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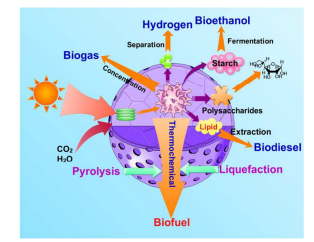


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Thermochemical conversion of low-lipid microalgae for the

production of liquid fuels

Yu Chen^a, Yulong Wu^{a,*}, Derun Hua^a, Chun Li^b, Michael P. Harold^{c,*}, Jianlong Wang^a and Mingde Yang^a



This *critical review* provides an investigation elaborated by recent references to different strategies for the sustainable conversion of microalgae to liquid fuels (bio-oil). Microalgal biomass is considered as the third generation of biofuel, which has notable advantage over other biomass in that it does not compete with food or cropland resources. The conversion of algal biomass into liquid fuels provides a long-term sustainable option for fuels production, which can be achieved in an environmentally compatible manner. This article reviewed recent developments in the field of algal biomass conversion into liquid fuels, especially thermochemical conversion of low-lipid microalgae. We start with a brief introduction of microalgae and its biochemical components. After an overview of the main strategies involved in algal biomass conversion, we focus on the thermochemical conversion of algae, including pyrolysis and hydrothermal liquefaction and compare the methods in detail. In addition, the catalytic upgrading of algae based crude bio-oil was also examined. Finally, challenges and opportunities for future research on the production of bio-oil from microalgae are analyzed.

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Review

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Thermochemical conversion of low-lipid microalgae for the production of liquid fuels: challenges and opportunities

Yu Chen^a, Yulong Wu^{a,d,*}, Derun Hua^a, Chun Li^b, Michael P. Harold^{c,*}, Jianlong Wang^a and Mingde Yang^a

The development of renewable biomass energy sources has attracted attention because of the potential for a sustainable fuel s with a low carbon intensity. Microalgae are considered as the third generation biofuel, and have a notable advantage over other biomass in that it does not compete with food or cropland resources. The conversion of algal biomass into liquid fuels provides a long-term sustainable option for fuels production, which can be achieved in an environmentally compatible manner. Among the microalgal conversion methods, thermochemical conversion, which can make full use of all components in the algae, is viewed as one of the best conversion method, especially for low-lipid microalgae. This article reviews recent developments in

10 the field of algal biomass conversion into liquid fuels, with particular attention focused on the thermochemical conversion of low-lipid microalgae. We start with a brief introduction of microalgae and its biochemical components. After an overview of the main strategies involved in algal biomass conversion, we focus on the thermochemical conversion of algae, including pyrolysis and hydrothermal liquefaction and compare the two methods in detail. In addition, the catalytic upgrading of algaederived crude bio-oil was also examined. An assessment is made of the challenges and opportunities of a commercial-scale

15 microalgae-to-fuels process in light of mitigating technical, environmental, and logistical issues.

1. Introduction

Concerns over geopolitical and environmental implications of petroleum availability, supply, and consumption have been ²⁰ growing in recent years.¹ With the eventual depletion of fossil fuels as a source for fuels and chemicals, the need for the development of sustainable renewable energy has become a global theme aimed at addressing the issues on climate change,

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technology.

Yulong Wu an Associate Professor since 2007. From 2007-2008, he worked as visiting scholar at Saarland University in Germany. He became New Century Excellent Talent of Ministry of Education in 2012, and he became Member of Directors Board of China Energy Society in 2010. His main scientific interests are conversion of biomass energy, heterogeneous catalysis and resource chemical engineering.

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energy security, and ever-increasing demand for limited 35 petroleum resources. This interest has prompted researchers to develop potentially viable approaches for the production of biofuel from biomass, notwithstanding recent advances in the recovery of petroleum and gas by advanced production methods like fracturing and horizontal drilling.^{2,3} The utilization of 40 bioresources for the production of alternative fuels provides one of long-term sustainable options for fuels production which may be accomplished in an environmentally compatible manner for many regions of the globe.4,5 Among the renewable biomass resources, microalgae are viewed as next generation fuel 45 feedstock due to its superior photosynthetic efficiency, higher growth rate, area-specific yield, and higher carbon dioxide utilization capabilities compared to terrestrial plants.



Yulong Wu is an Associate Professor of Institute of Nuclear and New Energy Technology (INET), Tsinghua University. Hereceived BS (1994) from Wuhan University of Technology, obtained MS and PhD degree in Chemical Engineering from Tianjin University in 1998 and 2001, respectively. He became a Postdoc at Department of Chemical Engineering, Tsinghua University, and then joined the faculty of INET, Tsinghua University. He has been

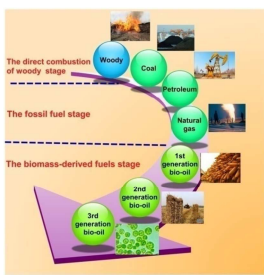


Fig. 1 Development of energy formation in the process of human history

Prior to the exploitation of low-cost fossil fuels, humankind was dependent on the direct combustion of woody biomass to 5 meet its energy demands (Fig. 1). The fossil fuel era commenced with the use of coal for heat and electricity. The discovery of petroleum crude oil provided an inexpensive liquid fuel source that significantly improved the standard of living and, by most accounts, catalyzed the industrial revolution.^{6,7} Biomass-derived ¹⁰ fuel, called "biofuel" or "bio-oil" for short, has attracting considerable attention and research activity because it is derived



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Derun Hua

Jianlong Wang Mingde Yang

from renewable biomass resources and potentially relieves the entrenched global dependence on petroleum-based fuels.⁸⁻¹⁰ Moreover, biofuels are a potentially carbon-neutral feedstock. At 15 the cornerstone of this green industrial revolution is the potential

for algae as one of biofuel feedstocks.⁴ The concept of using microalgae to produce fuels has already been discussed for more than half a century, but a concerted effort started from the first oil crisis in the 1970s.¹⁰

Microalgae are considered as an attractive optional 20 feedstock for several compelling reasons:¹¹⁻¹⁵ (1) Potential biofuel yields from certain microalgae strains are projected to be at least 60 times higher than that from soybeans, approximately 15 times more productive than terrestrial plants, and nearly 5 times of that 25 of palm oil per acre of land on an annual basis; (2) the growth cycle of microalgae is comparatively short with biomass yield that can double within 24 h; (3) the biomass production of microalgae is 5-30 times higher than that of traditional oil crops per unit surface area; (4) microalgae can be rich in oil, over 60% 30 by weight of dry biomass in some species; (5) microalgae does not pose a threat to traditional agricultural resources as they can be cultivated on non-arable land or waste water; (6) microalgal harvesting can be integrated with a fossil-fuel-fired power plants for capture and use of CO₂ via photosynthesis. Finally, the 35 cultivation of microalgae may be coupled with wastewater bioremediation via removing nitrogen, phosphorus, and heavy metals.



Michael P. Harold is the Chairman and the M. D. Anderson Professor of Chemical and Biomolecular Engineering at the University of Houston (UH). Harold, who received his chemical engineering education at Penn State (BS) and UH (PhD), is the author of more than 125 publications and has given over 275 presentations and invited lectures. He held earlier positions at University of Massachusetts (1985-1993) and DuPont Company (1993-2000).

Michael P. Harold

His honors include the Esther Farfel Award from UH and the ACS Richard A. Glenn Award. In 2011 Harold was appointed the 7th Editor-in-Chief of AIChE Journal.

Derun Hua earned his MS degree from University of Jinan in Shandong (2007) and his PhD degree from China University of Petroleum (2011), all in Chemical Engineering. He engaged in postdoctoral research at Tsinghua University in 2011. He has been an Associate Professor of Ganna Normal University in China since 2013. He has interests in conversion of biomass, and synthesis of catalyst. Jianlong Wang is a professor and deputy director of Institute of Nuclear and New Energy Technology, Tsinghua University. He received his PhD from Harbin Institute of Technology in 1993. He worked at Leeds University (England) as a visiting professor from 1998 to 1999 and worked at Berlin Technical University (Germany) as a Humboldt Researcher from 2000 to 2002. Prof. Wang's research focuses on water and wastewater treatment,

environmental biotechnology. Prof. Wang has published more than 300 papers in peer-reviewed academic journals. **Mingde Yang** received BS (1983) and MS (1986) degrees in Chemical Engineering from Heilongjiang University and Tsinghua University, respectively. Yang joined the faculty of the Institute of Nuclear and New Energy Technology at Tsinghua University, where he became Associate Professor in 1999, and was promoted to full professor in 2008. From 1991 to 1993, He was a visiting scientist at Karlsruhe University in Germany, and was a Visiting Research Scholar at Atomic Energy Research Institute in Japan in 1996. His main scientific interests are high efficient utilization of biomass resource, solvent extraction technology, and environmental treatment technology on catalysis, etc.

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The original concept for converting microalgal biomass into biofuels involved lipid extraction for the production of biodiesel via transesterification. However, compared to low-lipid algal strains with the content of lipid less than 15 wt.%, high-lipid 5 microalgal species typically have lower biomass productivity and

- growth rates, and require stringent and controlled cultivation. In contrast to the transesterification process in which only the lipids in the algae are utilized, thermochemical routes involve the conversion of the entire algal cell, including the proteins, ¹⁰ carbohydrates and lipids, into fuel oil.¹⁶ Indeed, the conversion of
- low-lipid microalgae into biofuel in the future is an active area of research.¹⁷⁻²¹

Microalgal proteins, lipids, and carbohydrates contain C, H, O, N and other elemental moieties. As a result, the ¹⁵ thermochemical conversion of microalgae is a challenge for its effective utilization, C and H are the primary elemental constituents of fossil fuels and conventional refining and petrochemical processing follows from that fact. Petroleum is under-functionalized, containing mostly C and H, and therefore ²⁰ requires of the addition of functional groups through oxidation,

amination, hydration, etc. To the contrary, microalgal biomass is

over-functionalized and requires removal of functionality, particularly the O- and N- containing groups.⁹ Indeed, this leads to the greatest challenge on the production of liquid fuel from ²⁵ lignocellulosic and microalgal biomass via thermochemical conversion.²²⁻³⁵

There have been earlier reviews on the conversion of microalgae to biofuels.^{1,16,27,36-38} Several reviews on the conversion of high-lipid microalgae to biodiesels covering topics ³⁰ like lipid extraction and transesterification have also appeared.^{8,13,15} In this review, our intent is to highlight the latest developments of microalgal biomass thermochemical conversion methods and future prospects for converting low-lipid microalgae into liquid fuels. This review primarily focuses on ³⁵ thermochemical conversion technologies (pyrolysis and hydrothermal liquefaction) of microalgae with/without catalysts as well as the catalytic upgrading of crude bio-oils into end products. Since the overwhelming majority of transportation fuels (excess 90%) is on the basic of liquid fuels, so we focus attention ⁴⁰ on liquid fuels obtained from microalgal biomass in this review.

Microalgae classification	Species of selected microalgae	Proteins	Carbohydrates	Lipids	Ref.
Anabaena	Anabaena cylindrica	43-56	25-30	4-7	[43]
Batrachospermum	Aphanizomenonflos-aquae	62	23	3	[43]
Chlamydomonas	Chlamydomonasrheinhardii	48	17	21	[43]
	Chlorella protothecoides	53	11	15	[44]
	Chlorella pyrenoidosa	71.3	22	0.1	[42]
Aegagropila	Chlorella spp.	30	15-17	9-13	[45]
	Chlorella vulgaris	42-58	12-17	14-22	[21,27,46,47]
Cladophora	Cladophora sp.	25	25	6	[48]
Desmodesmus	Desmodesmus sp.	38-44	13-20	10-14	[49]
	Dunaliellabioculata	49	4	8	[50]
Dunaliella	Dunaliellasalina	57	32	6	[43]
	Dunaliellatertiolecta	64	21	15	[28]
Euglena	Euglena gracilis	39-61	14-18	14-20	[43]
Chlorococcum	ChlorococcumLittorale	38	23	16	[51]
Oscillatoriopsis	Microcystisaeruginosa	31	12	13	[52]
*	Nannochloropsisoculata	42.6	6	24	[46]
	Nannochloropsissalina	37	33	12	[53]
Nannochloropsis	Nannochloropsis sp.	52	12	28	[35,54]
	Nannochoropsis oc.	57	8	32	[21,27]
	Nannocloropsisoculata	39	20	17	[55]
Porphyridium	Porphyridiumcruentum	28-39	40-57	9-14	[43,46]
Prymnesium	Prymnesiumparvum	28-45	25-33	22-38	[50]
	Scenedesmusdimorphus	8-18	21-52	16-40	[50]
	Scenedesmusobliquus	50-56	10-17	12-14	[43]
Scenedesmus	Scenedesmusquadricauda	47	-	2	[50]
	Scenedesmus sp.	60	10	20	[56]
Spirogyra	Spirogyra sp.	6-20	33-64	11-21	[43]
	Spirulina	55-70	17-23	4-13	[46,57-59]
	Spirulina maxima	60-71	13-16	6-7	[43]
Spirulina	Spirulinaplatensis	46-63	8-14	4-9	[43,44,53,60]
	S. platensis	48	30	13	[61]
	Lyngbya sp.	30	13	1	[48]
Synechococcus	Synechoccus sp.	63	15	11	[50]
Platymonas	Tetraselmismaculata	52	15	3	[50]

Table 1 Main composition in the dry biomass of various microalgal species (wt.% dry weight)

2. Characteristic components of microalgal biomass

Microalgal biomass has a number of advantages over other biomass types; these advantages include higher area yields, 5 higher oil content, lower water consumption, and ease-of-growth on non-arable lands. To this end, microalgal biomass is an intriguing feedstock for the production of biofuel.

Microalgae are photosynthetic microorganisms that can produce three major biochemical components; namely, lipids, ¹⁰ proteins and carbohydrates (Fig. 2, Table 1). The primary elemental constituents of algae, which can be converted into fuels, are C, H, O and N. This is an important distinguishing feature of microalgae.^{39,40} Compared to lignocellulosic biomass, for example, these functional groups afford the potential to be ¹⁵ made into high value-added specialty chemicals.

Lipids account for 7-23 wt.% of the weight of microalgae under specific cultural conditions, such as high C/N medium, or conditions of stress.⁴⁰ Microalgal lipids comprise saturated and polyunsaturated fatty acids with typically 14 to 20 carbon units

²⁰ for the former and >20 for the latter. Typically, lipids are in the form of triglycerides which are used to produce biodiesel via the transesterification.⁴¹ Conventional microalgae-to-biodiesel technology generally requires high-lipid strains, but these strains tend to have lower biomass productivity compared to low-lipid ²⁵ algal strains.⁴² Thus, low-lipid microalgal strains require

alternative conversion strategies.

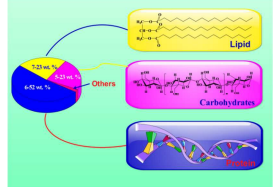


Fig. 2 Composition of microalgae with lipids, carbohydrates, proteins, and others represented

³⁰ Proteins make up 6-52 wt.% of the weight of microalgae.⁴⁰ Proteins, which are in the form of amino acids (Fig. 2), are the main source of nitrogen in microalgae (Table 1). Purified proteins are used in the food, animal feed, health, and specialty chemical markets. On the other hand, the protein fraction means that

35 conversion processes necessarily involve the nitrogen removal. This complicates any process of convert microalgae to liquid fuels.

Microalgae can accumulate a high carbohydrate content due to their relatively high photosynthetic efficiency.⁴ Carbohydrates, ⁴⁰ which are arguably the most important sources of energy and

- biological nutrients, comprise 5-23 wt.% of the algal feedstock.⁴⁰ Carbohydrates are homopolymers consisting of D-glucopyranose units linked via β -glycosidic bonds and/or α -glycosidic bonds (Fig. 2), and can be deconstructed into glucose monomers.
- ⁴⁵ Carbohydrates, accumulate in the plastids as reserve materials (starch), or become the main component of cell walls (cellulose, pectin, and sulfated polysaccharides).⁷

Overall, microalgal biomass belongs to a complex feedstock containing a large number of molecular functionalities that can be 50 exploited in a variety of uses. Indeed, the extraction and

- purification of the lipid, protein, and carbohydrate components are carried out through various chemical, thermal, and microbiological processes. A major challenge is to synthesize the desired products with sufficient purity and yield. Thus, the ss selectivity control and feedstock utilization are paramount. As for
- liquid fuels production, the major challenge in microalgal biomass conversion can be described as follows: on the one hand, much of the heteroatoms (O, N and S) in the microalgal biomass are removed. On the other hand, much of C and H remain in the 60 residue, and a large proportion of the remaining compositions are converted into liquid fuel.

For this reason, notwithstanding the obtained bio-oils from algae were similar with that of some lignocellulosic biomass via thermochemical conversion, algal bio-oils contained a number of ⁶⁵ N-compounds (most likely from protein degradation). In addition, compare to lignocellulosic biomass, there is no lignin existed in algal cell, so fewer aromatic compounds and their derivatives were present in algal bio-oils.

3. Strategies of microalgal conversion ⁷⁰ to liquid fuels

The growth of microalgae in water and its complex makeup distinguishes microalgae from conventional fossil fuels and other biomass types. With microalgal biomass being highly functionalized, conversion to a hydrocarbon-like fuel means that 75 energy-intensive defunctionalization steps are needed. For example, the removal of oxygen requires deoxygenation through the presumed use of a reductant. Such conversion steps involve a different set of challenges compared to the refining of coal, petroleum, or natural gas. The conversion processes must be done 80 selectively and efficiently to produce fuels with sufficient energy content while minimizing energy consumption.⁹ Unlike its

content while minimizing energy consumption.⁷ Unlike its lignocellulosic counterparts, algal biomass typically grows in water and must therefore requires extensive drying prior to further conversion.¹ Moreover, its aforementioned high protein ⁸⁵ content means that conversion processes requires the removal of

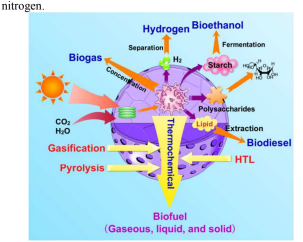


Fig. 3 Potential strategies for the production of fuels and chemicals from microalgae

Given the unique make-up of microalgae, researchers have investigated a variety of conversion methods that concentrate on

specific products.⁶² As depicted in Fig. 3, conversion methods include the production of biogas via concentration and/or catalysis, of biohydrogen via gasification and separation, bioethanol via fermentation, of monosaccharide via hydrolysis as 5 platform chemicals, and of biodiesel via extraction and

transesterification.^{8,11,13,63-78} Each approach exploits specific chemical functionality to produce different chemicals.

In contrast, thermochemical conversion has been devoted to transforming the entire algae for the production of ¹⁰ biofuel.^{1,8,11,12,14,16,25-32,35,39,40,42,47-60,79,80} Thermochemical conversion can be subdivided into gasification, pyrolysis, and hydrothermal liquefaction (HTL).^{81,82} Pyrolysis and HTL are two key routes converting microalgae into liquid fuels. During pyrolysis, microalgae as feedstock are heated in the absence of ¹⁵ oxygen to form bio-oil, solid char and gaseous products. On the other hand, HTL processed the microalgae at moderate temperatures produces liquid fuels in sub/supercritical water. In

contrast to pyrolysis, HTL has the advantage of minimizing undesired cross-linking related reactions because the solvent ²⁰ dilutes the concentration of the products at relatively low temperatures.

4. Biofuel production from microalgae via thermochemical conversion

The conversion of microalgae to biodiesel via the ²⁵ conventional transesterification route relies on microalgae with a sufficiently high fraction of lipids.²⁰ However, lipids comprise only a fraction of the total organic content of microalgae. The residual components consisting of proteins, carbohydrates, and unutilized lipids can be used to produce liquid, gaseous fuels ³⁰ and/or solid residue.⁸³ Lipid extraction and conversion is not viable for low lipid strains. Rather, thermochemical conversion of

the whole algae is carried out for biofuels production.^{31,36,42,83,84} Thus, thermochemical conversion is an option to process lowlipid microalgae or post-extraction residues of high-lipid ³⁵ microalgae (Fig. 4).

The thermochemical conversion (pyrolysis and HTL) of microalgae spans thermal decomposition and chemical reformation of the organic matter into biofuel.⁸⁵ A comparison of pyrolysis and HTL is provided in Table 2.³⁸ Pyrolysis refers to the ⁴⁰ anaerobic thermal decomposition of organic compounds into a mixture of gases, liquid, and chars.⁸⁶ Traditional pyrolysis is noncatalytic while recent attention has focused on the downstream catalytic upgrading of pyrolysis oils. HTL involves

the reaction of microalgae in a solvent at elevated temperatures 45 and pressures with or without catalyst.³¹ Pyrolysis is characterized by short gas residence times, operation at atmospheric pressure and relatively high temperatures. Prior to the pyrolysis, the algal feedstock must undergo dewatering and drying steps, which includes sedimentation, flocculation, dissolved air flotation, 50 filtration, and centrifugation. Drying is one of most dominant costs for algae harvest and may account for 30% of the total product costs, and the power consumption was equivalent to 15.8% of the energy of the recovered hydrocarbon. Energy costs climb steeply when the concentration of the slurries increases. In 55 contrast, HTL is usually performed at lower temperatures with longer residence time and much higher pressure. HTL converts the algal feedstock into biofuel in an aqueous phase, obviating the dewatering and drying procedures. Therefore, HTL is ideally suited to conversion of wet microalgae because it is tolerant to 60 the high moisture content of the feedstock.^{25,87}

In addition, HTL results in a higher quality of biofuel whereas pyrolysis, with the exception of fast pyrolysis, results in higher yield (Table 2).³⁸



Fig. 4 Possible microalgal processing options based on lipid content

This section focuses on pyrolysis and HTL process for the production of bio-oil from microalgae in detail. Published studies focused on microalgal conversion via pyrolysis and HTL to produce bio-oil with or without catalysts fall into this scope. ⁷⁰ Aside from the conventional thermochemical conversion, other assisted technologies, such as microwave assisted process and co-process will also be presented for comparison.

Madaala	The design of the second se	Reaction mechanism/process	Technique feasibility		
Methods	Treatment condition/requirement	description	Superiority	Drawback	
	Relatively high temperature (723-773 K);	Light small molecules are converted to oily	High biofuel yield up to	Feedstock need to be	
Pyrolysis	short residence time (~1 s); atmosphere	products through homogeneous reactions in	80 wt.% on dry feed;	dried prior to use. Poor	
	pressure; drying necessary	the gas phase	low capital cost	biofuel quality obtained.	
	L		Available for commercia	l Relatively low biofuel	
	Low temperature (573-673 K); long	Occurs in aqueous medium which involves	use; better quality of	yield (20-60 wt.%); need	
HTL	residence time (0.2-1.0 h); high pressure	complex sequences of reactions	biofuel (high HHV,	high pressure equipment	
	(5-20 MPa); drying unnecessary		low moisture content)	thus higher capital cost	
te: the bio-oil	yield from pyrolysis includes the ad	queous fraction whereas HTL excludes	the water soluble pro	ducts (aqueous fractio	

Table 2 Comparison of pyrolysis and HTL for biofuel production³⁸ Copyright 2012 Elsevier

75 4.1 Pyrolysis of microalgae for bio-oil production

Pyrolysis is a thermochemical change of organic matter in a

heated enclosure, usually in an oxygen-absent or very low oxygen level environment, to form a mixture of gases, liquid, and solids residue.⁸⁶ Most earlier applications have utilized coal and peat as feedtocks while lignocellulosic biomass has attracted attention in recent years.⁸⁸⁻⁹⁶ More recent studies have shown that pyrolysis of microalgae can produce bio-oil that is superior to bio-oil produced from lignocellulosic biomass.^{1,11-19} The pyrolysis of ⁵ microalgae aims at maximizing the production of energetically exploitable liquid and gaseous products.

An optional reaction for microalgae pyrolysis is conducted in a fixed-bed reactor that is externally heated by an electrical furnace with the temperature measured by a thermocouple

- ¹⁰ positioned inside the bed (Fig. 5).⁵⁶ The experiment steps are described as follows: (1) The reactor was filled with a certain amount of microalgal biomass that had been dried and ground to a particle size. (2) The reactor system was purged with carrier gas (e.g., N_2) which was also used during the reaction. (3) With the ¹⁵ carrier gas flown, the reactor system was heated at a certain
- heating rate until reached the reaction temperature, and then held for some minutes. (4) The product vapors were collected through condensation in flasks cooled by an ice bath. (5) The aqueous and organic phases were separated, and then detected using some 20 analysis methods (such as GC/MS, FT-IR, and elemental

analysis) to evaluate the bio-oil composition.

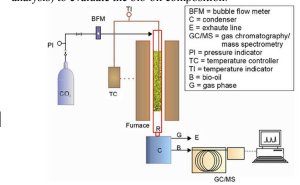


Fig. 5 Schematic reactor system for microalgae pyrolysis⁵⁶ Copyright 2012 Elsevier.

25 4.1.1 Pyrolysis behavior of microalgae

A thorough knowledge of the pyrolysis behavior of microalgae is critical to evaluate the potential and feasibility of pyrolysis for microalgal conversion. Questions such as at what temperature does algae pyrolyze, what is its major product, what ³⁰ is the bio-oil yield, and what is the effect of key operating

- parameters are practical questions are not easily answered, and involve complex underlying phenomena.
- Thermogravimetric analysis (TGA) is an established method for studying the thermal degradation mechanisms and ³⁵ kinetics of microalgae.^{40,44,45,97-103} TGA provides semiquantitative information about the pyrolysis temperature and kinetics, and phase distribution of the products, notwithstanding its rather unrealistic gas-solid contacting design. To illustrate the information gleaned from TGA, the pyrolysis of two microalgal
- ⁴⁰ species (*Spirulina platensis* and *Chlorella protothecoides*) were carried out by a conventional temperature ramp.^{44,97} TGA revealed the maximum weight loss from these two microalgae occurred over a several hundred degree temperature range (~420-830K) with minor differences for each species. An increase in the
- 45 heating rate caused a shift in the degradation to higher temperatures which was interpreted with a simple model that inferred a decrease in the activation energies for the

devolatilization stage and an increase in both the instantaneous maximum and average reaction rates.

- In general there are three primaries stages that occur during the pyrolysis of microalgae. Zou *et al.*⁹⁸ carried out the thermal pyrolysis of *D. tertiolecta*. They affirmed that moisture was removed in the first stage; most of the pyrolysis occurs in the second stage in which most of the organic material was ⁵⁵ decomposed based on the largest decrease in mass; remaining solid was slowly decomposes in the third stage. The initial temperature of pyrolysis and the temperature at which the pyrolysis rate reaches the peak value shifts to the higher range as the heating rate increased. Similar pyrolysis features have been ⁶⁰ reported for *Nannochloropsis gaditana*, *Chlorella spp.*,
- Nannochloropsis, Potamogeton crispus, Sargassum thunbergii, C. vulgaris, and Nannochloropsis sp.^{40,45,99-101}

The pyrolysis behavior of the common *Chlorella* species obtained using TGA show that the volatile species consist ⁶⁵ primarily of water and CO₂, as well as H₂ at high temperatures.¹⁰² The thermal pyrolysis of *Nannochloropsis sp.* indicated the different biochemical components of the microalgae in the absence of a catalyst seem to be decomposed in the following order: carbohydrates, proteins, and lipids.⁹⁹ The addition of ⁷⁰ catalysts with the microalgae can have an important effect in the presence of Na₂CO₃ during the pyrolysis process, the main weight decrease shifted to a lower temperature. This indicated that Na₂CO₃ may affect the decomposition of carbohydrates and proteins but not that of lipids. The Na⁺ from Na₂CO₃ can ⁷⁵ penetrate the biomass and break the hydrogen bonds, enabling pyrolysis to occur at lower temperatures.

The thermal behavior of six microalgal species (*Tetraselmischui*, *Chlorella like*, *C. vulgaris*, *Chaetocerous muelleri*, *D. tertiolecta* and *Synechococcus*) showed the results as follows: First, the ratio of evolved liquid, gas and char products varied markedly across all species with temperatures up to 773 K; second, the rate and temperature of evolution of these fractions was also inconsistent; third, the mix of combustible volatile gas compounds that evolved with temperature was also variable, resulting in differences in the its energy content (higher heating value, HHV).¹⁰³

In summary, the pyrolysis behavior of microalgae obtained from TGA can provide useful information for laboratory-scale pyrolysis. However, the measurements on pyrolysis behavior of microalgae have its limitations due to the unrealistic gas-solid contacting. These limitations activate the incentive to devise and apply alternative analysis methods that enable bridging of the gap between the lab- and full-scale pyrolysis, thereby enabling a fundamental study of pyrolysis.

95 4.1.2 Noncatalytic pyrolysis of microalgae

Noncatalytic pyrolysis, also known as direct pyrolysis, refers to the process in which microalgae are thermally degraded at moderate temperatures (623-973 K) in the absence of catalyst.¹⁰⁴ The heating rate and temperature have a significant ¹⁰⁰ effect on the products and yields. In general, at sufficiently high temperature the degradation process results in the production of pyrolysis vapor which upon downstream condensation yields a dark viscous fluid called bio-oil or pyrolysis oil, while the nonvolatile solid phase is called biochar.¹⁰⁴ Pyrolysis processes ¹⁰⁵ are often classified by their heating rate, with rates of 0.1-1 K ·s⁻¹

referred to as slow pyrolysis, whereas rates of 1-200 K \cdot s⁻¹ as fast pyrolysis (Table 3).^{81,82}

(1) Slow pyrolysis

In addition to the relatively slow heating rate, slow ⁵ pyrolysis is characterized by longer gas residence time.¹¹⁷ Slow pyrolysis of microalgae results primarily in the production of biochar and pyrolysis gas.^{102,105} CH₄ and CO₂ are the major components in the gaseous product.⁸⁶ Some examples follow to illustrate.

¹⁰ Slow pyrolysis of *C. protothecoides* resulted in a bio-oil

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yields exceeding 40 wt.%.¹⁰⁶ The gaseous product yield generally increased with temperatures and residence time due to the secondary reactions. When *Spirulina sp.* was used as feedstock via slow pyrolysis, the optimized temperatures with the highest ¹⁵ maximum biochar and bio-oil yields were 773 and 823 K, respectively.¹⁰⁷ In addition, the bio-oils obtained from slow pyrolysis of *Nannochloropsis sp.* Residue mainly consisted of long-carbon chain compounds with various terminal groups with an oxygen content of 30.1 wt.% and a HHV of 24.6 MJ·kg⁻¹.¹⁰⁸

	82	
Table 2	rating parameters, expected yields and product composition for pyrolysis processes ⁸² Copyright 2010 Elsevie	
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			Read	Reaction condition					
Pro	Process types		Heating rate/K·s	s ⁻¹ Temper	ature/K	с м	ain prod	lucts	
<u>Classa non a la si a</u>	Carbonization	Hours \sim days	Very low	573-	-673		Charcoa	al	
Slow pyrolysis	Conventional pyrolysis	5-30 min	0.1-1	573-	.973	Gas, liquid and charcoa		charcoal	
Frank manual la sia	Fast pyrolysis	0.5-5 s	1-200	773-	1073		Liquid		
Fast pyrolysis	Flash pyrolysis	<0.5 s	>10 ³	823-	1273	Liq	uid and/	or gas	
old et al. compar	ed the slow pyrolysis o	of Scenedesmus	feedstocks.	Moreover,	the	bio-oil	vield	increased	

Harold et al. compared the slow pyrolysis of Scenedesmus sp. with that of duckweed.⁵⁶ Scenedesmus sp. afforded a higher bio-oil yield than duckweed, whereas microalgal bio-oil had a higher HHV of 19 MJ·kg⁻¹ than that of duckweed bio-oil (15 ²⁵ MJ kg⁻¹). The authors stated that the thermolysis of proteins, carbohydrates and lipids required less stringent conditions than that of lignocelluloses and resulted in a product with higher HHV. In addition, they affirmed several reactions during the thermolysis including deamination, direct process methylation, 30 decarboxylation (DCO), dehydration, decarbonylation, cyclization, dimerization, and homonolysis. Another important reaction occurred in this process is Maillard reaction, which converts calloung with the second se converts carbohydrates and proteins to form amadori compounds,

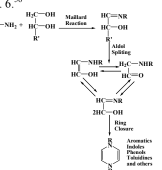


Fig. 6 Proposed reaction mechanism leading to the formation of amadori compounds⁵⁶ Copyright 2012 Elsevier.

Slow pyrolysis has the noted disadvantage of leading to secondary cracking, condensation and polymerization of ⁴⁰ pyrolysis products. These reactions lead to decreasing the bio-oil yield and have adverse effects on the bio-oil properties. Moreover, the lower HHV and the longer residence time tend to increase the energy requirements.

(2) Fast pyrolysis

- ⁴⁵ Since the aim for microalgal pyrolysis is the production of organic liquid phase (biofuel), fast pyrolysis is recommended (Table 3). The achievement of heating rates as high as 200 K·s⁻¹ requires high operating temperatures, short residence time, and fine particles (<1 mm).¹⁰⁴
- ⁵⁰ The fast pyrolysis of dried, ground *Scenedesmus sp.* resulted in a bio-oil which rivaled that from lignocellulosic

temperature up to a point and then decreased.¹¹⁰ The rather high bio-oil yield from microalgae suggests that fast pyrolysis is a ⁵⁵ potential method for converting algae to liquid.¹⁰⁵ When the conversion of *C. prothothecoides* and *M. aeruginosa* was carried out with fast pyrolysis, the bio-oil yields were 18 and 24 wt.%, respectively.⁵² Additionally, the bio-oil from fast pyrolysis of microalgae has a HHV of 29 MJ·kg⁻¹, which is about 1.4 times of ⁶⁰ that of wood. Liquid fuels from fast pyrolysis of microalgae may be used in many applications as direct substitutes for conventional fuels.⁵²

The bio-oil yield ranged from 25-30 wt.% with a HHV of 25 MJ·kg⁻¹ in the fast pyrolysis of microalgae in a falling solids ⁶⁵ reactor.¹¹¹ Fig. 7 shows the GC-MS-classified organic fractions of bio-oils obtained during pyrolysis in three atmospheres (N₂, steam and CO₂). The data indicate that the pyrolysis atmosphere has an important effect on the product distribution, and the presence of steam increases the fraction of hydrocarbons while ⁷⁰ decreases the oxygenate fraction. The C fraction from the steam swept pyrolysis exceeds that from the N₂-swept pyrolysis while the O fraction is lower. These trends reflect in a higher HHV for the bio-oil during the steam swept process. The authors speculated that the reactions occurring during steam pyrolysis are ⁷⁵ presumably steam reforming and deoxygenation.¹¹¹

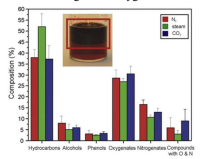


Fig. 7 Bio-oil composition of pyrolysis employed at three atmospheres: N_2 , steam, and CO_2^{111} Copyright 2012 Elsevier.

Process conditions: 773 K, N₂ flow=250 mL·min⁻¹, algae mass=7 g, dp<90mm

⁸⁰ The pyrolysis temperature plays a crucial role on the pyrolysis product distribution of blue-green algae blooms.¹¹² The maximum bio-oil yield of 55.0 wt.% was obtained at a pyrolysis temperature of 773 K, particle size below 0.25 mm and a sweep

gas flow rate of 100 mL·min⁻¹. The bio-oil has a HHV of 31.9 MJ·kg⁻¹ and an O/C molar ratio of 0.16 under optimum conditions. These results showed that the pyrolysis of algal biomass is a promising process for producing renewable fuel ⁵ while improving the quality of a freshwater lake.¹¹²

Pyrolysis through microwave-assisted power has been proposed for both microalgae and their extraction residues to obtain bio-oil.¹¹³ Microwave-assisted pyrolysis (MAP), initially developed by Tech-En Ltd. in Hainault, UK, has been

- ¹⁰ investigated in recent decades.^{114,115} Compared to the traditional processes, MAP offers several advantages including uniform internal heating of large biomass particles, ease of control, no need for agitation or fluidization and fewer particles (ashes) in the bio-oil.¹¹⁶
- ¹⁵ Noteworthy is the enhancing effect of the microwave power on the pyrolysis product yield. A maximum bio-oil yield of 32.0 wt.% was obtained at a microwave power of 750 W in the MAP of *Chlorella sp.* When the material was changed into *C. vulgaris*, it was found that a microwave power of 2250 W gave the highest
- ²⁰ bio-oil yield of 74.9 wt.%; a bio-oil yield as high as 87.4 wt.% was obtained when activated carbon was added as a catalyst.¹¹⁷ Notwithstanding current MAP techniques offering numerous advantages and showing excellent potential for enhancing microalgal bio-oil yield, the growth of industrial microwave
- ²⁵ heating applications is hampered by the lack of knowledge on the microwave systems and commercial necessary equipment for these pyrolysis processes. In addition, the economic assessment of the MAP has yet not been conducted.
- Fast pyrolysis requires a reactor configuration in which the ³⁰ residence time of microalgae is of order only a few seconds. As mentioned earlier, fast pyrolysis seems to be a viable technique for future replacement of fossil-fuel with biomass-derived liquid fuels because of the potential for high biomass-to-liquid yield. However, some technical challenges need to be solved because
- ³⁵ bio-oils from direct pyrolysis are acidic, unstable, viscous, and contain solids and chemically dissolved in water. Therefore, the bio-oil must be upgraded via hydrogenation or catalytic cracking to decrease oxygen content and remove alkalis.

4.1.3 Catalytic pyrolysis of microalgae

- The objective of pyrolysis is to convert the algal biomacromolecules into lower molecular weight species through a series of reactions including decarbonylation, dehydration and aromatization reactions. However, as stated earlier, the pyrolysis oil has the drawbacks of high oxygen content, high acidity, high
- ⁴⁵ viscosity, and a relatively low specific energy content, which inhibits its further application as a transportation fuel. Improvement of the bio-oil stability, decrease of the acidity and increase of the energy density can be achieved by removal of oxygen and nitrogen that cause these problems. The use of a
- ⁵⁰ catalyst may redirect the chemical reactions during the pyrolysis process result in the *in situ* upgrading of the bio-oil.¹⁰²

Compared to the noncatalytic pyrolysis of algal biomass, catalytic pyrolysis can not only upgrade the quality of bio-oil but also adjust the components of bio-oils to meet different ⁵⁵ demands.¹⁰⁸ This may be accomplished either by mixing biomass with the catalysts¹¹⁸ or flowing pyrolysis vapors over a catalyst positioned downstream from the pyrolysis zone. Catalytic pyrolysis generally produces biofuel of enhanced quality (higher

HHV, lower oxygen content, and higher aromatic hydrocarbon ⁶⁰ content) even at atmospheric pressure without the need for a reductant, which makes this a cost-effective upgrade.¹¹⁹ Previously, most of the applied catalysts focused on molecular sieves in the catalytic pyrolysis process of algae. However, the investigation of other catalysts may be considered to ameliorate ⁶⁵ biofuel quality.¹²⁰ Here we highlight some recent examples.

(1) Alkalis salt catalysis

Babich *et al.* studied the pyrolysis of *Chlorella* both with and without Na₂CO₃ as the_catalyst.¹⁰² The presence of the Na₂CO₃ catalyst lowered the initial degradation temperature, and ⁷⁰ produced a bio-oil with lower acidity and higher HHV than biooils produced without the catalyst. However, the Na₂CO₃ also promoted the gas yield and reduced the liquid yield.

(2) Molecular sieve catalysis

Early in 1990, Milne et al. first proposed the catalytic 75 conversion of entire microalgae over HZSM-5molecular sieve, but the obtained results were ambiguous.¹²¹ The formation of high value-added aromatics is enhanced considerably, which can be attributed to the acidity of the zeolite catalysts in the catalytic pyrolysis of Laminaria japonica over microporous zeolite ⁸⁰ catalysts (HZSM-5, Hβ and HY).¹²² HZSM-5, which, together with its Brönsted acidity and specific pore structure, showed the highest selectivity for aromatic production. Compared to noncatalytic pyrolysis, the catalytic pyrolysis of microalgae produced lower bio-oil yields because of the catalytic cracking of 85 bio-oil compounds to form gaseous products.¹²² Similar results were reported by Wang,⁴⁷ Du¹²³ and Gopakumar¹²⁴ when HZSM-5 was used for the catalytic pyrolysis of C. vulgaris. An increase of the aromatic hydrocarbon yield of 0.9 to 25.8 wt.% was achieved while the oxygen content in turn decreased from 30.1 to 90 19.5 wt.% and the HHV increased from 24.4 to 32.2 MJ·kg⁻¹.124 The catalytic pyrolysis of microalgal biomass produces more monocyclic aromatics than that of lignocellulosic biomass.⁴⁷ Pan

et al. pyrolyzed *Nannochloropsis sp.* using variable amounts of HZSM-5 over a range of temperatures.¹⁰⁸ The catalyst increased ⁹⁵ the HHV of the bio-oil from 24.6 to 32.7 MJ·kg⁻¹. The bio-oil obtained from catalytic pyrolysis is rich in aromatic hydrocarbons based on the GC-MS results.

A detailed study comparing noncatalytic and catalytic pyrolysis using exchanged cations ZSM-5 catalysts was carried ¹⁰⁰ out by Campanella and Harold.¹¹¹ Noncatalytic pyrolysis gave the highest total liquid content, whereas catalytic pyrolysis resulted in the highest hydrocarbon fraction. A comparison of four exchanged ZSM-5 catalysts (H-, Fe-, Cu- and Ni-) showed differences in the bio-oil yield and composition (Fig. 8).¹¹¹ 105 Among the studied catalysts, H-ZSM-5 gave the largest enhancement in the liquid product yield.¹¹¹ Fig. 8a shows the effect of these catalysts on the bio-oil yields under identical operating conditions. The solid residue vield remained unchanged or slightly decreased. The volatile product yield increased at the 110 expense of a decrease in the pyrolysis oil (bio-oil plus aqueous fraction). These results provide evidence for desired deoxygenation chemistry that produces CO, CO₂ and light hydrocarbons. Further, GC-MS measurements demonstrated that the zeolites catalyst enhanced the energy content associated with

¹¹⁵ the more favorable bio-oil composition (Fig. 8b). Notable trends included a decrease in the fraction of oxygenated species and an

increase in the fraction of hydrocarbons; meanwhile, the yield of nitrogen compounds remained the same or slightly decreased. Compared to noncatalytic pyrolysis, the catalytic pyrolysis provided an increase in phenols, which are high value-added ⁵ chemicals, and could increase the attractiveness of the catalytic pyrolysis of microalgae.¹¹¹ In addition, the enhanced aromatic fraction yield may owe to Dielse-Alder and condensation reactions.

In summary, the heteroatom derived from the microalgae

¹⁰ biochemical composition can only be partly removed in catalytic pyrolysis, while the noncatalytic pyrolysis followed by bio-oil upgrading (two-stage method) can remove almost all the heteroatoms to satisfy the requirements for use as transportation fuel. However, the two-stage method presents some ¹⁵ disadvantages, including numerous treatment steps, high storage cost, and low bio-oil yield, etc. Therefore, both of the two methods have their advantages and disadvantages.

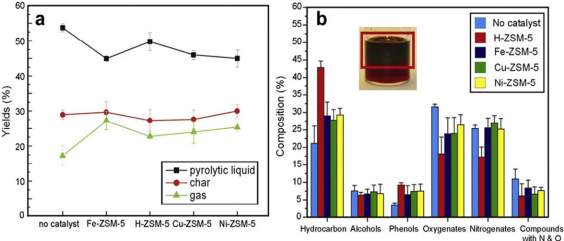


Fig. 8 Experimental results of catalytic pyrolysis reactions conducted using different zeolites¹¹¹ Copyright 2012 Elsevier. (a) Effect of catalyst on product yield in catalytic pyrolysis reactions. (b) Bio-oils product distribution.

 $Process \ conditions: \ 773 \ K, \ N_2 \ flow=250 \ mL \ min^{-1}, \ algae \ mass=7 \ g, \ WHSV=13.5 \ h^{-1}, \ GHSV=20000 \ h^{-1}, \ residence \ time \ \sim 1.5 \ s^{-1}, \ residence \ s^{-1}, \ ress \ s^{-1}, \ residence \$

4.1.4 Co-pyrolysis of microalgae with other substances

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As mentioned earlier, some challenges with bio-oils ²⁵ obtained from the pyrolysis of microalgae that may hinder their commercialization include high oxygen content, low HHV, low volatility, etc. Upgrading of these unstable bio-oils would require a downstream hydrogenation step, for example. Thus, there is a need to explore strategies of improving the bio-oil quality.¹²⁵ To

- ³⁰ this end, the co-pyrolysis of mixtures of biomass and other substances has received increasing attention.¹²⁶⁻¹³⁰ Co-pyrolysis of algae with other feedstocks offers an interesting approach to overcome certain disadvantages of algal-only feedstocks.^{129,131} Moreover, the co-pyrolysis may not require high-pressure
- ³⁵ hydrogenation, making the process safer. Attractive co-feeds include solid waste, coal, and even plastics. For example the transfer of hydrogen from a co-feed with a higher hydrogen-tocarbon ratio might enable de-oxygenation of the algal pyrolysis products leading to a higher quality biofuel.
- ⁴⁰ Tang *et al.* studied the co-pyrolysis characteristic of microalgae and municipal solid waste under N_2/O_2 and CO_2/O_2 atmospheres using TGA.¹³² As the blending ratio of microalgae increased from 10 to 70 wt.% under a N_2/O_2 atmosphere, the volatile release temperature decreased from 542 to 520 K, the
- ⁴⁵ temperature at the maximal peak decreased from 583 to 561 K, the maximum rate of weight loss decreased from 11.94 to 7.88 wt.% and the residual weight decreased from 30 to 20 wt.%.

The results from a TG- and modeling-based study using fresh water algae *Chlorococcum humicola* and a *Victorian* brown ⁵⁰ coal and their blends at different proportions showed that a

smaller amount of algae can be mixed with coal without

⁷g, whisv=13.5 if , Ohsv=20000 ff , residence time ×1.5 s
significantly changing in the pyrolysis characteristics.¹³³ Coal was mixed with algae to produce a slurry and then the combustion behavior of the coal-water slurry was investigated by Li *et al.*¹³⁴
⁵⁵ Yuan *et al.* studied the rapid pyrolysis of the aquatic biomass (blue-green algae and water hyacinth) with two coals (bituminous and anthracite).¹³⁵ During the co-pyrolysis of algal biomass and coal, the interactions between algae and coal decreased char-N yields and increased volatile-N yields, but the total yields of NH₃
⁶⁰ + HCN decreased. HCN formations consistently decreased, whereas NH₃ formations only decreased in the high-temperature range but increased in the low-temperature range.

4.1.5 Prospectives for pyrolysis of microalgae

Pyrolysis of microalgae faces many challenges that will ⁶⁵ require innovative solutions. But recent developments with catalytic pyrolysis, potentially with synergistic co-feeds, hold promise for eventual commercial application. Various life cycle analyses and related studies have been carried out for microalgae and other biomass feedstocks to produce biofuels.¹³⁶⁻¹³⁹ It is not ⁷⁰ our intent to do an exhaustive review but to provide some key highlights. The following are critical factors that will have to be addressed pertaining to microalgae.

- Energy content. While compared to other biomass sources, the energy content of algae on a dry basis is nominally half that of a fossil fuel. The rather high oxygen content is primarily responsible. The removal of oxygen through chemistries like decarbonylation is essential to reduce the reductant requirements for deoxygenation.
- Water content. Algae grow in water and therefore require dewatering and drying prior to pyrolysis. The latter is particularly energy intensive. The use of a fraction of the water during the pyrolysis as a source of hydrogen through reforming of pyrolysis products is one approach.

• Use of photo bioreactors (PBR). One way to address the land requirement is to grow algae in transparent vertical columns. Designs would be needed to maximize the utilization of sunlight and minimize the materials cost. The use of polymeric materials that withstand photocatalytic degradation is essential. The materials cost of the PBR is a

non-negligible factor in the overall economics.

Logistical issues. Notwithstanding the land and water requirements for growth, the infrastructure needed for harvesting and conversion would require a massive effort and investment. Bio-oil production facilities should be located in close proximity to the lands producing the algal biomass. A stabilized bio-oil could then be transported to refineries for conversion to transportation liquids. This poses a trade-off between transportation costs and economies-of-scale afforded by larger pyrolysis facilities.

4.2 HTL of microalgae for bio-oil production

As described above, microalgal pyrolysis converts whole algae into liquid fuels.²⁰ However, a large amount of water ²⁰ accompanies algae with typical algal cell density of 1 g·L⁻¹.¹ As a result, the economics of microalgal pyrolysis are undermined by the costly dewatering and drying steps.²⁸

- As alternative to pyrolysis is hydrothermal liquefaction (HTL) which involves conversion of microalgae in slurry ²⁵ comprising a liquid solvent and is carried out at moderate temperatures and sufficient pressure to keep the solvent in the liquid phase. HTL is of interest because it eliminates the need to expend the energy to dewater and dry algae, as required in other thermochemical conversions.¹⁴⁰ To this end, the conversion of
- ³⁰ algae via HTL has received increasing interest in recent years even though biomass HTL is a rather mature technology, first described in the 1940s, with technology improvements in the 1980s by Shell researchers.¹⁴¹

HTL utilizes a variety of solvents, including water as the ³⁵ reaction medium. The use of water of course presents several

advantages over other solvents because it is ecologically safer, cheaper, and readily available, not to mention that it is the growth medium for algae.¹⁴² It is noted that the ionic product of water under high temperature and pressure conditions below the critical ⁴⁰ point of water is up to three orders of magnitude higher than that under ambient conditions (Fig. 9).¹⁴³ A high ionic product is favorable for acid- or base- catalyzed reactions. In addition, water can act as an acid or base catalyst precursor due to the relatively high concentrations of H_3O^+ and OH^- ions from the self 45 dissociation.¹⁴³ When water is heated and compressed, the hydrogen bonds are weakened, resulting in a change in dielectric constant, acidity, and polarity, and increasing the reactivity of water. For example, the dielectric constant of water decreases from 78.85 to 13.96 when the temperature increases from 25 to 50 250 °C, resulting in the transition of water molecules from very polar to fairly nonpolar.¹⁴⁴ The dissociation constant of water (K_w) increases from 10⁻¹⁴ to 10⁻¹¹ just below 250°C, resulting in an enhancement in the rates of acid- and base-catalyzed reactions in water.145

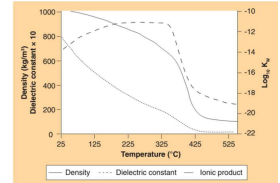
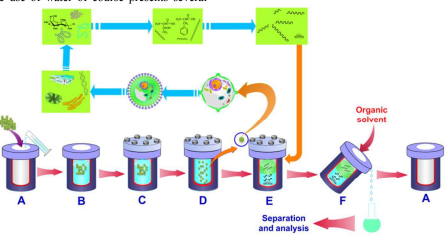


Fig. 9 Density, static dielectric constant at 30 MPa and ionic product of water at 25 MPa¹⁴³ Copyright 2012 Future Science Group



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Fig. 10 Experimental procedure of HTL for microalgae in closed batch reactor A: Reactor, B: Charging, C: Sealing, D: Heating, E: Cooling, F: Washing

HTL of microalgae proceeds through a multi-step procedure as depicted in Fig. 10, and are as follows. (1) During the initial stages the algal cell membrane and wall are disrupted chemically in the high temperature and pressure water. Numerous reactions

65 between organic compounds in the cell membrane and wall occur, and some small molecules are produced. (2) Once the microalgae are lysed, the intracell components, lipids, proteins, and carbohydrates, participate into the reaction process. Hydrolysis and depolymerization occur, converting carbohydrates into ⁷⁰ monosaccharides and polysaccharides, proteins into peptides and amino acids, and lipids into fatty acids and glycerol. (3) Repolymerization/self-condensation reactions occur. This includes the conversion of lipids into fatty acids, proteins into nitrogen heterocycles, pyrroles, and indoles; and carbohydrates ⁷⁵ into cyclic ketones and phenols. Others chemical reactions such as Maillard reaction between the smaller molecules may be also

present at this stage. (4) Following the liquefaction process, algae are eventually converted into a series of products including liquid, gas and solid residue. The long-chain nonpolar molecules generally named as bio-oil are formed. Short chain polar

- 5 molecules are dissolved into water medium and the formed aqueous phase, which is mostly used to cultivate microalgae. The gaseous products, mainly CO₂, are directly vented to the atmosphere in most cases.
- In real HTL process, bio-oil can be separated easily from 10 water phase with the addition of the organic solvents (such as trichloromethane, dichloromethane, tetrahvdrofuran, and nhexane). The separation procedure of HTL products is described in published papers.^{42,49,53,54}
- Microalgal derived HTL bio-oil has several desirable 15 attributes. First, the HTL conversion of microalgae into bio-oil has an efficiency of 30-75% and a net positive energy yield that is 3-10 times of the input heat energy.¹⁴⁶ Based on the principle of green chemistry, Zhang et al. proposed the concept of Environment-Enhancing-Energy (E²-Energy) to integrate bio-oil
- ²⁰ production and wastewater treatment (Fig. 11).⁴² According to E²-Energy, microalgae are grown to serve as a wastewater treatment method to uptake nutrients and capture CO₂ from HTL products.¹² Subsequently, the resulting algae are further converted into bio-oil via HTL. The algae species that grow in the post-HTL

25 water are expected to have low lipid content because the N and C contents of the HTL aqueous stream are high.⁴² That said, the biooil obtained from the HTL of microalgae still contains a fraction of O and N. Such bio-oil has a lower quality and HHV compared to the actual transportation fuel.

To improve the quality of bio-oil, two methods have been 30 developed. One method involves HTL with homogeneous and/or heterogeneous catalysts. The other method involves the use of organic solvents or co-processing with other substances for microalgal HTL. In this section, we highlight the developments 35 of noncatalytic HTL firstly, followed by catalytic HTL, and then by HTL in co-solvents. In addition, selected contents that elucidate the mechanism of microalgal HTL also fall within scope of this section.





					Conditions			
Туре	Ref.	Catalysts	Feedstocks	Reaction medium	Temperature (K)	Holding Tim (min)	- Max. bio-oil yield (wt.%)	Max. HHV (MJ·kg ⁻¹)
	[148]	/	Dunaliella tertiolecta	Water	633	30	36.9	26.6
	[54]	/	Nannochloropsis sp	Water	473-773	60	43.0	39.0
-	[49]	/	Desmodesmus sp.	Water	448-723	5-60	49.0	36.0
	[53]	/	Spirulina Nannochloropsis salina	Water	493-648	30	38.0 46.0	35.2 38.1
	[149]	/	Nannochloropsis sp.	Water	873	1	66.0	37.0
Noncatalytic	[86]	/	Scenedesmus Spirulina	Water	573	30	45.0 31.0	35.5 35.8
Liquefaction	[42]	/	Chlorella pyrenoidosa	Water	553	120	39.4	35.4
(direct	[150]	/	Chlorella pyrenoidosa	Water	553	120	39.4	35.4
liquefaction)	[58]	/	Spirulina	Water	573	30	32.6	34.7
	[151]	/	Spirulina platensis	Water	623	60	39.9	39.9
	[61]	/	Spirulina platensis	Water	623	60	41.0	34.2
	[57]	/	Spirulina	Ethanol	633	-	35.4-45.3	32.6
_	[59]	/	Spirulina	Ethanol Methanol 1, 4-dioxane	653	20	54.0 55.1 56.6	38.3 39.8 36.8
	[152]	/	Chlorella pyrenoidosa	Ethanol	443-643	5-120	64.6	38.9
	[29]	Na ₂ CO ₃	Botryococcus braunii	Water	473~613	60	64.0	-
	[28]	Na ₂ CO ₃	Dunaliella tertiolecta	Water	473~613	5 and 60	43.8	36.0
	[153]	Na ₂ CO ₃	Enteromorpha prolifera	Water	493-593	5-60	23.0	30.0
Homogenous	[33]	Na ₂ CO ₃	Microcystis viridis	Water	573 and 613	30 and 60	33.0	30.0
catalysis	[32]	Na ₂ CO ₃	Dunaliella tertiolecta	Water	633	50	25.8	30.7
	[31]	Na ₂ CO ₃ KOH CH ₃ COOH	Chlorella vulgaris Spirulina	Water	573 and 623	60	27.3 20.0	37.9 39.9

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		НСООН						
		Na ₂ CO ₃	Chlorella vulgaris, Nannochloropsis occulata		623		35.8 34.3	37.1 34.3
	[27]	НСООН	Porphyridium cruentum	Water		60	20.0	36.3
			Spirulina				29.0	36.8
		Na ₂ CO ₃					51.6	36.3
	[60]	$Ca_3(PO_4)_2$	Spirulina platensis	Water	573-623	30-60	34.5	38.4
_		NiO					30.2	35.1
		FeSO ₄						
	[57]	FeS	Spirulina	Ethanol	633	_	46.0	37.1
	[37]	Na ₂ CO ₃	spiruuna	Ethalion 055	055		40.0	57.1
_		NaOH						
			Chlorella vulgaris				27.0	33.2
	[27]	[27] HCOOH ¹	Spirulina	Water	623	60	29.0	35.1
	[2,]	neoon	Nannochloropsis occulta		025	00	26.0	39.6
_			Porphyridium creuntum				27.1	36.3
	[154]	H_2SO_4	Sargassum polycystum	Ethylene glycol	443	15	87.7 ^a	-
	[155]	H_2SO_4	Dunaliella tertiolecta	Ethylene glycol	443	33	45.0	28.4
	[86]	HZSM-5 Raney-Ni	Chlorella pyrenoidosa	Ethanol	473-573	30	71.3	36.2
-		Pd/C Pt/C						
	[35]	Ru/C Ni/SiO ₂ -Al ₂ O ₃	Nannochloropsis sp.	Water	623	60	57.0	38.0
Heterogeneous		CoMo/y-Al ₂ O ₃						
catalysis		zeolite						
	[156]	Ni/REHY	Dunaliella salina	Water	473	60	72.0	30.1
_		Co/Mo/Al ₂ O ₃	Chlorella vulgaris				38.7	39.7
	[21]	Ni/Al/Al ₂ O ₃	Nannochloropsis occulta	Water	623	60	30.0	42.0
_		Pt/Al/Al ₂ O ₃	ivannocnioropsis occulta				38.9	38.2
	[157]	Fe(CO) ₅ -S	Spirulina	Water	573 and 613	30 and 60	78.3	33.0

4.2.1 Noncatalytic HTL of microalgae

The high moisture content and their small particle size of microalgae (ca. microns) makes HTL a convenient microalgal s conversion process.¹⁵⁸ Over the past years, a variety of microalgal species have been studied as feedstocks for HTL to bio-oil (Table 4). Some of these species include *B. braunii*, *D. tertiolecta*, *D. salina*, *C. vulgaris*, *N. occulta*, *S. platensis*, and *Spirulina*, etc.^{21,28,29,31,32,151,156} A wide range of operating conditions have led to highly variable bio-oil yields and HHVs as well as overall energy balances compared to pyrolysis oil. In general, HTL of microalgae has resulted in bio-oil yields that are 5-30 wt.% than the initial lipid content.^{27,28,31,32,54,1451,159,160} This is a result of the conversion of other microalgal components into bio-oil.^{151,161,162}

¹⁵ The produced gas from HTL consists mainly of methane and carbon dioxide, such as the study of HTL of *B. braunii*.¹⁶³ Similar results were reported by Minowa *et al*.²⁸ who obtained a bio-oil yield of 37 wt.% with a HHV of 36 MJ·kg⁻¹ from *D. tertiolecta*. Other low-lipid microalgae, like *Spirulina*, were used

²⁰ as the feedstock, and the biofuel yield could reach as high as 78.3 wt.%.¹⁵⁷ A bio-oil yield of 49 wt.% was obtained from the HTL of *Desmodesmus sp.* as the feedstock. About 75 % of the HHV (22-36 MJ·kg⁻¹) in microalgae was transformed into bio-oil.⁴⁹

Researchers have reported that the highest bio-oil yields are ²⁵ obtained during HTL under subcritical conditions. Brown *et al.* converted the microalga *Nannochloropsis sp.* into bio-oil via HTL at different temperatures.⁵⁴ The highest bio-oil yield of 43 wt.% was obtained at 623 K which corresponded to subcritical water. The bio-oil yields decreased from this maximum value in the ³⁰ 673-723 K temperature range. By 773 K, the bio-oil yield was nearly half the maximum value as a result of oil-range molecules reacting to form lighter and more volatile compounds which are not captured in the oil fraction. Moreover, at higher temperatures (supercritical conditions), higher molecular-weight compounds ³⁵ derived from oil-range molecules react together to form solid products.⁵⁴

Zou *et al.* studied in detail the factors influencing HTL of the microalga *D. tertiolecta* for the production of bio-oil under various conditions.¹⁴⁸ The maximum bio-oil yield was ⁴⁰ approximately 36.9 wt.% at a reaction temperature of 633 K and a holdup time of 30 min, with a feedstock ratio of materials to water of 1:10. The empirical formula of bio-oil with a HHV of 26.6 MJ·kg⁻¹ was established as CH_{1.38}O_{0.43}N_{0.07}, and the bio-oil included species such as hexadecanoic acid, palmitamide, and ⁴⁵ fatty acid methyl esters.¹⁴⁸

The HTL of low-lipid microalga *C. pyrenoidosa* resulted in a peak bio-oil yield of 65.4 wt.% when carried out at 553 K and a reaction time of 120 min. The resulting HHV was 35.4 MJ·kg⁻¹, which increased to 38.5 MJ·kg⁻¹ at 573 K and 30 min reaction ⁵⁰ time, suggesting lower O and N contents in the bio-oil.⁴²

In summary, studies of HTL of microalgae shows that a

rather high bio-oil yield (20-66 wt.%) can be obtained, which is a highly viscous bio-oil with are relatively high N content of 1-5 wt.% and a HHV of 20-40 $MJ \cdot kg^{-1}$. The optimum operating conditions for obtaining the maximum bio-oil yield is in the 573-

- 5 623 K temperature range with reaction times of 15-120 min. However, the operating conditions are highly specific to strain and system. To obtain a bio-oil with lower nitrogen content, lower reaction temperature and shorter reaction time should be used or the protein fraction should be removed prior to HTL.
- ¹⁰ Most of the aforementioned studies have used small bench-scale reactors with capacity less than 10 g of microalgae as feedstock. Jena *et al.* conducted a relatively large scale experiment with *S. platensis* in a 1.8-L batch reactor.¹⁵¹ A reaction temperature of 623 K, holdup time of 60 min, and solids content of 20 wt.%
- ¹⁵ were identified as the optimal conditions for achieving the bio-oil yield of ~40 wt.%. Meanwhile, 98.3 wt.% of carbon was converted, and the obtained bio-oil had a HHV of $39.9 \text{ MJ} \cdot \text{kg}^{-1}$.

HTL is typically performed with slow heating and/or long reaction time (tens of minutes or longer). However, some recent ²⁰ results have suggested that shorter reaction times may be sufficient. A decrease in the reaction time would greatly reduce the reactor volume required for continuous bio-oil production, subsequently reducing the capital costs of such a process. Savage *et al.* investigated the fast HTL of the green marine alga

- ²⁵ Nannochloropsis sp. at batch reaction times of 1, 3, and 5 min within the temperature range of 573-873 K. Conventional liquefaction was carried out for 60 min at the same temperature range as a comparison.¹⁴⁹ The bio-oil yield of 66 wt.% was obtained at a reaction time of 1 min and temperature of 873 K.
- ³⁰ This yield exceeded any previous report on the HTL of the same algal species. The bio-oil produced by fast HTL has carbon contents and HHV similar to those produced by the traditional isothermal liquefaction process, which involves treatment for tens of minutes. Moreover, the authors affirmed that the reaction ³⁵ ordinate is a useful parameter for interpreting the results from algae liquefaction performed at different temperatures and

reaction times.¹⁴⁹ Almost all the results to date have been for batch systems,

and most studies have used organic solvents to recover the bio-oil 40 fraction, which is perhaps not necessary in a continuous process.

The introduction of organic solvents can increase the bio-oil recovery and will affect the water phase composition. Therefore, batch experiments only give partial insights into a continuous process; nevertheless, studies are useful for obtaining the ⁴⁵ optimum operating conditions and reaction pathways.

The large volume requirements of biomass conversion would rule out the use of a batch reactor. Thus, continuous operation will be required to make the bio-oil production more economically feasible. Moreover, the heat recovery from a

- ⁵⁰ continuous process increases the overall energy efficiency.¹⁶⁴ Jazrawi *et al.* developed a continuous-flow, pilot-scale HTL reactor system for the HTL of microalgae (*Chlorella* and *Spirulina*) as a follow-up to earlier research.¹⁶⁴ It was anticipated that the maximal bio-oil yield may be obtained at shorter
- 55 residence times under continuous-flow HTL processing. The investigators demonstrated the successful operation of a continuous-flow, pilot-scale HTL reactor system and provided insight into the processing of microalgae under subcritical

conditions. The bio-oil yields reached a maximum of 41.7 wt.%
⁶⁰ for *Chlorella* processed with a 10 wt.% solid concentration at 623 K, 3 min residence time, and 20 MPa. Continuous-flow HTL process provides a basis as well as technical parameters for its further industrialization. With increasing temperature, the oxygen content of the bio-oil decreased and the nitrogen content
⁶⁵ increased due to conversion of the algal protein fraction.

Recently, Elliott *et al.* reported a continuous-flow reactor system at relatively low temperature (623 K) and moderate pressure (20 MPa) to produce bio-oil from algae via HTL process.¹⁴⁰ A high bio-oil yield was obtained from the continuous 70 HTL of whole algae. An analysis of the bio-oil composition revealed lipid-derived alkane products and heterocyclics derived from other biomass components.

4.2.2 Catalytic HTL of microalgae

HTL technology can effectively convert wet microalgae ⁷⁵ into bio-oil. However, the obtained bio-oil has much higher concentrations of O and N compared to conventional petrochemical-based transportation fuels. In addition, not more than half the feedstock is converted to bio-oil, and the availability of feedstock is not reliable. This has motivated research of ⁸⁰ catalytic HTL intended to improve the bio-oil quality and yield. To date, the majority of studies that have carried out catalytic HTL have used homogenous catalysts, only a few articles have used heterogeneous catalysts (Table 4). A major challenge in catalytic HTL regards the catalyst durability in terms of leaching, ⁸⁵ deactivation, and regeneration under the aggressive conditions characteristic of HTL. Here we highlight some of the recent developments in catalytic HTL of algae.

(1) Homogenous catalysts

Homogenous catalysts have received more attention for ⁹⁰ liquefaction of algae than heterogeneous catalysts.²⁹ The addition of alkali salts has a positive effect on HTL. To date, the moststudied homogeneous catalyst for the HTL of microalgae has been Na₂CO₃, which has been shown to improve gasification rates, accelerate the water gas shift reaction, and increase overall ⁹⁵ bio-oil yields.^{27-29,31-33,57,60,153,155,163,165-169} In addition, the base catalysts raise the pH, inhibiting the dehydration of biomass monomers. Deoxygenation through dehydration (generally catalyzed by acid) instead of decarboxylization (DCO) give unsaturated compounds which easily polymerize to undesired ¹⁰⁰ char and tar.³⁶ Thus, alkali suppresses char and tar formation. Notwithstanding their advantages, homogeneous catalysts are more difficult to separate and recover after reaction.²⁹

Minowa *et al.* carried out the earliest work on homogenously catalyzed liquefaction of microalgae using 5% Na₂CO₃ solution for the strains on *B. braunii* and *D. tertiolecta*.^{29,163} The addition of Na₂CO₃ increased the bio-oil yield and energy yield. The HTL of *M. viridis* with 5% Na₂CO₃ as the catalyst resulted in the bio-oil yield increasing from 28.0 to 33.0 wt.% while the energy yield increasing from 29.4 to 39.5 ¹¹⁰ wt.%.³³ In addition, the decrease in the oxygen content of the biooil was from 24.2 to 19.7 wt.%.³³ The effect of the catalyst was more pronounced at lower temperature and shorter residence time. Similar results were obtained during the HTL of *D. tertiolecta*, *B. braunii* using 5% Na₂CO₃ as a catalyst; in fact, the ¹¹⁵ obtained bio-oil was nearly equivalent in quality to that of petroleum oil.^{32,163,169}

According to the influence of the content of microalgae on

the yields and product distribution, the reported bio-oil yield obtained from HTL microalgae exceeds the lipid content of the algae. The bio-oil yield followed the trend lipids > proteins > carbohydrates.²⁷ Both proteins and lipids were efficiently ⁵ converted to bio-oil without catalysts, whereas the carbohydrates conversion was enhanced with Na₂CO₃. The carbohydrate and protein fractions of microalgae in water were converted into biooil with efficiencies of nearly 10.0 wt.% and 20.0 wt.%, respectively.²⁷ The effectiveness of Na₂CO₃ reportedly depends to

¹⁰ a large extent on the reaction temperature. Dote²⁹, Minowa²⁸, and Inoue³⁰ found that increasing the liquefaction temperature from 573 to 613 K with Na₂CO₃ as a catalyst decreased the bio-oil yield. Yang³³ and Ross³¹ reported the opposite effect. These apparently inconsistent effects may be due to the different ¹⁵ biochemical compositions of the microalgae, which, as

mentioned above, respond differently to the presence of Na₂CO₃. The lipid content reported by Dote²⁹ and Minowa²⁸ was higher than that for the algae used by Ross³¹, supporting this hypothesis. In some cases the addition of a catalyst may not increase

20 the bio-oil yield but will alter the product distribution. For example, a common catalysts FeS had no effect on the bio-oil yield but significantly altered the proportion of the dominant compound, ethyl hexadecanoate.

The effect of catalyst type on the HTL of algae is another ²⁵ critical issue. For example, the HTL of *C. vulgaris* and *Spirulina* using alkalis (KOH and Na₂CO₃) and organic acids (CH₃COOH and HCOOH) as the catalysts were reported by Biller *et al.*³¹ The catalysts enhanced the bio-oil yield in the order of Na₂CO₃> CH₃COOH > KOH > HCOOH. The use of organic acids can

³⁰ improve the flow properties and lower the boiling point of the bio-oil.³³ However, the underlying mechanism for the apparent catalytic promotion by Na₂CO₃ or formic acid during the liquefaction of biological molecule like proteins remains unclear. The common catalyst Na₂CO₃ can have positive effects on the

³⁵ liquefaction of carbohydrates as well. For this reason, catalyzed HTL of high-carbohydrate microalgae resulted in higher yields with the addition of a carbonate catalyst as compared to the noncatalytic process. On the other hand, the use of alkali with high-lipid feedstocks can induce saponification reactions, which

⁴⁰ leads to soap formation and reduced the bio-oil yield. However, long-chain alkanes can be obtained from lipids with the use of Na₂CO₃. Model protein components investigated were preferably processed in water alone and exhibited the highest yields and energy content at the used conditions. These results suggest that

⁴⁵ high carbohydrate-containing algae should be processed in alkali, whereas high-protein and high-lipid algae are best processed in water alone or in formic acid because these conditions can reduce the boiling point and increase the flow properties.

Another group of catalysts, including alkaline earth metal ⁵⁰ [Ca₃(PO₄)₂] and transition metal oxide (NiO) were studied for the effect on the bio-oil yield from HTL of the microalga *S. platensis* for comparisons with alkali metal (Na₂CO₃).⁶⁰ Na₂CO₃ increased the bio-oil yield to 51.6 wt.%, which was 29.2 wt.% higher than that under noncatalytic conditions. In addition, the presence of

⁵⁵ NiO and Ca₃(PO₄)₂ increased the yields of gaseous products, while catalytic HTL using Na₂CO₃ produced lower gaseous yields than noncatalytic conditions. The use of the Ca- and Nibased catalysts increased the gaseous yields, and decreased the bio-oil formation.

In addition to the alkali Na₂CO₃ acidic species such as 60 H₂SO₄ have been used. Zou et al. liquefied the microalga D. tertiolecta at 393-473 K using 0-3.0 wt.% H₂SO₄ as the catalyst in ethyleneglycol (EG).¹⁵⁵ A statistical analysis of their data showed that a maximum liquefaction yield of 45.0 wt.% could be 65 obtained at the optimized conditions of 2.4 wt.% H₂SO₄ at 443 K for 33 min. The direct liquefaction of Sargassum polycystum C. Agardh in EG with H₂SO₄ as the catalyst along with microwaveassisted liquefaction resulted in bio-oil yield of 87.7 wt.%. In fact, the bio-oil was mainly composed of fatty acid methyl esters ⁷⁰ and alkanes with chain lengths from C_{17} to C_{20} .¹⁵⁴ When the feedstock was changed into Ulva prolifera, the maximum liquefaction yield of U. prolifera was 84.8 wt.% with a HHV of 15.1 MJ·kg⁻¹, which was obtained under a microwave power of 600 W using 6.0% H₂SO₄ as the catalyst via microwave-assisted 75 direct liquefaction.¹⁷⁰ The bio-oil was composed of benzene carboxylic acid, diethyl phthalate, long-chain fatty acids (C13 to C_{18}), fatty acid methyl esters, and water.

The use of homogeneous catalysts does not always have a positive effect on the bio-oil yields and properties, especially if ⁸⁰ the additional cost is considered. Furthermore, the recovery of the homogenous catalysts is a problem.

(2) Heterogeneous catalysts

Heterogeneous catalysts provide a more attractive option than homogeneous catalysts in HTL. Their practical advantage is ⁸⁵ their separation is accomplished by simple filtration. Moreover, solid catalysts are commonly used in low-temperature water gasification of biomass, and gasification is crucial during HTL because oxygen is removed during this process.¹⁶¹

A large number of studies have been carried out using ⁹⁰ catalysts to improve bio-oil yield during HTL of microalgae. The choice of catalyst depends on the specific composition of the algal strain. In addition, heterogeneous catalysts tend to undergo coking during HTL process. Moreover, regeneration of the deactivated heterogeneous catalysts was difficult because the ⁹⁵ catalysts alone cannot be separated from the solid residue. These shortcomings greatly limit their practical application in the HTL of algae, and further works should be performed to overcome these disadvantages.

The most extensive reports on the influence of ¹⁰⁰ heterogeneous catalysis on HTL were published by Duan and Savage, who produced crude bio-oils from the microalga *Nannochloropsis sp.* via HTL in the presence of six heterogeneous catalysts [Pd/C, Pt/C, Ru/C, Ni/SiO₂-Al₂O₃, CoMo/γ-Al₂O₃ (sulfided), and zeolite].³⁵ The bio-oil yield in the ¹⁰⁵ absence of catalysts was 35.0 wt.%, but increased to 57.0 wt.% when the Pd/C catalyst was used without hydrogen. Ni/SiO₂-Al₂O₃ was the most active catalyst for desulfurization. The bio-oil produced with in the presence of Pd, Pt, Ru and Co-Mo catalysts exhibited a lower viscosity and lighter color than the uncatalyzed ¹¹⁰ or zeolite-catalyzed samples. Meanwhile the presence of Ni, Pt, and Co-Mo decreased the O/C ratio. This suggests that catalytic deoxygenation was promoted.

In another study, NiO was used to catalytic HTL of both single (*Spirulina*) and mixed algal (from open ponds with ¹¹⁵ wastewater) cultures.¹⁵⁸ Unexpectedly, the added NiO decreased bio-oil yields. The maximum bio-oil yield is up to 40.0 wt.% in

the presence of alumina-supported transition metal catalysts in the temperature range of 573 to 623 K.²⁷ The liquefaction procedure carried out at 623 K gave a bio-oil with a HHV of 39.0 MJ·kg⁻¹. Biller *et al.* investigated three catalysts: an aluminas supported Co/Mo catalyst, an alumina-supported Ni catalyst and

- an alumina supported Pt catalyst, ²¹ The results indicated that the bio-oil yield from the HTL of *C. vulgaris* and *N. occulta* increased slightly with the use of heterogeneous catalysts; however, the increase of HHV was up to 10%. The HTL results of
- ¹⁰ a low-lipid microalga *C. pyrenoidosa* using heterogeneous catalysts showed that Raney-Ni and HZSM-5 catalysts had no significant effect on the HTL process. H_2 as the processing gas slightly improved the bio-oil yield and quality, whereas catalysts have no significant effect.⁸⁴
- ¹⁵ Matsui *et al.* studied the liquefaction of *Spirulina* with various concentrations of $Fe(CO)_5$ -S catalyst.¹⁵⁷ Reactions in 1-methylnaphthalene with a small amount of water in CO and $Fe(CO)_5$ -S gave conversions greater than 96.0 wt.% and total amount of bio-oil, gas and water yields was up to 83.0 wt.%.
- In summary, catalytic HTL conversion of algae can produce hydrocarbons for liquid fuels and hydrogen/methane-rich product gases. Thus, this field has tremendous potential and a bright outlook. Most recent studies on producing liquid fuels from the HTL of algae have focused on homogeneous catalysis by metal
- ²⁵ salts or alkali. More work is needed to identify better heterogeneous catalysts for these applications. In particular, the development of nonprecious metal-based catalysts is of particular interest. Finally, active catalytic materials that resist deactivation during HTL are needed.

30 4.2.3 HTL of microalgae in co-solvents

Water has been the most popular medium for the liquefaction of microalgae. However, HTL with water as the sole medium has drawbacks. First, the liquefaction is relatively harsh, involving rather high temperatures (523-623 K) and high

³⁵ pressures (10-20 MPa). Second, the bio-oil product is of inferior quality because it contains a relatively high oxygen and nitrogen content, which decreases the HHV and storage stability. Third, the conditions are unfavorable for maximizing the bio-oil yield. A recent study showed that only 40% carbon and 35% hydrogen

⁴⁰ in the feedstock are converted to bio-oil while a large fraction of organic species remained in the aqueous phase, thus resulting in a relatively low yield of bio-oil.³³

To address these drawbacks, organic solvents have been studied as alternative media during microalgal liquefaction. ⁴⁵ Organic solvents used in microalgal liquefaction can dissolve or stabilize the weak polar or even nonpolar intermediates because of their lower dielectric constant compared to water.¹⁷¹ Thus, such solvents can produce higher bio-oil yield. Another advantage of using organic solvents is that more moderate operating conditions ⁵⁰ can be used.

The HTL of the low-lipid microalga *C. pyrenoidosa* was processed under sub/supercritical ethanol⁸⁴ The highest bio-oil yield of algae was 71.3 wt.% at 513 K, whereas the highest HHV of the bio-oil was 36.2 MJ·kg⁻¹ at 573 K. Supercritical ethanol ⁵⁵ condition (>513 K) is essential for the conversion of *C. pyrenoidosa*, and higher temperature facilitates deoxygenation.⁸⁴

In addition, publication studies have shown that the solvent polarity can have a significant impact on the liquefaction features

of microalgae.¹⁷² When ethanol as the reaction medium for 60 liquefaction of C. pvrenoidosa, the HHV of the bio-oils produced under different reaction conditions ranged from 27.7 to 36.5 MJ·kg⁻¹. The bio-oil yield increased from 9.8 to 64.6 wt.% while the solid residue decreased from 60.1 to 11.9 wt.% as the temperature was increased from 443 to 623 K. The solvent type is 65 significantly affected the bio-oil product distribution. The dominant components of the bio-oils were fatty acid methyl and ethyl esters when methanol and ethanol were used as solvents, respectively. In contrast, the primary product was hexadecanenitrile when 1, 4-dioxane was used as solvent.⁵⁹ In 70 addition to solvent polarity, operating variables, such as temperature, reaction time, solvent/microalgae ratio, and catalyst type and dosage, also influenced the conversion and yield of the bio-oil. As reported by Matsui et al., reactions in tetralin and hydrogen significantly increased the bio-oil vield as the 75 temperature was increased from 573 to 698 K.¹⁵⁷

Additionally, previous studies had proved that the use of a co-solvent with water was advantageous to microalgae HTL on the bio-oil yield and product distribution.¹⁷³⁻¹⁷⁶ Brennecke *et al.* found that the solubility of supercritical fluids could be greatly ⁸⁰ improved by adding a small amount of a second solvent, which is commonly called co-solvent.¹⁷⁴ Ethanol, 2-propanol, and methanol are often used as co-solvents for microalgal conversion in HTL.^{175,176} Significantly higher bio-oil yield can be achieved in co-solvent-water mixture than in water alone as the reaction ⁸⁵ medium. Investigation of hydrophobic hydration in methanol-water mixtures under supercritical conditions is of a great practical importance for chemical engineering.^{177,178}

Chen *et al.* reported the production of bio-oil by direct liquefaction of *D. tertiolecta* with sub/supercritical ethanol-water ⁹⁰ as the reaction medium at high temperature and pressure.¹⁷⁵ The bio-oil and solid residue (SR) yields as well as the conversion, are shown as a function of ethanol content in Fig. 12. The results indicate that using either ethanol or water as the medium is less effective for the conversion or bio-oil yield compared to the ⁹⁵ ethanol-water mixture. Ethanol and water showed synergistic effects on the liquefaction of *D. tertiolecta*. Sub/supercritical water can provide ionic, polar nonionic and free radical for the production of bio-oil from microalgae.¹⁷⁹

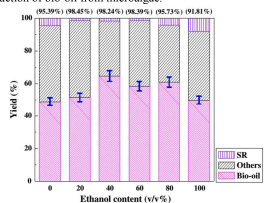


Fig. 12 Effect of ethanol content on the bio-oil, others and SR yields for the microalgae liquefaction in ethanol-water cosolvent¹⁷⁵ Copyright 2012 Elsevier

Other studies have reported the function of ethanol as hydrogen-donors in algae liquefaction for bio-oil production.¹⁷⁶ In spite of the interesting synergistic effects afforded by the solvent

mixture, the liquefaction of algae with sub/supercritical ethanolwater as the medium is a complex process, and many reactions could occur. The hydrogen-donor function of ethanol cannot be regarded as a simple dehydrogenation of ethanol. Direct evidence 5 of the hydrogen-donor effect could be obtained by the hydrogen

isotopic tracer method. Clearly, further research is needed. 4.2.4 Co-liquefaction of microalgae with other substances

(1) Co-liquefaction of microalgae and coal

Coal is an attractive long-term energy source because of its comparatively large reserves. However, its larger sulfur, and metals content calls for the development of clean conversion technologies. Along these lines, direct coal liquefaction is an interesting technology. However, coal liquefaction requires harsh reaction conditions and the requisite hydrogen increases the cost and deters wide-spread adoption. To this end, the co-liquefaction

of coal and biomass has gained increasing attention because it takes full advantage of the hydrogen of biomass, which could reduce hydrogen consumption, and result in milder operation 20 compared to direct coal liquefaction.

Earlier studies have investigated the co-liquefaction of microalgae (*Chlorella*, *Spirulina*, and *Littorale*) with coal (*Australian Yallourn* brown coal and *Illinois* No. 6 coal) under pressurized H₂ in 1-methylnaphthalene at 623-673 K for 60 min ²⁵ with various catalysts.⁵¹ Co-liquefaction of *Chlorella* with

²⁵ with various catalysts. Co-inqueraction of *Chioretta* with *Yallourn* coal resulted in 99.8 wt.% conversion with a 65.5 wt.% of hexane-soluble fraction obtained at 673 K using a $Fe_{1-x}S$ catalyst. When *Littorale* and *Spirulina* were used, similar results were obtained with an iron-based catalyst. Conversely, the oil

³⁰ yield in the co-liquefaction with *Illinois* No. 6 coal was close to the additivity of the respective reaction with Fe(CO)₅-S, even at S/Fe=2. Ru₃(CO)₁₂was also effective for the co-liquefaction of microalgae with coal.⁵¹

Yang *et al.* reported the co-liquefaction of *D. tertiolecta* and ³⁵ coal to produce liquid fuel with sub/supercritical water-ethanol as the reaction solvent.¹⁸⁰ The optimal conversion and oil yield were 70.6 and 40.3 wt.%, respectively. The results showed that an obvious synergetic effect existed between *D. tertiolecta* and coal, which not only improved the conversion and oil yield but also

⁴⁰ enhanced the oil quality. The synergetic effect values of conversion and bio-oil yield were 15.7 and 12.5 wt.%, respectively, under the optimal reaction conditions.

(2) Co-liquefaction of microalgae and plastic

- The rapid growth of plastics use worldwide has led to a 45 concomitant increase in the amounts of plastics waste which is bulky and resistant to degradation. Thus, the conversion of waste plastic to liquid fuels is an intriguing approach especially given its typically high HHV (approximately 40 MJ·kg⁻¹) as a result of its high hydrogen and carbon content. The co-processing of waste
- ⁵⁰ polymer with biomass has received considerable attention.¹⁸¹ Plastics could provide hydrogen during co-processing with biomass, increase oil production, and improve oil quality because of the high hydrogen content in plastics (approximately 14 wt.% for PP and PE). Furthermore, the degradation of polymer could be ⁵⁵ improved via mixing with biomass.

A study on the co-liquefaction of microalgae (*Spirulina*) and synthetic polymer (high-density polyethylene, HDPE) in sub/supercritical ethanol showed that the decomposition of

Spirulina and HDPE were mutually improved.¹⁸¹ The addition of
Spirulina reduced the requisite degradation conditions for HDPE liquefaction and resulted in a high conversion of the HDPE. Synergetic effects were reported for the co-liquefaction of *Spirulina* and HDPE. For example, with a Spirulina/HDPE feed ratio of 4/6, the oil yield obtained at 613 K increased by 44.8
wt.%. The oil from *Spirulina*/HDPE co-liquefaction had higher carbon and hydrogen content but lower oxygen content, resulting in a HHV of 48.4 MJ·kg⁻¹, a level comparable to fossil fuel. Moreover, the chemical compositions of the oil from co-liquefaction of *Spirulina*/HDPE blends were similar to that from ⁷⁰ sole HDPE liquefaction, in which aliphatic hydrocarbons dominated.¹⁸¹

In summary, microalgae can have obvious different role when they co-liquefying with different kinds of substances. Microalgae could act as the hydrogen donor in the co-processing of algae with coal because of their higher hydrogen contents. In view of the co-processing with plastics, algae act as the hydrogen receiver because they have less hydrogen than plastics. The obvious synergetic effect exists during co-processing; as a result, both the oil yield and quality have been improved.

80 4.2.5 Mechanistic pathways of microalgal HTL

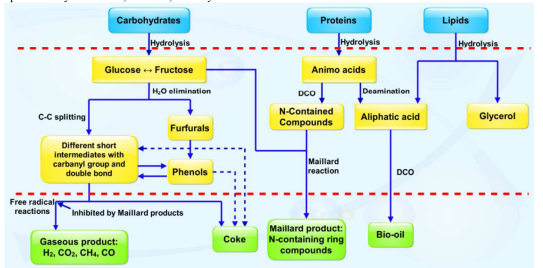
In HTL, water simultaneously acts as reactant and catalyst, making this process different from pyrolysis. Under conditions close to the critical point, water has many interesting properties, such as low viscosity, high solubility of organic substances, high 85 ionic product, etc. During HTL, a series of reactions occur, depolymerization, including hydrolysis, and repolymerization/self-condensation.¹⁸¹ Microalgae, consisting of proteins, carbohydrates, and lipids, produce distinct compounds during HTL. Generally, lipids are transformed into fatty acids; 90 proteins into nitrogen heterocycles, pyrroles, and indoles; and carbohydrates into cyclic ketones and phenols.²⁷ The products of HTL consist of bio-oil, water-soluble fractions (containing polar organic compounds), gaseous and solid residue fractions. In addition, a substantial part of the oxygen in the microalgae can be 95 removed by dehydration or decarboxylation (DCO). In spite this general understanding of overall effects, the mechanisms for the HTL reactions of microalgae to bio-oil are not well understood. It goes without saying that understanding the microalgal HTL mechanisms and kinetics is essential for the design and 100 optimization of the overall HTL process. Below we highlight the current level of understanding.

The carbohydrates in algae include polysaccharides, celluloses, hemicelluloses and starches. During HTL, carbohydrates are rapidly hydrolyzed to monosaccharides with ¹⁰⁵ glucose as one of the main products (Fig. 13).^{161,182} The glucose is readily converted to fructose, an isomer of glucose. The fructose subsequently undergoes degradation with fragmentation products (e.g., glycolaldehydes and glyceraldehydes). Some short intermediates can form volatile product (e.g., H₂, CH₄, CO, etc.) ¹¹⁰ and coke via further reaction.

Lipids mainly consist of fatty acid triglycerides (TAGs), which are nonpolar compounds with aliphatic characteristics. In HTL, TAGs are hydrolyzed to fatty acids and glycerol. The glycerol is subsequently converted to water-soluble soluble 115 compounds. Free fatty acids are relatively stable but partially degrade to produce long-chain hydrocarbons for transportation fuels via DCO (Fig. 13).

Most proteins are composed of linear polymers of amino acids, and they both have structural and metabolic functions. The peptide C-N bond links the amino acids together between the 5 carboxyl and amine groups; this bond will be hydrolyzed under HTL conditions resulting in the production of amino acids (Fig. 13). The amino acids rapidly undergo DCO and deamination, and consequently produce hydrocarbons, amines, aldehydes and

acids. Some of these products are the same as those from the 10 hydrolysis of carbohydrate. The interaction between the hydrolysates from carbohydrates and proteins can react with each other to generate N-containing ring compounds. This process is recognized as Maillard reaction, which is confirmed by many other published papers.^{37,84,161,182-186}



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Fig. 13 Simplified reaction pathways for HTL of carbohydrates, proteins, and lipids altogether ¹⁶¹Copyright 2011 Elsevier, ¹⁸²Copyright 2010 Elsevier With the aforementioned main chemical pathways identified, the HTL of algae is a complex process that is strongly 20 affected by the feedstock type and the HTL experimental conditions. Knowledge about the composition of HTL bio-oils is important to understand the HTL mechanism. Torri et al. provided the HTL mechanisms based on the chemical compounds in the bio-oils obtained from the HTL of Desmodesmus sp.¹⁸⁷ The 25 possible mechanisms are described as follows (Fig. 14).¹⁸⁷

(1) The HTL-derived bio-oil is a mixture of a large number of compounds and macromolecular constituents, ranging from peptides to long-chain hydrocarbons. HTL at relatively low temperature (473-523 K) allows the extraction of the solvent 30 soluble part of the cell constituents. Therefore, lipids, some shortchain algaenans, and some hydrophobic protein fragments end up in the organic solvent phase. However, most proteins and carbohydrates are not converted to water-insoluble products. Below 523 K, HTL is accompanied by a certain degree of thermal

35 degradation, and the extraction of lipids and algaenan is improved to a certain degree. Therefore, the reactivity between proteins and carbohydrates is crucial during HTL. In general, proteins and carbohydrates can interact between themselves and/or with lipids through various ways.

(2) At 573-648 K, proteins and celluloses start to break 40 down, giving diketopiperazines (DKP), amino-acid derivatives, and carbohydrates derivatives (e.g., furans), the products from the cross reaction of those species, and asphaltene-like components. Protein degradation increases the bio-oil yields, but the

45 hydrophobic portion of the protein-carbohydrate derived components increase the nitrogen content of the bio-oil. The main chemical route for the protein conversion is probably through depolymerization. Peptide depolymerization can be described as a progressive cyclization with the formation of gradually smaller

50 cyclic oligo-peptides, with the final product being DKP. Even at 473 K, carbohydrates and proteins can be fragmented into smaller products (e.g., amines and aldehydes), which may be able to form melanoid in-like materials and asphaltene-like materials.

(3) As the reaction proceeds, a strong increase is observed 55 in "pyrolysis-like" products produced from thermal fragmentation of proteins and carbohydrates (e.g., amino acids side chains or 2methyl-cyclopentenone) and of smaller products from Maillard reactions (e.g., pyrroles), at the expense of peptides and DKP, which are probably converted into amino acids and/or to other 60 by-products.¹⁸⁷

As mentioned earlier, introducing organic compounds as the processing solvents in HTL can improve the quality of bio-oil. The HTL mechanism of microalgae in ethanol is different from that in water. Based on GC-MS results and published reports, 65 Zhang et al. provided a potential HTL mechanism of algae (Fig. 15):⁸⁴ (1) Carbohydrates, proteins, and lipids first break down to their corresponding monomers, such as glucose, xylose, phenols, amino acids, and fatty acids under HTL conditions. These monomers further decompose, and then to form various types of 70 intermediates. (2) The monomers and their intermediates undergo a series of reactions. As illustrated in Fig. 15, the amino acids undergo DCO and deamination reactions to form the corresponding amines and keto acids, respectively. The keto acids undergo DCO to form ketones, which are abundant in the liquid 75 products. The fatty acids undergo DCO to form aliphatic hydrocarbons or react with ethanol through esterification to form fatty acid esters. The fatty acids also react with amines or ammonia through acylation to form amides. The monosaccharides react with amino acids through Maillard 80 chemistry to form melanoidin (nitrogenous polymers) and solid products. The monosaccharides further decompose to form small

acids and furfural derivatives. The furfural derivatives and phenols undergo repolymerization to form large molecular components and solid products.⁸⁴ The competitive reaction of the

carboxyl group and ethanol to form esters, which are more stable ⁵ under HTL conditions, suppress DCO and which results in a decrease in the volatile product yield.

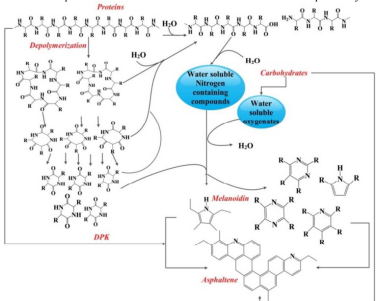
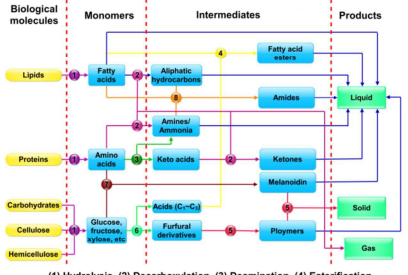


Fig. 14 Possible mechanisms for HTL oil formation from protein/carbohydrate macromolecules¹⁸⁷ Copyright 2012 American Chemical Society



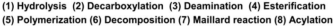


Fig. 15 Potential reaction mechanism of HTL for algae⁸⁴ Copyright 2013 Elsevier

Water plays a crucial role during the HTL of algae. Of particular importance is hydrolysis which converts proteins, carbohydrates, and lipids into a variety of components. To this end, the use of a cosolvent with water has attracted considerable 15 attention because of the possible redirection reaction pathways.

- ²⁰ carbohydrates, and lipids in *D. tertiolecta* is acid-catalyzed. Under acidic conditions, proteins first form a long peptide chain, which is then hydrolyzed to form amino acids. The amino acids undergo cracking, condensation, DCO, deamination, etc. to form liquefied product. Carbohydrates undergo dehydration to form
- ²⁵ monosaccharides, a part of which may then react with ethanol to form the ether that exists in the solid residues. Most monosaccharides may further react to generate carboxylic acid or other organic compounds that undergo the acid-catalyzed process. Lipids undergo dehydration to form carboxylic acid and glycerol.
- ³⁰ A competitive reaction with carboxylic acid occurs between ammonia and ethanol to form amides and esters during HTL. With sufficient ethanol present, amides react with ethanol. The presence of ethanol significantly influences the composition of liquefied products. The carboxylic acid can then react with ³⁵ ethanol via alcoholysis to form carboxylic acid esters and undergo ammonolysis with ammonia to form amides. The amides can also undergo alcoholysis to generate carboxylic acid esters when the concentration of ethanol is high. Ammonia, which is

produced during the acid-catalyzed decomposition of proteins,

may be used as an ammonolysis reagent in direct liquefaction.¹⁷⁵ Ester, Hydrocarbon, etc.

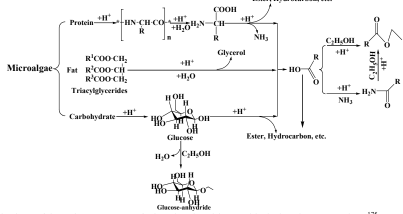


Fig. 16 HTL mechanisms of the main components of microalgae in sub/supercritical ethanol-water cosolvent¹⁷⁵ Copyright 2012 Elsevier

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- In summary, macromolecules in the microalgae first hydrolyze into small fragments (fatty acids, amino acids, and glucose), which are then converted into even lower molecular weight compounds. Particularly, amino acids undergo DCO and deamination to produce hydrocarbons, amines, aldehydes, and
- 10 acids. The generated compounds (intermediates) are unstable and rearrange into larger compounds via condensation, cyclization, and polymerization. Although many researchers have studied the mechanism of HTL, understanding the detailed interactions during the HTL of microalgae and their kinetics warrant further

15 research.

4.2.6 Prospectives for HTL of microalgae

HTL is a desired conversion process that is specifically suitable for producing biofuel from wet feedstock because the energy intensive drying process is not necessary. Compared to

- 20 biofuel produced from other thermochemical conversion process, such as pyrolysis, HTL biofuel has a higher energy density because it is more effective in terms of oxygen removal from the biomass feedstock via decarboxylation. Undoubtedly, HTL of microalgae are still in their early stages, in spite there is a large
- 25 number of superiority in the biofuel production via microalgae HTL, the goal to provide a sufficient and cost-competitive method for the biofuel production without government subsidy has not been achieved yet.

Microalgal biomass, which has potential to serve as a 30 renewable source of energy, is relatively new and unexplored. HTL will most likely find its place in an energy system where algae are used because of its safe and cheap reactant (water). Any commercial application of algae HTL would require the use of a continuous process where heat recovery can be incorporated in

- 35 order to be sufficiently energy efficient. To achieve high yields of high quality bio-oil, the use of catalysts and potentially co-solvent is needed. Further research is needed to identify these catalysts and to understand the reaction pathways and to quantify the HTL kinetics. The following are recommended key factors that will ⁴⁰ have to be addressed pertaining to HTL microalgae:
 - Catalyst. Catalytic HTL is one of the most ideal methods of microalgal liquefaction, and a suitable catalyst can increase biofuel yield and improve quality of bio-oil as well as. However, homogenous catalysts present a serious problem on
- the recovery of the catalysts, and the hydrothermal stability of 45 heterogeneous catalysts must be taken into account when

water as the reaction medium during HTL microalgae process. Further studies on more catalysts are necessary to identify supports and active materials that can better resist deactivation in HTL.

- Mild reaction conditions. HTL with sole water as the medium has potential drawbacks including relatively harsh with rather high temperatures (523-623 K) and high pressures (10-20 MPa). To solve these problems, organic solvents or organic solvents-water mixture instead of water have been applied 55 during microalgal HTL. Replacing water with organic solvent or mixture as the reaction medium for HTL is theoretically feasible. The main advantage of using organic solvents or mixture is that more moderate operating conditions can be obtained. Organic solvents that have been used for 60 microalgae liquefaction include alcohol, dioxane, tetralin, 1methylnaphthalene, toluene and chloroform, etc.
- Development of pilot-scale plant. Process development work related to continuous operation and scale up has been reported. Process development, design, and optimization are 65 facilitated by the availability of mathematical models that faithfully describe the process chemistry. The development of economic pilot-scale plant of HTL microalgae to produce biooil is one trend of HTL development.
- 70 Reaction mechanisms and kinetics. Full knowledge of the mechanisms involved in the HTL process is critical. A systemic approach in determining the effects of various feedstock components and different reaction conditions on bio-oil yield and quality is needed because numerous algae species could be used as feedstock and different reaction 75 conditions could be carried out.

4.3 Comparison of pyrolysis and HTL of algae

As mentioned above, HTL is a low-temperature (523-623 K) and high-pressure (5-20 MPa) process particularly suited for 80 high-moisture feedstocks. In contrast, pyrolysis is accomplished at moderate to high temperatures (673-873 K) and atmospheric pressure but requires feedstock drying. Considerable attention has been given to HTL and pyrolysis for algal conversion. However, evaluating these two competing methods is complicated by the 85 myriad of different algae strains. For this reason, studies that have directly compared the two methods using the same feedstock can provide valuable information.

One such study was conducted by Jena and Das who

compared the liquefaction and pyrolysis of *S. platensis*.⁶¹ The conversion was performed with a slurry containing 20.0 wt.% of algal biomass and a reaction time of 60 min. Detailed comparison results are listed in Table 5. The energy consumption ratio (ECR) ⁵ of HTL was found to be 0.70, indicating that this process was a net energy producer. In contrast, the ECR value of pyrolysis

suggested that the pyrolysis consumed more energy than what could be produced from algal feedstock. Moreover, the bio-oil obtained from HTL had a higher energy density and superior fuel ¹⁰ properties such as thermal and storage stability, compared to that obtained from pyrolysis.¹⁸⁸

	Liquefaction	Slow pyroly	vsis
Reaction temperature/K	623	623	773
Conversion/wt.%	93.0	60.0-72.0	-
Bio-oil yield/wt.%	40.7	23.8	28.5
HHV/MJ·kg ⁻¹	34.2	29.3	33.6
Energy recovery from the original algae/%	67.9	33.9	46.7
ECR	0.7	2.11	1.56

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 Table 5 Comparison between HTL and slow pyrolysis of algae⁶¹ Copyright 2011 American Chemical Society

The compositions of bio-oil from HTL and pyrolysis have notable differences. Bio-oils generated from pyrolysis, especially 15 at 623 K, have higher percentages of nitrogenous compounds and aromatic heterocycles compared to those generated from HTL. Bio-oils generated from HTL are easier to upgrade than those from pyrolysis because of the higher abundance of straight-chain

- compounds than that of the former. FT-IR spectra of the bio-oil ²⁰ samples obtained from HTL and pyrolysis of microalgae are shown in Fig. 17.⁶¹ A distinct band at approximately 3300 cm⁻¹ for pyrolytic bio-oils corresponds to N-H functional groups and represents a higher abundance of nitrogenous compounds than
- that for HTL bio-oils. Lower peaks in bio-oil from HTL at 1670 $_{25}$ cm⁻¹ (C=O) represent less abundance of carboxylic acids, esters,

or aryl ketones than that in bio-oil from pyrolysis.

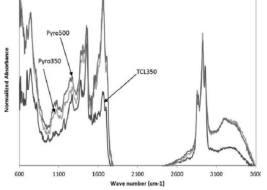


Fig. 17 Infrared spectra of bio-oil samples obtained from HTL and pyrolysis of microalgae⁶¹ Copyright 2011 American Chemical Society

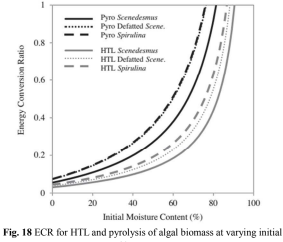
³⁰Bio-oil from HTL had higher amount of inorganic elements than that from pyrolysis. This result can be attributed to HTL being a high-pressure process that results in more intense reactions compared to pyrolysis. Greater amounts of inorganic elements could have leached from solids ending up in the ³⁵ liquids/bio-oil fraction. In comparison, the bio-oil vapor from

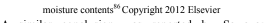
pyrolysis was collected in a set of condensers, leaving the solids inside the main reactor, as the reaction further proceeded.

In a second study, Vardon *et al.* converted *Scenedesmus* (raw and defatted) and *Spirulina* to bio-oil by HTL (573 K and ⁴⁰ 10-12 MPa) and slow pyrolysis (heated to 723 K at a rate of 50K·min⁻¹), and then compared the produced bio-oils.⁸⁶ Both the two conversion routes produced energy-dense of bio-oil (35-37 MJ·kg⁻¹) similar to shale oil (41 MJ·kg⁻¹). However, bio-oil yields (24-45 wt.%) and physicochemical characteristics were ⁴⁵ greatly influenced by conversion route and algal strains. Notable differences were observed in the mean bio-oil molecular weight (pyrolysis: 280-360 Da; HTL: 700-1330 Da) and the percentage of low-boiling compounds (bp<673 K) (pyrolysis: 62-66 wt.%; HTL: 45-54 wt.%). HTL and slow pyrolysis of algae produced ⁵⁰ bio-oils with similar HHV, heteroatom content, and functionality. Nevertheless, pyrolytic bio-oil displayed a significantly higher percentage of cyclic oxygenates (16-24 wt.%) compared to HTL bio-oil (8-12 wt.%) in the form of phenolic compounds.

The energy efficiency of the two conversion routes was also 55 discussed. Analysis of ECR also indicated that HTL is more favorable (ECR 0.4-0.6) than pyrolysis (ECR 0.9-1.2) for processing wet algal biomass (80% moisture content) because the latter requires water volatilization. However, pyrolysis is energetically favorable if the starting algal biomass has low 60 moisture content (Fig. 18).⁸⁶

Finally, HTL and pyrolysis have different dewater requirement and schedule. HTL is favorable for wet algal conversion because of its integration with wet microalgae slurry (10-20% solids), as opposed to pyrolysis, which requires dried ⁶⁵ microalgae (80% solids). The dewatering requirements to achieve the percent solids required for the HTL necessitates the use of bio-flocculation, dissolved air filtration and centrifugation for water removal. The pyrolysis pathway requires the remaining water to be removed by using thermal methods. Drying of ⁷⁰ microalgae requires substantial energy, accounting for nearly half of the overall net energy ratio (NER) for the pyrolysis pathway modeled at the industrial-scale.¹⁸⁹





A similar conclusion was reported by Sawayama.185

According to their calculations, the energy required for algae liquefaction is only $6.7 \text{ MJ} \cdot \text{kg}^{-1}$ of the bio-oil produced. Therefore, HTL is preferred over pyrolysis for the conversion of algae to bio-oil because of its energetic and economic advantages.

- Although HTL may be more attractive than pyrolysis with respect to energy efficiency, not all studies agree that HTL has a positive ECR. Biller and Ross employed HTL on four algal species (*C. vulgaris, Spirulina, Nannochloropsis oc.* and *P. cruentum*) and indicated that only the HTL of *C. vulgaris* without
- ¹⁰ catalyst produced an ECR of 0.8. The ECRs of all other species were all equal to or larger than 1.0, suggesting that more energy is needed for liquefaction than what can be gained from the biooil.²⁷ Therefore, algal conversion to bio-oil by HTL must be performed with caution.

5. Upgrading of crude bio-oil from algae

As we have discussed, bio-oil derived from algae via the thermochemical conversion processes of pyrolysis and HTL has a higher oxygen content, lower stability and lower HHV than ²⁰ petroleum. This is in spite of the substantial advances made in catalytic pyrolysis and HTL towards the enhancement of bio-oil yield and quality. The upshot is that the bio-oil requires further upgrading to put in on par with conventional petroleum feedstock

- for subsequent refining into transportation fuel. As discussed ²⁵ above, generally HTL bio-oil has lower oxygen and nitrogen content than bio-oil obtained by pyrolysis.³¹ The ultimate aim of upgrading is to improve the quality of bio-oil by decreasing the fraction of organic acids, aldehydes, and other reactive compounds because they increase corrosiveness and acidity.
- ³⁰ Moreover, compared to bio-oil derived from lignocellulosic biomass, microalgal bio-oil contains significant amounts of nitrogen rooted from protein in microalgae, which is undesirable in the final product.

Because of the high diversity of compounds in the bio-oil, ³⁵ the upgrading of bio-oil is a complex reaction network; representive reactions include cracking, decarbonylation, DCO, hydrocracking, hydrodeoxygenation (HDO), and hydrogenation. ¹⁹⁰⁻¹⁹⁴ These are discussed next.

Catalytic cracking with various zeolite catalysts is an ⁴⁰ attractive method for upgrading triglycerides to produce fuels.¹⁹⁵⁻¹⁹⁸ Cracking does not require H₂, which is a significant advantage over deoxygenation using hydrotreating catalysts. However, cracking has the disadvantage of coking of the catalyst, which therefore requires regeneration to maintain activity.

⁴⁵ DCO and decarboxylation yield hydrocarbon chains with one carbon atom less as compared to the reactant, that is to say, carbon chain length is reduced, which is generally undesirable for fuels. Decarbonylation yields olefins, and DCO produces paraffins. By contrast, the HDO route selectively cleaves C=O

⁵⁰ bonds, while the C-C bonds remain intact. Furthermore, the HDO reaction eliminates oxygen by producing H₂O instead of CO₂, rendering it more environmentally friendly than DCO, and attract more attentions.

It is worth mentioning, in order to improve fuel combustion ⁵⁵ performance it may be desirable to retain some oxygen in the fuel. These methods include esterification and selectivehydrogenation.¹⁹⁹ Wan *et al.* reported one-step hydrogenationesterification of aldehyde and acid to ester over bifunctional Pt catalysts.²⁰⁰ However, production of oxygenated biofuel only be ⁶⁰ reported in upgrading of bio-oil derived from lignocellulosic biomass, and there are no published data for algae-base bio-oil, it may be important topic worth studying in the future.

5.1 Upgrading of model compounds

Algal-derived bio-oil contains unsaturated and saturated 65 long-chain fatty acids (C_{16} - C_{18}), so it is of interest to understand the relevant upgrading chemistries for these species. Here we report on selected studies.

In recent work, the mechanisms for decarbonylation and DCO were investigated using palmitic or oleic acids as model $_{70}$ compounds on different catalysts.²⁰¹⁻²⁰⁵ The main products from palmitic acid were C₈-C₁₅ n-alkanes, with pentadecane the alkane with the highest yield. Hydrogenation of the double bond in oleic acid was faster than its DCO to form C₁₇ hydrocarbons.

Noble metal-based catalysts have high activity and have ⁷⁵ been extensively investigated. Na et al. investigated the deoxygenation of bio-oil obtained by pyrolysis of microalgae for the production of hydrocarbon fuel by metal-supported catalysts.²⁰⁶ The distribution of the products depends on the reaction conditions and catalyst used.²⁰⁷ To reduce the cost, ⁸⁰ tungsten-based catalysts were explored.²⁰⁸ Both tungsten oxide and tungsten carbide based catalysts allow an upgrading of the feed to higher-value products. The activated carbon is promising and inexpensive catalytic materials for converting fatty acids to alkanes.^{209,210}

Upgrading via hydrogenation is a promising route and 85 another research focus.¹⁹⁷ In a series of related works, Lercher and coworkers²¹¹⁻²¹³ reported that the extracted microalgae oil from algae cell is found to be triglycerides predominant. The upgrading process begins with the hydrogenolysis of the 90 triglycerides to produce fatty acids. The research showed that microalgal oil can be nearly quantitatively hydrodeoxygenated to alkanes by a cascade of reactions on bifunctional catalysts containing Ni and an acidic zeolite (HZSM-5 and HBeta).²¹¹ The reaction pathway proceeds through an initial hydrogenolysis of 95 triglyceride leading to fatty acid and propane. The subsequent hydrogenation of the carboxylic group of fatty acid leads to the corresponding aldehyde; for example, octadecanal (ratedetermining step), followed by either decarbonylation of octadecanal to n-heptadecane and carbon monoxide (minor route) 100 or hydrogenation of octadecanal to 1-octadecanol (major route). Subsequently, the produced 1-octadecanol undergoes sequential acid-catalyzed dehydration and metal-catalyzed hydrogenation leading to the final n-octadecane. The overall reaction pathway proposed for microalgae oil transformation is shown in Fig. ¹⁰⁵ 19A.²¹¹ In summary, Ni catalyzes efficiently the hydrogenolysis of the fatty acid ester, the decarbonylation of aldehyde intermediates, and the hydrogenation of -COOH, -CHO, and C=C double bonds in reactants and intermediates, and the acid function catalyzes the dehydration of alcohol intermediates.

¹¹⁰ It was shown that the HDO rate for palmitic acid was greater on Ni/ZrO₂ than on Ni/SiO₂ or Ni/Al₂O₃ but was slower than that on H-zeolite-supported Ni.²¹³ In the absence of H₂, ketonization catalyzed by ZrO₂ is the dominant reaction (Fig. 19B). Pd/C favors direct DCO (-CO₂), whereas Pt/C and Raney ¹¹⁵ Ni favor decarbonylation pathway (-CO). The deoxygenation rate of palmitic acid decreases in the sequence of r(Pt black) \approx r(Pd

black) > r(Raney Ni) without H₂. The reaction mechanism is the same for either noble mental or transition metal. The development of more affordable catalysts with similar performance and durability is of great interest from an industrial ⁵ standpoint because noble metals are costly.

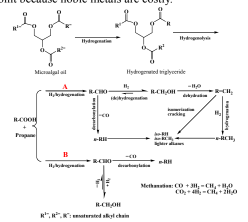


Fig. 19 Proposed reaction pathway for the transformation of microalgal oil to alkanes over different catalysts

A: Bifunctional Ni/HBeta catalysts²¹¹ Copyright 2009 Wiley; B: Ni/ZrO₂ catalyst²¹³ Copyright 2012 Wiley

5.2 Upgrading of real bio-oil from HTL microalgae

Bio-oils produced from microalgae via HTL are formed in an aqueous environment. From the view of engineering, oil ¹⁵ upgrading under the same environment may be more advantageous. Thus, bio-oil treatment in sub- or supercritical aqueous environment may be an effective approach for oil upgrading.

Duan published a series of related works^{35,214-216}. Table 6 ²⁰ provides a summary of Duan's contribution to bio-oil research. The combination of high temperature and H_2 increase the HHV of

bio-oil. In addition, the performance of carbon-based catalysts is superior to that of other catalysts.

In Duan's work, the upgraded bio-oil had better properties ²⁵ after processing with Pt/C catalyst and high-pressure H₂ in SCW. The bio-oil had a HHV of 43 MJ·kg⁻¹, and the total acid number of the upgraded oil (25) was considerably lower than that of original feed (256). Finally, the upgraded bio-oil had a very high content of hydrocarbon molecules, including alkanes and ³⁰ aromatic compounds. Overall, the properties of the upgraded oil obtained from the catalytic treatment in SCW are similar to those of hydrocarbon fuels derived from petroleum. This work shows that the crude bio-oil from the HTL of microalgae can be effectively upgraded in SCW in the presence of a Pt/C catalyst.

On the other hand, molecular breakdown of the crude and upgraded oils are clearly different. The chromatogram for the crude bio-oil shows minimal material eluting prior to 40 min. By contrast, the upgraded oil shows some large peaks at retention times shorter than 12 min and many regularly spaced peaks, 40 which correspond to a series of n-alkanes starting at about C₉. It indicated the catalytic hydrothermal upgrading process produced oil with more low-boiling species.

During upgrading of crude algal bio-oil in SCW, different types of reactions occur simultaneously. In addition to 45 hydrogenation, hydration and/or oxygenation reaction also occur, which involve water in the upgrading chemistry, leading to the incorporation of H and O atoms into the upgraded oil.

The catalyst must endure erosion from sub/supercritical system, and the effect of H₂O on bio-oil treated, which was not ⁵⁰ found in the published papers, should be considered. Most results were obtained on a small scale, and product characterization has mainly been restricted to GC-MS. Bio-oils from sub/supercritical system treatment are still far away from direct use as transportation fuel because the product oil still contained some ⁵⁵ oxygenated compounds and nitrogen-containing compounds. Therefore, additional treatment and process optimization is needed.

	*	-			
No.	Catalyst	T/K	Ambience	HHV/MJ·Mg ⁻¹	Ref.
1	Pd/C, Pt/C, Ru/C,Ni/SiO ₂ -Al ₂ O ₃ , CoMo/γ-Al ₂ O ₃	623	He	38.0	[35]
2	Pt/C, Mo ₂ C, HZSM-5	703-803	He	42.0-43.0	[214]
3	Pt/C	673	H_2	43.0	[215]
4	Pt/γ-Al ₂ O ₃	673	H_2	-	[216]

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Table 6 Comparison of Duan's contribution to catalytic upgrading of algal bio-oil

6 Conclusions and perspectives

⁶⁰ In recent years, the production of renewable fuel obtained from microalgae has attracted considerable attention because of algal fast growth rate, minimal competition with food crops, and other factors. Therefore, microalgal-based biofuels have a paramount role to play in combating energy shortage, global ⁶⁵ warming and climate changes. Thus, there is tremendous potential for this field and the outlook is bright.

Nevertheless, research for the production of biofuels from microalgae is in the early stages, we are still on some way from realizing the potential to produce commercially viable microalgal

⁷⁰ biofuels at a large scale. Sustained, in-depth research is needed to accelerate the practical use of microalgae as energy feedstock, and enhancement of bio-oil production yield and energy efficiency of the process is our overall goal.

In summary, thermochemical conversion of microalgal 75 feedstock is still in the developmental stage, and challenges in the coming years include the following:

• Feedstock provide and algal cultivation.

The main hurdle of algae-based biofuels to be solved is the cost gap, since the technologies (including cultivation, harvesting and conversion) accessible today cannot make microalgal biofuels production economical. The cost of the infrastructure facilities and the energy required for microalgae cultivation and harvesting are high, for example, the drying operation of the microalgal biomass consume intense input energy.

The growth rate of algae is fast in comparison to other plants, but in the context of the overall economics higher rates will be needed to reduce the aforementioned land

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requirements. Genetic engineering of algal strains with high growth rate and energy content is critical. In addition, it 60 appears that choice of an appropriate algal strain and cultivation conditions possibly incorporating flue gas as a source of carbon

• Life cycle analysis

Microalgae obviously require CO_2 for growth, which raises the importance of coupling their growth with CO_2 producing facilities, such as cement and coal-fired power

¹⁰ plants. For example, the use of CO_2 absorption columns with algae growth and harvesting comprises a synergistic approach that provides integrated solutions, which could bring costs of CO_2 capture and utilization down.

Production volume, land footprint and water requirement. The sheer volume of transportation fuel consumption worldwide is daunting. For example, the U. S. alone consumes about 19 million barrels of petroleum products per day. Of that total, about two-thirds are used for transportation. Based on a conservative estimate for growth

rate of 81 g dry algae/m² day, and assuming that 25% of the intrinsic energy content is utilized towards bio-oil, 25,000 km² (6.2 million acres) of land area, 8.466E+8 m³ saline water and 7.914E+9 m³ fresh water would be needed to produce the equivalent of 1,000,000 barrels of oil equivalent/day. Thus, a vast land area and water would be needed.

Process coupling and co-processing.

Due to the high cost, algal cultivation for biofuel production alone is hard to achieve cost effectiveness and a positive energy balance. One possible solution to reduce the algae production cost and process energy cost is to combine the algal cultivation with current wastewater treatment process.

In addition, co-processing of microalgae with other feedstocks could also help reduce the overall cost, co-process with waste plastics is such a good example. Waste plastics caused serious environment pollution, which can be regarded as "white pollution". Addition of a fraction of the waste plastics during the thermochemical conversion of microalgae

40 can obtain higher yield and higher oil quality due to role of waste plastic as hydrogen source and presence of a synergistic effect in co-process.

• Specific issue in conversion process

a. Catalyst.

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- 45 Catalysts are crucial in the production of bio-oil from algae. The selection of proper catalysts for the conversion of algae to desired products is a complex process. Promising catalyst candidates must be identified from vast numbers of trial compounds to provide useful information
- about active sites, optimized structures and composition, and possible synthesis routes. The majority of the work to date on producing liquid fuels from algal HTL has focused on homogeneous catalysis by metal salts or alkali. The more recent studies, however, are beginning to examine heterogeneous catalysts due to advantages in separation and selectivity of the catalyst. In particular, the development of non-precious metal based catalysts would provide a major

advance. Finally, there is a need for more catalyst

development work to identify supports and active components that better resist deactivation in hot compressed water or supercritical water.

b. Reaction mechanisms and kinetics.

Understanding the mechanisms of algal thermochemical conversions, especially catalysis mechanism, is the critical issue for improving conversion processes and guide the experimental research, and further studies could provide insights into the catalysis mechanism, especially concerning the catalyst deactivation.

In addition, process kinetics is extremely to optimized algal conversion reaction and design the pilot-scale reactor in the next step. The effects of various feedstock components and operating parameters on the yield and quality of bio-oil must be further investigated to identify the optimal processing conditions because numerous algae species could be used as feedstock and different reaction condition could be conducted.

c. Continuous pilot-scale reactor

Development of pilot-scale plant is needed to accelerate algal conversion process profitability and economic feasibility. However, most of the algal conversion reactions reported have been on a small scale (less than 10 g algae), and only very few attempted algal pyrolysis and HTL have been carried out in continuous reactor systems. Designing a large reactor is very difficult because the mechanisms involves complex process reactions are not yet clearly understood. Finally, during scale up of the HTL process, technical difficulties arise when pumping algae slurry into high pressure reactors and

d. Product characterization

Understanding the chemical composition of bio-oil is required to evaluating conversion process and optimizing the upgrading of crude-oil. However, the small scale of most algal conversion reactions results in too little sample being obtained to distill the liquid product. Thus product characterization has been limited to GC-MS and elemental analysis for the liquids. When larger amount of sample be obtained, other analytic method, such as ¹H NMR spectra, can be conducted, ¹H NMR can provide more detailed information about the composition of the total liquid product.

e. Computational modelling

The computational modelling for algal thermochemical conversion so far is still very limited due to lack of accumulated knowledge in this relatively new field, most related work has been conducted on simplified model systems. More simulation research in this field is expected to be performed in the future, and direct simulation of real algal systems is highly desired. To this end, algae and its conversion process should be understood more deeply and efficient new methodologies, which are capable of simulating large real algal conversion systems, should be developed.

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