

A Review on Oil Spill Clean-up Using Bio-Sorbent Materials with Special Emphasis on Utilization of Kenaf Core Fibers

Jen Yen Tan,^a Siew Yan Low,^a Zhen Hong Ban,^{a,b} and Parthiban Siwayanan^{a,b,*}

Thousands of oil spill incidents regrettably have occurred on a daily basis around the globe. Oil spills are a form of pollution that poses major catastrophic threats to marine ecosystems. Oil spill incidents commonly occur in the middle of the ocean, and the process to remove the oil becomes a great challenge. Absorbents, due to their good oil-absorbing characteristics, are becoming more popular nowadays as one of the effective oil spill clean-up methods. Among the many types of oil absorbents, plant fibers were also being studied to remove the oil spills from the sea surface. This paper reviews factors leading to oil spills, the detrimental effect of the oil spill on the environment, the oil spill removal methods, and the application of plant fibers as oil absorbent material. The paper also will highlight the latest development on the utilization of kenaf core fibers as oil-absorbent material and the use of oil absorbed kenaf core fibers as fuel briquette by taking advantage the high calorific value of the oil spill.

Keywords: Oil spill; Marine pollution; Oil sorbents; Kenaf core fiber

Contact information: a: School of Energy and Chemical Engineering, Xiamen University Malaysia Jalan Sunsuria, Bandar Sunsuria, 43900 Sepang, Selangor, MALAYSIA; b: College of Chemistry and Chemical Engineering, Xiamen University, Xiamen 361005, Fujian, CHINA;

* Correspondence to: parthiban.siwayanan@xmu.edu.my

INTRODUCTION

Factors Causing Marine Oil Spills

In recent years, the world still revolves around fossil fuel products, especially petroleum derivatives. These fossil fuel products require transportation using sea routes. Unfortunately, some accidents such as tanker wrecks and oil exploration release a large amount of oil into the sea and cause a huge impact on the marine ecosystem. Marine species such as birds and marine invertebrates are life-threatened by such accidents, not to mention clean-up work that is intensively carried out requires huge amount of manpower (Alonso-Alvarez *et al.* 2007; Aguilera *et al.* 2010).

The Deepwater Horizon oil spill that occurred in 2012 spilled 4 million barrels of oil into the sea and caused irreversible damage to the environment. The spilled oil even reached the wetlands of Louisiana in the United States and damaged the wetland ecosystem. It is estimated that hundreds of thousands of seabirds died due to starving and drowning, as their feathers were coated with oil (Balmer 2014; Potenza 2016). There is a decreasing trend on oil spills due to the increasing awareness of safety management and technical inspection on oil tankers. Despite that, there is room for improvements to be made regarding oil spill governance. In Malaysia, the amount of crude oil and petroleum products that are transported through the Strait of Malacca is increasing steadily throughout the years

(Villar and Hamilton 2017). Sources for oil spills include oil and gas exploration, transportation and storage, oil tanker accidents and leakages, accidents during ship loading/discharging, ship ballasting/de-ballasting, bunkering, discharging effluent from the petrochemical industry, or bursting pipelines (Ossai *et al.* 2020). Most of the oil spillages are occurring in the Atlantic Ocean, as the ship routes are concentrated there. Complex water situations can be caused by collision between oil tankers or ships, technical errors, bad weather, fire accidents due to electrical sparks, or wrong decisions made by the authorities (Galieriková and Materna 2020). The shipping industry is prone to oil spills. An analysis done by Chen *et al.* (2018) shows that an oil spill easily can occur during refuelling operations, causing ignition of fire and explosion. In the event of anchoring in open water or inland water, the ships are highly prone to collision. Other than that, the failing operating equipment and fire/explosion during cargo loading/unloading are crucial factors contributing to oil spillage.

Effects of Marine Oil Spill to Environment

Oil spills impose adverse impacts on aquatic life, threaten fisheries, and lead to loss of jobs. For example, the mode of nutrition of shellfish will lead to biological accumulation of oil spills particulates and/or chemicals in the oil spill area (Ferguson *et al.* 2020). When petroleum hydrocarbon enters the water, higher molecular weight hydrocarbons will sink to the bottom while the lighter and less viscous crude oil will float on the water surface. Weathering process will spread, evaporate, disperse, or cause emulsification to the oil, affecting the spreading rate of the thin surface films and slicks (Mishra and Kumar 2015).

Different composition of the petroleum hydrocarbon imposes different effects on the environment. Polycyclic aromatic hydrocarbons (PAHs) are one of the components in oil spills. PAHs are toxic and have mutagenic and/or carcinogenic properties. As they are highly lipid soluble, they are readily absorbed from the gastrointestinal tract when accidentally consumed by the aquatic life (Abdel-Shafy and Mansour 2016). Other components present include straight and branched chain alkanes, alkenes, alkynes, sulphur compound, oxygen compound, nitrogen compound, and certain metals. These components cause health effect such as haemotoxicity, mutagenicity, and cytotoxicity to humans who consume contaminated seafood (Ossai *et al.* 2020).

Marine Oil Waste Removal Techniques/ Technologies/ Processes

Physical removal of oil waste is one of the available methods. An oil skimmer is capable of collecting and removing a variety of oil on the water (Song *et al.* 2015). Containment booms are applied on the oil spill area to confine or prevent it from spreading to other areas. They can be used to change the direction flow or concentrate the oil spill for further action such as burning (Fingas 2011). *In-situ* burning is suitable for application in severe oil spill events to burn and remove the oil from the water surface at high rate and efficiency. Biological methods include biodegradation and bioremediation of oil spill by applying microorganisms to degrade the hydrocarbon in the target area (Atlas and Atlas 1991). Dispersants is another class of chemical that can be used for oil spill treatment. Detergent-like products are applied onto the layers of oil and diffuse to the oil/water interface. As a result, the surface tension between oil and water droplets are broken, hence the oil-water layers can be scattered into tiny droplets (Lessard and DeMarco 2000). Other methods include coagulation, flocculation, coalescences, membrane filtration and vacuuming (Dave and Ghaly 2011). Each oil spill removal method faces its own challenges in oil spill removal.

Although mechanical and chemical treatment are commonly used, they require higher cost. Mechanical treatment faces challenges under rough sea waves along with high wind velocity. Chemical treatment requires post-treatment such as dispersion to reduce toxicity effect on marine creatures (Alaa El-Din *et al.* 2018). *In-situ* burning releases carbon dioxide into the atmosphere upon combustion. This method is less applicable when the spilled oil forms a stable emulsion which consists of more than 25% of water, as such oil slicks are unable to be ignited (Mullin and Champ 2003). Bioremediation has many advantages in terms of environmental aspect and economic properties. However, the biodegradation rate of oil spill by the microorganisms depends on many factors such as concentration of oil, availability of other nutrients, environmental factors, as well as different activities of microorganisms. During the initial application of microorganism where the oil concentration is high, the biodegradation process shows undesirable result. Another major concern for this method is the time needed to clean up the oil spill (Dave and Ghaly 2011). While methods involve coagulation have high oil removal efficiency, they require high operating cost and they are labor intensive as complex operation are required. Some of these methods will generate a second source of pollutant (Wahi *et al.* 2013).

Oil spill sorbents are relatively insensitive to sea conditions, easy to use, and economic feasible (Al-Majed *et al.* 2012). There are two types of sorbent, namely natural and synthetic sorbents. Each type of sorbent has its disadvantage. Natural organic sorbents are very difficult to be collected and labor intensive because they will absorb some of the seawater. Many of the natural inorganic sorbent are loose material and can be difficult to be applied in windy conditions. Synthetic sorbents, which are sometimes modified into super-hydrophobic materials, can be suitable to be reused many times. However, many are chemically synthesized and not biodegradable. In addition, sorbent collection after oil sorption requires high manpower (Dave and Ghaly 2011). The oil-soaked absorbent will be disposed of by burning in an industrial incinerator. The density and viscosity of the oil normally will be altered after the weathering process. Sea conditions such as waves and currents will further disperse the oil slick will be further dispersed, changing their physical and chemical properties. Photo-oxidation of the oil will produce a variety of oxidized compounds which become more water-soluble (Rogowska and Namiesnik 2010).

Researchers are continually researching and modifying synthetic sorbents to improve their performance. For example, Kukkar *et al.* (2020) engineered carbon nanotube (CNT) materials to super-hydrophobic products such as CNT aerogel/sponge sorbents, which showed excellent oil sorption capacity (80 to 180g/g) and can be reused up to 1000 cycles. Light hydrophobic sorbents can be synthesized to deal with the light oil in the oil spill. For example, cellulose/graphene aerogels show high sorption capacity against many types of oil, including light oil such as hexane (Mi *et al.* 2018). Therefore, it is crucial to synthesize sorbent that could deal with different oil compounds. Reduced graphene oxide (rGO) coated on porous polyurethane (PU) is a cost-effective and highly hydrophobic sorbent. It has been shown to be effective in cleaning up oil spill, reusable and can absorb various organic solvents (Zhu *et al.* 2016). However, sorbent is often used in small spills and needs to be combined with other methods. Once being used in oil absorption, they need to be removed from water and they are re-used, recycled, or disposed (Rengasamy *et al.* 2011a). As mentioned above, one of the problems for using sorbents is the collection after oil sorption. To ease in the collection process, the sorbents can be wrapped inside a net that will keep them afloat (Bayat *et al.* 2005). On the other hand, mixing of different oil removal techniques can be proposed. The advantages of booms are easy to handle and flexible for

towing. For example, fence booms and curtain booms can be used. However, the collected oil need further treatment. Sorbents could be applied together with booms to increase the oil spill cleanup efficiency (Dave and Ghaly 2011). Skimmers are applied in the area to recover the oil spill when there is a floating layer. To deal with the sea environment, oil spill sorbent can be applied in the targeted area. Elevating skimmers can be combined with sorbent for more effective and faster processing (Hoang *et al.* 2018).

UTILIZATION OF PLANT FIBER AS OIL SPILL SORBENT

Application of Bio-sorbent in Oil Spill

This section focuses on environmentally friendly natural sorbents. Natural sorbents can be classified into organic and inorganic sorbents. An example of an inorganic sorbent is a natural occurring mineral, such as perlite. Perlite is an environmentally friendly oil sorbent that shows good oil sorption capacity (Teas *et al.* 2001). On the other hand, organic sorbent can be further classified into animal origin and plant origin. Organic sorbents or bio-sorbents are gaining attention, as they have shown comparable oil sorption capacity with synthetic sorbents. They seem promising due to their feasibility and effectiveness, as no additional chemicals are needed to demulsify the oil (Wahi *et al.* 2013). Animal origin sorbents are obtained without harming their lives. For example, molting cuticle from tarantulas is being studied as a practical option for oil sorption. Based on average spider biomass per square meter, the global spider community has estimated that the spiders can produce 2 and 6 million tons of cuticle (depending on the frequency of molting) annually, and this waste material can be collected from 190,000 spider breeders worldwide. The oil sorption mechanism can be explained by physical trapping of crude oil drops *via* the brush-like structure of the cuticle. The irregular branched hairs trap and retain the crude oil. The presence of wax on the spider cuticle improves its hydrophobicity by promoting the action of van der Waals forces and other chemical interactions during the oil sorption (Machalowski *et al.* 2020).

Raw natural wool that shows high oil sorption capacity and good buoyancy is being tested for oil spill clean-up. Ciufu *et al.* (2019) obtained a sorption result of 2.66 to 11.89 g/g using crude oil. The good performance of wool is linked with the natural covering of wool grease, lanolin. During oil sorption, a thin layer of oil will be retained on the wool fibers, making them more hydrophobic (Johnson *et al.* 1973). According to Choi and Moreau (1993), the presence of oleophilic wax on the wool fibers aid in the adsorption of oil onto it. Next, the scaly structure of wool forms a deposition area for the oil as well as forming capillary bridges between oil and fiber. The medulla honeycomb structure also provides an area for oil retention. Human hair (50 to 100 μm diameter) shows good result in oil sorption. Human hair contains porous cortex, and its hydrophobicity varies with different type of hair. African origin hair has higher oil sorption capacity than Asian origin and European origin. It shows twice of oil sorption in comparison to the latter hair types. This is due to its coarse and spongy nature which contains more macro-pores. On the other hand, ground hair shows higher oil sorption capacity as its oil sorption surface area increased (Ifelebuegu *et al.* 2015). Researcher are researching different natural sorbent for oil spill removal as they are easily available. This reduces the usage of synthetic sorbent (Dave and Ghaly 2011). As heavier oil such as heavy crude oil can adhere to the sorbent easily regardless of natural or synthetic sorbent, natural sorbent can be used as greener alternative (Bayat *et al.* 2005).

Characteristics of Sorbent Material for Liquid Hydrocarbon Removal

Wettability and hydrophobicity

Natural fibers that are sourced from plants generally are more abundant and more sustainable. They are chemical-free, biodegradable, and some even show comparable density with synthetic sorbent. Many of the raw natural sorbents exhibit excellent sorption and they have been widely studied for the substitution of synthetic sorbent in oil spill. The most commonly investigated natural oil sorbents are kapok, milkweed fiber, sugarcane bagasse, rice husk, and various plant materials (Wahi *et al.* 2013). For example, cattail fiber and kapok fiber absorbed 827 and 1107 mg oil /g sorbent respectively, compared to 1008 mg oil /g sorbent of polyester fiber (Khan *et al.* 2004). Other plant fibers that have been tested for oil spill treatment include cotton grass fiber (Suni *et al.* 2004), vegetable fibers (Annunciado *et al.* 2005), and straw biomass waste (Tijani *et al.* 2016).

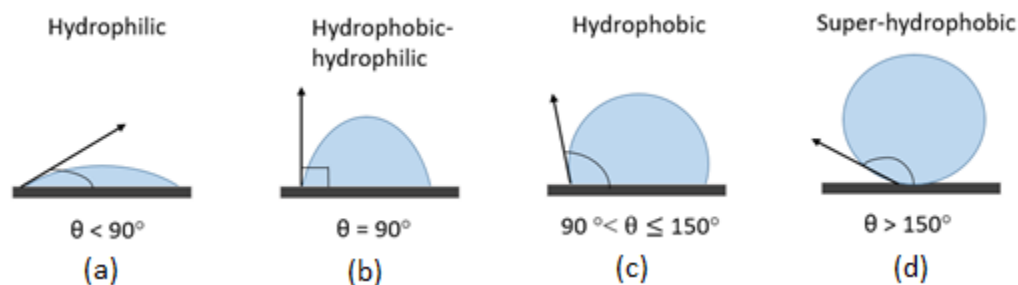


Fig. 1. Water contact angle on the fiber and its wettability (Doshi *et al.* 2018; Creative Commons CC-BY-NC-ND)

The wettability of water on the sorbent surface determines the oil sorption ability. Herein, wettability is measured by the water contact angle. The sorbent is considered as hydrophilic if the water droplet forms a contact angle on its flat surface that is less than 90° (Fig. 1a). The sorbent is regarded to be intermediate-wet if the contact angle is equal to 90° (Fig. 1b). For the contact angle between 90° and 150° (Fig. 1c), it is hydrophobic and if the contact angle exceeds 150° (Fig. 1d), then the sorbent is considered as super-hydrophobic. A sorbent with good hydrophobicity enhances the oil sorption capacity when water is present (Doshi *et al.* 2018). The finding of Lim and Huang (2007a) are consistent with this statement. The hydrophobic surface of kapok fiber is extracted and evenly deposited on a microscope glass. The contact angle for water droplet is observed to be 117° , whereas the contact angle for a diesel droplet is 13° . Kapok fiber is concluded to be oleophilic. Blue-colored water droplets are observed to form a large contact angle with the *Calotropis gigantea* fiber. The water droplets remain nearly spherical on the fibers with the contact angle of 129° ; this represents excellent hydrophobic behavior (Zheng *et al.* 2016).

A hydrophobic natural sorbent encourages high sorption of oil in the oil-water environment. Natural hydrophobic fibers are associated with the presence of wax on the fiber surfaces (Khan *et al.* 2004; Dong *et al.* 2015). Wax present on the fiber surface promotes interaction between oil and fiber surface *via* hydrophobic interaction and van der Waals forces. According to Cui *et al.* (2012), the presence of surface wax on fiber repels water droplets and reduces the contact angle between oil droplets and fiber surface. This reduces the surface tension, which leads to minimum energy barrier for oil sorption. The importance of fiber wax can be seen in kapok fiber, as water cannot penetrate into its lumen. The large contact angle ($>90^\circ$) between fiber wall and water forms negative capillary entry pressure and large surface tension against air in the lumen. This reduces the absorption of

water (Rengasamy *et al.* 2011a). Fibers such as sugarcane bagasse and rice husk that are lacking hydrophobic wax have high affinity for water sorption, thus reducing its oil sorption capacity. Rice husk after absorbing water even sunk to the bottom of container, thus eliminating its possibility to be used as sorbent in oil spill (Ali *et al.* 2012).

Oil sorption mechanism

Oil absorption has been mainly attributed to the capillary force *via* the lumen inside plant fibers (Lim and Huang 2007b). Fiber that shows good oil sorption capacity generally has hollow lumens, which act as the main site for capillary action to take place. A non-collapsing lumen such as the lumen in milkweed and kapok is essential for high oil sorption capacity. Choi and Moreau (1993) reported the movement of oil during oil sorption in plant fibers using ESEM micrographs. They eliminated other factors such as deposition or evaporation and confirmed that capillary action took place. Various researchers also confirmed that capillary action is the main sorption mechanism. Hydrophobic surfaces are the “promoter” of oil sorption. After diffusion of oil into the fiber *via* pores or lumen, capillary forces act as the main driving force (Dong *et al.* 2015; Fasanya *et al.* 2020; Haussard *et al.* 2003). Capillary forces are also responsible for the entrapment of oil inside the fiber, otherwise known as the oil retention capacity. There are three fractions of retention of oil in the fiber. The first oil fraction can be retained by weak capillary forces, but it is quickly released under gravitational force. The second oil fraction is retained by capillary force and can be retained under gravitational force. The third oil fraction is retained by capillary force and adsorptive force, which will not be released under gravitational and centrifugal forces (Cojocar *et al.* 2011).

On the other hand, the viscosity of the oil has huge impact on the oil sorption performance. Oil with high viscosity adheres easily onto the surface of fiber (Choi and Cloud 1992). Higher oil viscosity leads to higher initial sorption ratio of the fiber. This is because viscous oil is heavier (with higher specific gravity) and are absorbed onto the fiber surface easily. Consequently, viscous oil also desorbs easily during the drainage period (Rengasamy *et al.* 2011a). However, this does not mean that plant fiber oil sorption performance is good in the case of a highly viscous oil spill. When oil viscosity increases, the oil sorption capacity of the fiber decreases. A more viscous oil decreases the sorption kinetics as capillary action is sensitive towards oil viscosity. Hence, the rate of oil sorption is different for each fiber, depending on the fiber natural properties (Deschamps *et al.* 2003). This finding is consistent with the findings of Cao *et al.* (2016). Their results indicate that more viscous oil spreads at a lower rate on the surface of plant fiber. In their research, the most viscous engine oil took the longest time to be completely adsorbed by cattail fiber. For the sorption kinetics, oil sorption speed is relatively higher at the initial phase. Then, it gradually decreases when the process reach sorption equilibrium. Thilagavathi *et al.* (2018) explained that small diameter pores are filled up first, which resulted in high initial sorption rate. Xu *et al.* (2019) concluded that the area of available sorption site decreases after some time, resulted in decreasing amount of oil uptake after the initial quick sorption period.

Adsorption capacity for oil spill uptake plays an important role in determining the oil sorption efficiency. The desorption isotherm of a volatile hydrocarbon, which can be determined gravimetrically, could provide some insights of the oil retention mechanism. Generally, capillary retained liquid has the tendency to evaporate faster than the adsorbed film of liquid. Rotaru *et al.* (2014) carried out benzene desorption isotherm analysis and reported that capillary action is the dominant sorption mechanism compared to adsorption in sorbent. This finding is consistent with the finding of Cojocar *et al.* (2017). They carried

out a cyclohexane desorption isotherm experiment to determine which mechanism has larger impact and concluded that capillary action is the dominant mechanism. Furthermore, most of the liquid retained by capillary forces can be removed by centrifugation or squeezing, while the liquid retained by adsorptive force cannot be removed by centrifugation. In natural plant fiber, adsorption takes place to a lesser degree. Researchers often modify plant fiber in order to increase the oil adsorption. Lv *et al.* (2018) modified the jute fiber by introducing SiO₂ to the fiber surface. The modified fiber exhibited a higher specific area than raw fiber, which provided a higher adsorption area for oil. On the other hand, water adsorption capacity of fiber decreases. The similar trend is reported by Sidik *et al.* (2012). Compared to modified oil-palm leaves, the raw oil-palm leaves showed less adsorption of oil. FTIR spectra showed that untreated oil-palm leaves have less crude oil adsorption *via* less intensified peak across several regions. Adsorption capacity of plant fiber depends on the availability of fine fibrous structure forming the capillaries and sorbent surface structure (Angelova *et al.* 2011).

Porosity and fiber structure

Porosity plays important roles in oil sorption capacity. The oil sorption capacity is directly proportional to the porosity of a fiber, but only up to a certain extent. Capillary forces will be overcome by gravitational force in fiber with porosities larger than 0.98 (volume fraction of non-fiber), hence lowering the oil sorption capacity (Rengasamy *et al.* 2011b). By increasing the loose fiber assembly porosity up to 0.97, the oil sorption capacity increases (Thilagavathi *et al.* 2018). Beside porosity, the surface structures of fiber play important roles in determining fiber oil sorption capacity. A rougher surface generally has higher retention capability of absorbed oil. Coarse surface on the fiber increases the surface area for adhesion of oil (Wang *et al.* 2013). The aquatic plant *Salvinia sp.* contains a large amount of hydrophobic fine hairs on its surface. The “hydrophobic space” in the hairs are able to penetrate into the water-oil emulsion while only allowing oil to break through this space (Ribeiro *et al.* 2003). In the case of cattail fiber, its cross-section looks like a belt that is contracted in the middle and protruded on both sides, or “>--<”. Due to this special structure, cattail fiber has a larger surface area for absorption and retention of oil (Cui *et al.* 2012). Biomass biproducts such as fruit peel seem to have promising oil adsorption capacity, as their cellulose structure can retain the heavy oil such as lubricant oil and petrol oil (M. Abdullah *et al.* 2016). However, when it comes to reusability, natural fiber oil sorption capacity drops after the first absorption cycle. For example, there is still 20 to 23% oil trapped inside fiber lumen even after desorption. This causes the deformation of fiber structure as the inter-fiber distance is already contracted after the first oil sorption. Trapped oil also reduces the oil sorption capacity as they are still present in some of the lumens (Abdullah *et al.* 2010). Compared to synthetic sorbent, plant fiber oil sorption efficiency decreases with usage cycle. This is because the fiber structure is altered after tearing or crushing in the squeezing operation (Abdelwahab 2014).

Retention Profiles

Oil retention capacity is an important parameter in evaluating the sorbent. In many research articles the focus is on dynamic oil retention, oil retention, retention ratio, or retention profile. Chai *et al.* (2015) compared the weight of wet material to determine the oil retention capacity of different modified pomelo peel. The results indicated that the oil can be drained out faster initially before achieving equilibrium. According to Lim and Huang (2007b), oil on the external surfaces of the fiber has the tendency to drip out

instantaneously, and in the process of achieving equilibrium, extra-lumen liquid will be drained out slowly due to the insufficient capillary pressure holding against the weight of oil. Packing density affects the oil retention capacity of a fiber as higher packing density will result in higher oil retained in the fiber. Less viscous oil will be drained at a faster rate compared to more viscous oil, but it will achieve equilibrium earlier. Bazargan *et al.* (2015) proposed an equation to model the unsteady state loss of the liquid from the dripping sorbent. The instantaneous dripping of liquid is from the external surfaces of the sorbent. When the looser oil is drained, liquid released rate decreases along with stronger capillary forces. Rate of dripping also changes with the structure of the sorbent. Fibrous structure reduces the rate of release of liquid. In this proposed model, capillary force and sorbent structure are taken into account for the sorbent liquid retention. Research done by Radetic *et al.* (2008) is consistent with the model. The dripping liquid can be divided into three zones. In the first zone, the liquid retention drops the highest due to rapid liquid loss. In the second zone, the loss rate decreases while in the third zone, the liquid retention stabilizes. Similar oil retention curves are also reported by Al Zubaidi *et al.* (2016) and Rotaru *et al.* (2014). Wei *et al.* (2003) concluded that light oil tends to be released from sorbent faster with higher release rate compared to the heavy oil. High porosity sorbent has high oil sorption capacity but poor retention capacity. Fine fiber structure reduces the rate of release.

Issues using Plant Fiber as Oil Spill Sorbent

Most of the plant fiber will absorb water after some time. Plant fiber that is richer in cellulose and hemicellulose tends to be hydrophilic. The presence of –OH bonds in the cellulose will form hydrogen bonds with the water. Removal of this bond will alter the fiber characteristics, *i.e.* becoming more hydrophobic (Ramadevi 2012). Even hydrophobic fiber will eventually absorb water due to the presence of cellulose. Raw cotton that exhibits excellent oil sorption capacity sinks into water after being put in water for some time. Most of the natural fibers are partly hydrophilic and they are lacking stability in aqueous medium over time after oil sorption (Deschamps *et al.* 2003). Bagasse for example, has both hydrophilic and hydrophobic sites. Although the surface of bagasse is hydrophobic, its cellulosic pulp absorbs water when being put in water-oil mixture. It is observed that its amount of water absorbed increases with the increasing sorption time (Said Ael *et al.* 2009).

Treatment to Enhance Hydrophobicity

Cellulosic materials can be treated to acquire hydrophobic character. Common modification are chemical treatments such as sodium chlorite treatment, acetylation, mercerization, benzylation, graft polymerization, enzymatic treatment, peroxide treatment, and plasma treatment (Ali *et al.* 2016). Herein, low cost, less chemical usage treatment methods are discussed as the goal is to produce cost effective and green sorbent for oil spill absorption.

Chemical Treatment

In acetylation, the OH bonds are reduced and replaced with ester bonds (C=O) and acetyl (C-O) bonds. The latter two bonds have higher hydrophobicity than the hydroxyl groups, which are available for hydrogen bonding with the water. The acetylation process of hydroxyl group turns the raw corn silk into better oil sorption sorbent by enhancing its hydrophobicity. The oil sorption capacity of modified corn increases and it can be reused up to five cycles (Liu *et al.* 2016). Teli and Valia (2013) used acetic anhydride with N-bromosuccinimide (NBS) (2%) as catalyst in the acetylation of banana fiber. The cellulose

content of raw banana fiber was reduced from 57.64% to 50.61% and showed lower water sorption. Sugarcane bagasse, an agricultural waste was acetylated by introducing acetyl groups into the cellulose structure with acetic anhydride and NBS (1%). The hydrophilicity of bleached softwood kraft fibers, on the other hand, can be reduced when these fibers are treated with lignin or alkenylsuccinic anhydride. The ability of such fibers to take up oil, even when wet by water, can be increased by treating them with these reagents (Payne *et al.* 2012).

Modified sugarcane bagasse has oil sorption capacity ranges from 13.5 to 20.2 g/g and desorbs quickly when being squeezed (Sun *et al.* 2004). Mercerization is the treatment of fiber with alkali that break down the fibers bundle into smaller strain. This process often develops rougher surface topography for the fibers (Ali *et al.* 2016). According to Hasim *et al.* (2012), the changes of natural crystalline structure is dependent on the type of alkali and concentration used. Cellulose and other non-cellulose constituent such as hemicellulose, lignin, and pectin are removed. Anuzyte and Vaisis (2018) treated *Rhytidiadelphus triquetrus* (a moss species) with NaOH (5%) and observed that the oil (diesel) sorption capacity was increased by 22% after alkali treatment. The fiber surface became coarser, providing more surface area for oil sorption. Nevertheless, the removal of the cellulose component depends on the concentration of NaOH. For example, increasing NaOH concentration in the alkali treatment increases the abaca hydrophobicity, resulting in lesser absorption of different types of water moisture (Ramadevi 2012).

Grafting

Grafting is a simple and effective method. Natural fiber can be modified by grafting oleophilic groups on the fiber for easier attachment of oil. This process will increase the hydrophobic properties of the fiber. Jarrah *et al.* (2018) modified cotton through silylation and increased its crude oil sorption capacity by more than two folds. The hydroxyl groups on cotton surface are turned into hydrophobic alkylsilyl ether chains. Huang *et al.* (2016) grafted epoxidized soybean oil (ESO) onto the surface of cellulose fiber through epoxide ring-opening polymerization process. The hydroxyl groups on the fiber are replaced with hydrophobic alkanes group. Other than that, cellulose fibers in its wet state can also be modified. Layered double hydroxide (LDH) is grafted into the wet, hydrophilic fibers so that organic groups can directly interact with the fiber *via* electrostatic interactions (Sobhana *et al.* 2017).

Aerogel modification

An aerogel is a solid porous material. Due to its high porosity (up to about 99%) and high surface area (more than 100 m²/g), such materials are good candidates for oil spill sorption. Aerogels can be made from inorganic source or biomaterials (Doshi *et al.* 2018). Herein, low-cost, degradable plant fiber with less chemical derived aerogels will be discussed. They are known as cellulose-based aerogels (CBA). Yin *et al.* (2017) used *Eichhornia crassipes* as the cellulose source and polyvinyl alcohol (PVA) as cross-linking agent to produce CBA. The results for its oil sorption capacity were 77.08 and 91.82 g/g sorbent for crude oil and diesel oil, respectively, and it can be re-used up to 16 cycles. Thai *et al.* (2020) ground sugarcane bagasse before mixed with PVA, followed by freezing drying at -70 °C. As sugarcane bagasse aerogel was still hydrophilic due to the presence of hydroxyl groups on its surface, it was further coated with methyltrimethoxysilane (MTMS). Lightweight, high porosity and hydrophobic twisted carbon fibers (TCF) are produced *via*

pyrolysis of raw cotton. TCF can absorb oil from 50 to 190 times of its weight while remain floating on water surface after sorption of heptane (hydrocarbon) (Bi *et al.* 2013)

High temperature treatment

Hydrothermal treatment is an environmental friendly, simple, and cost effective method for the removal of hydrophilic cellulose components. Nordin *et al.* (2013) removed hemicellulose from palm fiber *via* superheated steam treatment. Anuzyte and Vaisis (2018) used hot water to modify the fiber surface. Higher contact surface area improves the adhesion of oil onto the modified fibers surface but also removes waxy coating and some other volatile components at the same time. Pyrolysis is another option in thermal treatment. Carbonized rice husk has better buoyancy and higher content of SiO₂ oleophilic groups. However, the effect of pyrolysis depends on the natural content of the plant fiber (Angelova *et al.* 2011).

UTILIZATION OF KENAF CORE FIBER AS OIL SPILL SORBENT

Kenaf Plant in Malaysia

Kenaf (*Hibiscus cannabinus* L.) has been studied for the possibility of being used as a natural sorbent for oil spill removal. Kenaf plant is a traditional crop which has high concentration of fiber and is widely available. The Malaysian government even established research and development program to further exploit the commercial value of kenaf. For example, the National Kenaf and Tobacco Board (LKTN) promoted the plantation of local kenaf plant by allocating 2000 hectares of land for this purpose. They estimated that in year 2025, the demand for Kenaf is RM 3.5 billion (Habibu 2018).

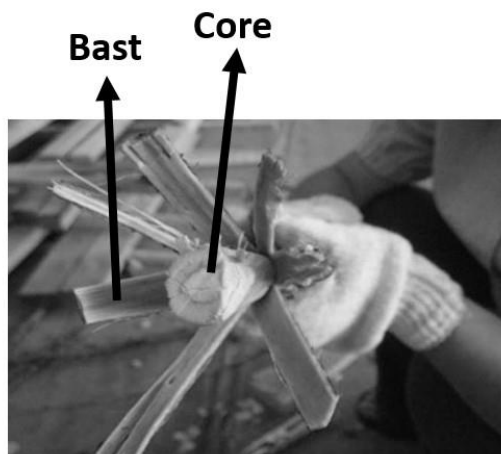


Fig. 2. Kenaf bast and core fiber

Characteristics of Kenaf Bast and Core Fibers

The two main fibers in kenaf are kenaf bast and kenaf core fibers. The bast fibers could be separated from the core fibers through two processes, which are decorticating and retting. Decorticating is the separation of the bast fiber from the core fiber using a decorticator. Retting is the separation of bast fiber from the core fiber through the facilitation of water or chemical soaking. The growth rate for kenaf bast fibers is higher than that of kenaf core fibers, which explains why bast fibers are relatively longer and

stronger than core fiber (Kamal 2014). Malaysian kenaf consists of about 35% bast and 65% core. Its core and bast are rich in lignin and cellulose. Based on the properties of locally obtained kenaf stem, it can be deduced that kenaf core will generate more fibers in term of weight percentage, as kenaf core made up larger proportions of the kenaf stem (Abdul Khalil *et al.* 2010). Kenaf core is more woody compared to kenaf bast, as it contains more lignin. It is mainly used in absorbent applications, filler, animal bedding, and composites (Dilip 2004). The bast fiber, which is rich in cellulose content, is an important material for the paper manufacturing industry. It can be obtained in bundles and can be cut into specific length. Ashori (2006) obtained 45 to 51% pulp yield through kraft pulping process using local kenaf fiber. The bast pulp has excellent density, tear index, as well as dry zero-span breaking length. Kenaf bast fiber is a promising material in the paper and pulp industry.

Table 1 & 2. Characteristics and Properties of Kenaf Stems

Characteristics/ Properties		Bast fiber	Core fiber
Proportion (%)	Cross-section area	21.96	78.04
	Dry Weight	32.2	68.5
Acidity (pH)		7.13	5.21
Reference		(Abdul Khalil <i>et al.</i> 2010)	

	Bast fiber	Core fiber
Cellulose (%)	59.8	37.6 - 49.0
Lignin (%)	17	15.0 – 21.0
Hemi Cellulose (%)	11.6	-
Ether Soluble Extract (%)	1.2	-
Cell Wall Contents	10.4	-
Reference	(Dilip 2004)	

Abdul Khalil *et al.* (2010) carried out FT-IR analysis using local grown kenaf fiber. Peaks at 3390 and 1190 cm^{-1} indicate that there are O-H stretching and O-H bending frequencies. Peak at 2910 cm^{-1} show there is $-\text{CH}_2$ stretching. These peaks indicate the presence of hydroxyl group of cellulose, hemicelluloses, and lignin. Absorbance peaks in the range of 1739 to 1650 cm^{-1} indicate carbonyl stretching (C=O) in the acetyl group in hemicelluloses and aldehyde group in lignin.

Studies on Absorption of Liquid Hydrocarbon using Kenaf Core Fibers as Sorbent Material

Research done by other researchers

Studies on oil spill uptake can be carried out by using the basic standard method, ASTM F726: Standard Test Method for Sorbent Performance of Adsorbents. According to this standard method, 15 min of contact time between liquid and sorbent is suggested, while 15 sec or 2 min are suggested for the dripping time. However, this standard is not that popular among researchers due to some of the shortcomings associated with the standard, which include the non-specified mesh size of net and large scale of sorbent needed (Bazargan *et al.* 2015). However, this standard method can be useful to provide a general insight on oil sorption capability of the kenaf core fibers. Studies have showed that kenaf core fibers has the ability to be used as oil spill sorbent, but the oil recovery is low. Anthony (1994) compared the absorption of oil (non-detergent oil) between kenaf core and bast

fibers at a sorption time of 10 minutes. The result shows that oil recovered (oil recovered from the sorbent by hand squeezing as a percent of oil absorbed) for kenaf core fiber is lesser than that of bast fiber. This can be explained by the absorbed and retained oil in kenaf core fiber. The first run for oil recovery of raw kenaf bast was 29.9%, while for raw kenaf core it was 2.9%. Lee *et al.* (1999) compared the oil sorption capacities of kenaf bast fiber and kenaf core fiber by manipulating the kenaf fiber's physical properties. For example, it is practical to refine them, extract them, and reduce their sizes. Kenaf core fiber shows better oil sorption capacity than that of kenaf bast fiber and smaller fiber size shows higher sorption capacity.

Table 3. Results of Experiment Conducted by Lee *et al.* (1999)

Oil Type	Refined Kenaf Bast (g/g)	Refined Kenaf Core (g/g)	Contact time (min)	Dripping time (min)
Oil/water	1.54	2.74	10	1
Oil/pure	7.08	7.16	10	1
	Ground Kenaf Bast (g/g)	Ground Kenaf Core (g/g)	Contact time (min)	Dripping time (min)
Oil/water	0.45	3.95	10	1
Oil/pure	2.34	6.36	10	1
	Extracted Kenaf Bast (g/g)	Extracted Kenaf Core (g/g)	Contact time (min)	Dripping time (min)
Oil/water	0.31	2.69	10	1
Oil/pure	8.15	7.94	10	1

Oloruntoyin and Olayinka (2016) determined different oil sorption capacities by varying parameters such as particle size, plant genotypes, and type of kenaf fiber. Important parameters such as buoyancy and amount of oil retained were also recorded. The mixture of genotypes records higher oil sorption capacity than single genotype. The results are shown in Table 4.

Table 4. Results of Experiment Conducted by Oloruntoyin and Olayinka (2016)

Product type	Buoyancy	Sorption (g sorbent/ g oil)	Retention (g sorbent/ g oil)	Contact time (hr)	Dripping time (hr)
Spun fiber (2-5 cm particle length)	Sunk	1.66	0.75	4	12
Powdered core	Partially floated	3.00	2.61	4	12
Rough core (1-1.2cm particle length)	Partially floated	1.76	1.56	4	12
Milled core/milled fiber mix (unbagged)	Floated	3.09	2.75	4	12
Whole stem grinded (20 mixed, bagged)	Floated	8.16	5.56	4	12
Whole stem grinded (1 genotype, bagged)	Floated	6.83	6.31	4	12

According to Siwayanan *et al.* (2020), kenaf core fibers tends to absorb crude oil four times of its mass, and 20 mesh kenaf core fiber absorbed more crude oil than 40 mesh

kenaf core fiber. In the published research, the longer the contact time of kenaf core fiber in the crude oil-seawater mixture, the lesser the absorption of oil by the kenaf core fiber. In contrast, the water sorption capacity increases when the contact time increases. The same trend is observed for both 20 and 40 mesh kenaf core fiber.

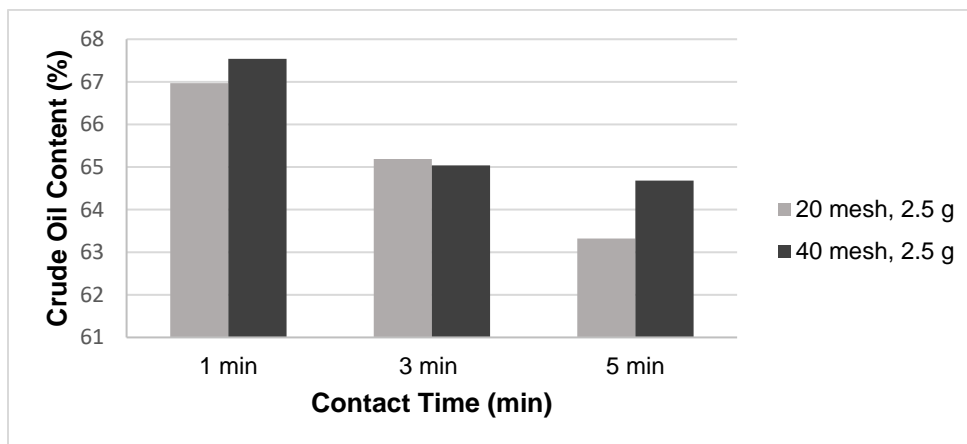


Fig. 3. Crude oil content (%) for 2.5g of 20 and 40 mesh kenaf core fibers at 1 min, 3 min and 5 min of contact time (Siwayanan *et al.* 2020; Creative Commons Attribution 3.0 licence)

Result from researchers in Xiamen University Malaysia (XMUM)

Oil sorption results for kenaf core fibers were obtained from experiments conducted by researchers in XMUM. Kenaf core fibers were either packed in a sachet for easier collection or ground into powder. The kenaf sachet and powder were placed in crude oil, seawater, and crude oil-seawater environments, respectively. The crude oil was obtained from a local refinery plant and seawater is obtained from a close-by sea. In seawater or crude oil environment alone, the maximum absorbed liquid started to appear at around 8 minutes. However, in oil and seawater mixture, prolonged contact time will increase the amount of seawater absorbed, but not the oil. Although 15 minutes of contact time is suggested by ASTM F726, the contact time was controlled at around 2 minutes as it is the optimum contact time of kenaf core fiber in a seawater-crude oil environment. The contact time was optimized in order to achieve the best oil sorption performance for kenaf core fiber.

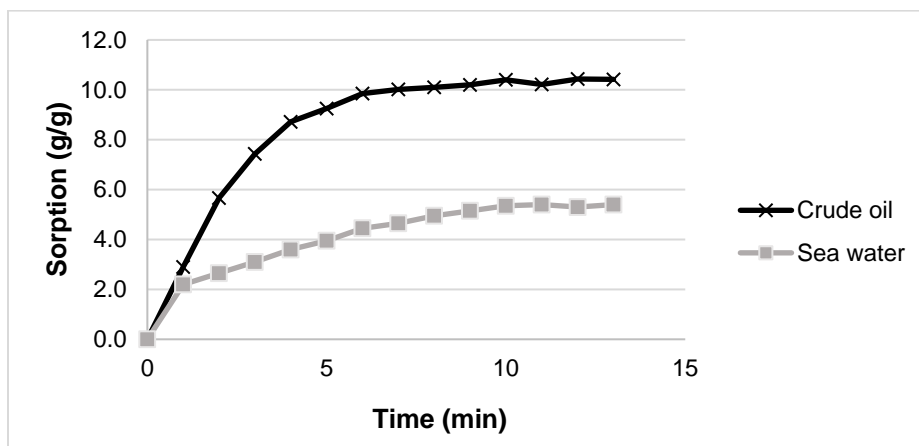


Fig. 4. Sorption capacity of kenaf core fiber against time

Table 5. Result by XMUM Researchers

	Water sorption capacity (g/g)		Oil sorption capacity (g/g)		Contact time (min)	Dripping time (s)
	Kenaf sachet	Kenaf powder	Kenaf sachet	Kenaf powder		
Crude oil environment	N/A	N/A	6.70	8.05	2	15
Sea water environment	4.32	6.25	N/A	N/A	2	15
Crude oil-seawater environment	2.60	5.10	3.60	1.40	2	15

Absorption Mechanism of Kenaf Core Fibers

First, the oil will diffuse and attach onto the kenaf core fiber surface. After that, oil will accumulate in the pores or scaly/rough structures in the sorbent. As discussed before, the main challenge for natural sorbent is that they have high water uptake. Negative ionic charge due to dissociation of the carboxyl groups of the kenaf fiber, brought about by the OH^- ions, will lead to swelling of fiber as well (Nosbi *et al.* 2011). The presence of wax on plant fiber enhances hydrophobic interaction between oil and the fiber (Wahi *et al.* 2013). The presence of wax on kenaf fiber has been verified through Soxhlet extraction using diethyl solvent (Abdul Khalil *et al.* 2010). Most of the hydrophobic substances will be dissolved in the solvent. Hence, it can be concluded that the presence of wax minimizes the surface tension and energy barrier for oil absorption (Abdullah *et al.* 2010). The internal capillary movement would be the main driving force for the absorption of oil within the plant fiber. The capillary action that occurred during oil sorption can be confirmed through SEM image analysis. The crude oil disrupts the orientation in the fiber wall, expanding the inter-fiber pores during absorption (Fig. 5) and resulted in the swelling effect of kenaf core fiber (Siwayanan *et al.* 2020).

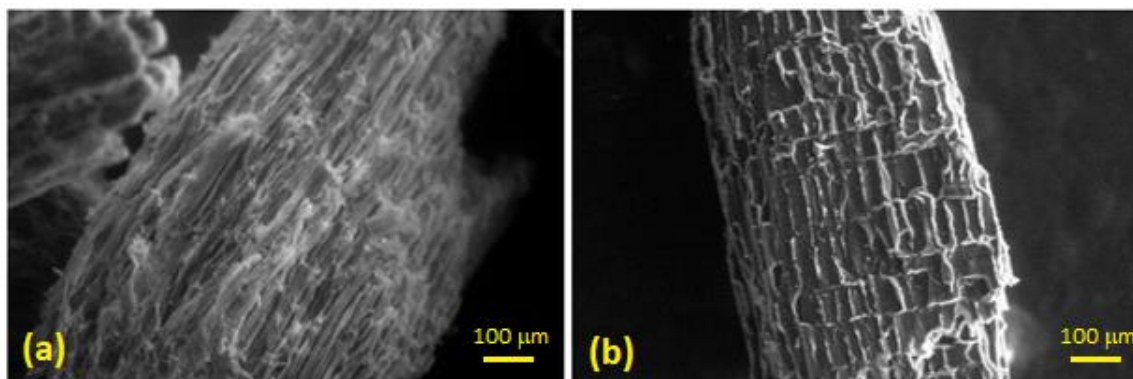


Fig. 5. (a) 20 mesh kenaf core fibers before and (b) after oil absorption, $\times 100$ magnification (Siwayanan *et al.* 2020; Creative Commons Attribution 3.0 licence)

Various studies discussed above showed that kenaf core fiber is a better sorbent than kenaf bast fiber for oil spill sorption. Kenaf bast fiber sank shortly after being put into the water bath, but kenaf core fiber remained floating throughout the experiment. This shows that kenaf core fiber has better buoyancy than kenaf bast fiber and can be easily applied and collected in oil spill environment. The differences in the buoyancy forces

between these two types of kenaf fibers can be attributed by their respective density, porosity, and absorption characteristics. Kenaf core fiber is rich in lignocelluloses components. The transverse section of the kenaf core fiber shows the presence of small hollow fibers and porous structure (Lips *et al.* 2009). This resulted in large absorption area. Cutting the dried kenaf core fiber into smaller pieces increases the exposed surface area. In other words, absorption and retention area are increased. Experimental results above showed that smaller kenaf core fibers have higher oil sorption capacity. From the results obtained from XMUM researchers, the kenaf core fiber seawater sorption capacity increases along with contact time. The presence of hydroxyl bonds facilitates in the absorption of water. It is important to control the contact time for oil spill sorption. This observation is consistent with the result of Siwayanan *et al.* (2020).

Common Disposal Methods for Oil Absorbed Fiber

Using non-buoyant materials as oil adsorbent could transfer the hazardous liquid to the sediment and threaten the food chain and ecosystem of the oil spill area. It is important to test the buoyancy of the sorbent. Only sorbent which passes the buoyancy test is suitable to be used as oil spill sorbent (Oloruntoyin and Olayinka 2016). Bio-sorbent sorption capacity decreases after reusing. The oil absorbed fiber is commonly squeezed, leading to structural damage of the sorbent. Unlike synthetic sorbent which can be reused many times, bio-sorbent such as kenaf core fiber has to be disposed after oil-spilled treatment. Incineration of the oil absorbed fiber could be done, but due to the presence of water in oil-absorbed fiber, evaporation is needed (Hubbe *et al.* 2013). Kenaf core fibers that are soaked with liquid hydrocarbon contain toxic hydrocarbon derivatives compounds such as benzene and polyaromatics. Upon disposing on the land, vapor-phase contamination could lead to odor issues. Over the time, hydrocarbon derivatives could leach directly into the soil and may contaminate groundwater. This might induce more cost to treat the polluted soil (Speight 2010). To evaluate the impacts of treatment of contaminated soil, multiple analysis would be required and this increases the analytical costs. It is difficult to formulate treatment strategies for the contaminated site to degrade the contaminant to safe level (Mrayyan and Battikhi 2005). Incineration of contaminated oil has many disadvantages, such as high operational cost due to requirement of high energy, large space, and environmental pollution (Okoh *et al.* 2019). Alternatively, composting of the polluted soil can be carried out. Composting involves biodegradation of the hydrocarbon contaminants in the soil using micro-organisms. This method has shown some proven viability but is restricted by the time needed and cost (Hubbe *et al.* 2013).

To recover the fuel value of oil, oil absorbed kenaf core fiber can be converted into fuel briquettes. In Malaysia, the government is promoting the use of biomass fuel briquettes in power generating plants (Chiew and Shimada 2013; Sing and Aris 2013). Kenaf fiber can be used as fuel for combustion in boilers, as it is rich in cellulose and lignin compounds (Saba *et al.* 2015). Kenaf core fiber powder after absorbing oil/ hydrocarbon compound in the petroleum-contaminated wastewater has the potential to be used as fuel. There is a research about utilizing kenaf fiber in water treatment. The sludge obtained after treatment is dewatered and dried, such that it can be used as biomass fuel (Tiemeyer 2005). Briquetting of the oil absorbed kenaf core fiber may require binder. Binder can adhere to the solid particles surfaces and generate strong bonds between the particles. Viscous binders harden and form solid bridges after cooling. Thin adsorbed binding layers can result in smoother layers, which will increase the particle contact area and decrease the inter-particles distances. As a result, the molecular bonding increases. Example of binders

are starch, lignin, resin, molasses lime and sodium bicarbonate. Proteins can be also used as binders, as they can be plasticized to increase the strength of the densified product. Example of protein binder derivatives are soybean meal, wheat, rye and barley (Kaliyan and Vance Morey 2009).

CONCLUSION

Plant fibers are gaining in popularity as potential adsorbents for oils, and there is an increasing demand for cheap and environmental friendly sorbents to clean up the oil spills in the sea. Some plant fibers show comparable oil sorption capacity to synthetic sorbent. The hydrophobicity of some plant fibers can be attributed to the presence of wax on their surfaces, porosity, and different type of fiber structures. However, due to the presence of hydroxyl groups, they will absorb water after some time in contact with water. Hence, less chemically intensive treatment methods can be considered to increase their hydrophobicity. Kenaf core fiber is easily available in Malaysia and it is suitable to be used as sorbent for oil spill. Results from various researches show that it has good oil sorption capacity. In summary, the cost and its oleophilicity make kenaf core fiber a suitable oil spill sorbent candidate.

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