case study partners will determine their approach based on the particular situation of the case.

### Rapid appraisal

Rapid appraisal firstly involves collecting data from existing written sources. Secondly, ‘key informants’ are recruited to help obtain the views of local people. Key informants are local people who have a good knowledge of the local area. Their opinions are sought and they are asked to identify further informants, and if willing can join the research team to assist in interviewing other local people. The final stage of the process is a validation workshop, which provides an opportunity to feedback on findings and identifies any remaining gaps.

2.7 Determining significance and mitigation

After analysis of impacts it is important to evaluate whether the (negative) impacts are acceptable. If negative impacts are below an acceptable threshold, proper mitigation measures must be taken or ultimately the project should be terminated. The acceptable threshold can be determined using traditional and local knowledge, community based knowledge, standards, guidelines, policy statements, and biomass sustainability certification systems. In many cases mitigation measures can be identified and discussed with impacted communities, governments and other stakeholders. From the positive view, measures can be taken to benefit optimally from the positive impacts of the project. Management of the mitigation needs to occur and on-going monitoring of the projects effects must be carried out to ensure thresholds are not crossed.
A. LIST OF SOCIAL IMPACTS

(UNEP 2002) provides a long list of possible social impacts on individual and household level and on community and institutional level.

**Individual and household level**

1. death, death of family member
2. arrest, imprisonment, detention, torture, intimidation or other abuse of human rights inflicted on individual
3. reduced availability of food and adequate nutrition
4. reduced control over fertility (availability of contraception, and empowerment)
5. reduced level of health and fertility (ability to conceive)
6. reduced mental health increased stress, anxiety, alienation, apathy, depression
7. uncertainty about impacts, development possibilities, and social change
8. actual personal safety, hazard exposure
9. experience of stigmatisation and deviance labelling
10. reduction in perceived quality of life
11. reduction in standard of living, level of affluence
12. worsening of economic situation, level of income property values
13. decreased autonomy, independence, security of livelihood
14. change in status or type of employment, or becoming unemployed
15. decrease in occupational opportunities potential diversity flexibility in employment
16. moral outrage, blasphemy, religious affront, violation of sacred sites
17. upset (objection/opposition to the project), NIMBY (not in my back yard)
18. dissatisfaction due to failure of a project to achieve heightened expectations
19. annoyance (dust, noise, strangers, more people)
20. disruption to daily living, way of life (having to do things differently)
21. reduction in environmental amenity value
22. perception of community, community cohesion, integration
23. community identification and connection to place (do I belong here?)
24. changed attitude towards local community, level of satisfaction with the neighbourhood
25. disruption to social networks
26. alteration in family structure and stability (divorce)
27. family violence
28. gender relations within the household
29. changed cultural values
30. changed perceptions about personal health and safety, risk, fear of crime
31. changed leisure opportunities
32. quality of housing
33. homeliness
34. density and crowding
35. aesthetic quality, outlook, visual impacts
36. workload, amount of work needed to be undertaken to survive/live reasonably
Community and institutional level
1. death of people in the community
2. violation of human rights, freedom of speech
3. adequacy of physical infrastructure (water supply, sewerage, services and utilities)
4. adequacy of community social infrastructure, health, welfare, education, libraries, etc.
5. adequacy of housing in the community
6. workload of institutions, local government, regulatory bodies
7. cultural integrity (continuation of local culture, tradition, rites)
8. rights over, and access to, resources
9. influences on heritage and other sites of archaeological, cultural or historical significance
10. loss of local language or dialect
11. debasement of culture
12. equity (economic, social, cultural)
13. changed equity/social justice issues in relation to minority or indigenous groups
14. gender relations in the community
15. economic prosperity
16. dependency/autonomy/diversity/viability of the community
17. unemployment level in the community
18. opportunity costs (loss of other options)
19. actual crime
20. actual violence
21. social tensions, conflict or serious divisions within the community
22. corruption, credibility and integrity of government
23. level of community participation in decision making
24. social values about heritage and biodiversity.
B. EXAMPLE DATA FORMAT FOR CBA (COST BENEFIT ANALYSIS)

Table 2 and Table 3 show the factors that should be taken into account when making a CBA.

**Table 2: crop specific data**

<table>
<thead>
<tr>
<th>Expenses per hectare ($/ha)</th>
<th>Labour requirements (hours/ha)¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yield (m³ or kg/ha)</td>
<td>Land clearing</td>
</tr>
<tr>
<td>Field clearing</td>
<td>Field preparation</td>
</tr>
<tr>
<td>Field preparation material</td>
<td>Planting</td>
</tr>
<tr>
<td>Irrigation</td>
<td>Irrigation</td>
</tr>
<tr>
<td>transplanted</td>
<td>Weed control</td>
</tr>
<tr>
<td>for cultivation</td>
<td>Pruning</td>
</tr>
<tr>
<td>farmer</td>
<td>Fertilising</td>
</tr>
<tr>
<td>Pesticides (quantity as well)</td>
<td>Pest and disease</td>
</tr>
<tr>
<td>for PHA</td>
<td>Harvesting</td>
</tr>
<tr>
<td>for storage</td>
<td>Packing/transport</td>
</tr>
<tr>
<td>packing</td>
<td>PHA (post harvest preparation)</td>
</tr>
<tr>
<td>transport</td>
<td>Packing/transport*</td>
</tr>
<tr>
<td>Year 0</td>
<td></td>
</tr>
<tr>
<td>Year 1</td>
<td></td>
</tr>
<tr>
<td>Year 2</td>
<td></td>
</tr>
<tr>
<td>Year 3</td>
<td></td>
</tr>
<tr>
<td>Year 4</td>
<td></td>
</tr>
<tr>
<td>Year 5</td>
<td></td>
</tr>
<tr>
<td>Etc. until Year 24</td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
</tr>
</tbody>
</table>

¹ Indicate the type of labour: unskilled, semi-skilled or high skilled

² Relates to yield
It is also important to indicate how often tasks like weeding are performed (per year or per lifetime of the crop). Other crop-specific factors that influence the profitability:
- Market price, (farm gate price, wholesale dealer price, consumer price)
- Transport distance and costs to processing unit, conversion costs

Table 3: Country specific data

<table>
<thead>
<tr>
<th></th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Labour expenses</strong></td>
<td></td>
</tr>
<tr>
<td>Average wage rate</td>
<td>$/day</td>
</tr>
<tr>
<td>Unskilled labour</td>
<td>$/hour</td>
</tr>
<tr>
<td>Semi-skilled labour</td>
<td>$/hour</td>
</tr>
<tr>
<td>Skilled labour</td>
<td>$/hour</td>
</tr>
<tr>
<td>Agricultural sector</td>
<td>$/hour</td>
</tr>
<tr>
<td>Manufacturing sector</td>
<td>$/hour</td>
</tr>
<tr>
<td><strong>Land costs</strong></td>
<td></td>
</tr>
<tr>
<td>Land rent</td>
<td>$/ha</td>
</tr>
<tr>
<td>Land rent commercial</td>
<td>$/ha</td>
</tr>
<tr>
<td>Opportunity cost of land</td>
<td>$/ha</td>
</tr>
<tr>
<td><strong>Utility costs</strong></td>
<td></td>
</tr>
<tr>
<td>Diesel, Petrol, kerosene</td>
<td>$/l</td>
</tr>
<tr>
<td>Taxes (excise duty, road toll)</td>
<td>% and $</td>
</tr>
<tr>
<td>Electricity</td>
<td>$/GJ</td>
</tr>
<tr>
<td><strong>Financial costs</strong></td>
<td></td>
</tr>
<tr>
<td>Exchange rate</td>
<td>$</td>
</tr>
<tr>
<td>Inflation</td>
<td>%</td>
</tr>
<tr>
<td>Discount rate</td>
<td>%</td>
</tr>
</tbody>
</table>

The costs listed in Table 2 and Table 3 can also contribute to an Input Output Analysis. For an I/O analysis additional data is required such as: total imports, direct-, indirect- and induced labour and so on, see e.g. (Wicke, Smeets et al. 2009).
REFERENCES


