Article
Partial Upgrading of Athabasca Bitumen Using Thermal Cracking

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Abstract: The current industry practice is to mix bitumen with a diluent in order to reduce its viscosity before it can be pumped to refineries and upgraders. The recovery of the diluent and its recycling to the producers, on the other hand, pose major environmental and economic concerns. Hence, onsite partial upgrading of the extracted bitumen to pipeline specifications presents an attractive alternative. In this work, thermal cracking of Athabasca bitumen was carried out in an autoclave at 400 °C, 420 °C and 440 °C in presence and absence of drill cuttings catalyst. At 400 °C, despite no coke formation, the reduction in viscosity was insufficient, whereas at 440 °C, the coke yield was significant, ~20 wt.%. A balance between yield and viscosity was found at 420 °C, with 88 ± 5 wt.% liquid, ~5 wt.% coke and a liquid viscosity and °API gravity of 60 ± 20 cSt and 23 ± 3, respectively. Additionally, the sulfur content and the Conradson carbon residue were reduced by 25% and 10%, respectively. The catalytic thermal cracking at 420 °C further improved the quality of the liquid product to 40 ± 6 cSt and 25 ± 2 °API gravity, however at slightly lower liquid yield of 86 ± 6 wt.%. Both catalytic and non-catalytic cracking provide a stable liquid product, which by far exceeds pipeline standards. Although small relative to the energy required for upgrading in general, the pumping energy requirement for the partially upgraded bitumen was 3 times lower than that for diluted bitumen. Lastly, a 5-lump, 6-reaction, kinetic model developed earlier by our group successfully predicted the conversion of the bitumen to the different cuts.

Keywords: thermal cracking; bitumen; asphaltene; diluent; drill cuttings; partial upgrading

1. Introduction

Bitumen type petroleum makes up 15% of the current proven oil reserves [1,2]. The high content of asphaltenes and other heavy molecules in bitumen leads to major logistical problems during their production, transportation and handling [3]. Despite its large abundance, the cost of bitumen production, transportation and upgrading must be kept to a minimum for the resource to be profitable [4]. Several upgrading processes exist, falling under either carbon rejection or hydrogen addition processes. Carbon rejection processes, such as thermal cracking, are typically much more economical at the cost of product quality [5]. The addition of hydrogen, typically in the form of hydrocracking, while applicable to a wider range of feedstock and simultaneously achieves a more desirable octane number product, requires a high-pressure hydrogen atmosphere [5,6]. Thus, an upgrading process can be selected depending on the need and the balance between capital cost and product quality. Regardless of the upgrading route, the general guideline is to maximize the liquid yield, by minimizing coke and gas yields, while attaining high quality, stable liquid product [6,7]. In both types of processes, coke formation during upgrading poses significant operational problems such as catalyst deactivation [8,9]. In addition, the carbon dense coke product is of little value, despite the recent efforts to find markets for petroleum coke [10–13]. Gaseous type products, on the other
hand, while compromising the liquid yield, could still hold some value as they could potentially be used for process heating.

Bitumen transportation is another aspect of bitumen handling that should be addressed. Due to the chemical nature of bitumen, its immediate transport following recovery is not feasible [14,15]. The current, most common, industry practice is to sufficiently reduce the viscosity of bitumen by mixing it with diluents [7,14,16]. Diluents are small chain hydrocarbons, e.g., natural gas condensates largely made up of naphtha [17], that are mixed with bitumen in order to reduce its viscosity and boost its °API gravity to pipeline standards. Pipeline requirements, while slightly varying, are typically a minimum acceptable viscosity of 350 cSt and a maximum acceptable density of 940 kg m\(^{-3}\) (°API gravity = 19) at 15.6 °C [18]. Therefore, the bitumen, a near solid substance at ambient conditions (viscosity > 200,000 cSt), must be mixed with sufficient amounts of diluent, typically 30 vol.%, in order to meet pipeline specifications [6,7,19]. The recovery of diluents is usually carried out through distillation, or otherwise fed together with the bitumen into the upgrader resulting in a constant demand for resupply [17,20]. Subsequently, an inexpensive, simple to operate, onsite upgrading unit to achieve sufficient viscosity and density reduction offers a very attractive alternative.

Partial upgrading, specifically for heavier petroleum sources such as bitumen, has gained much prominence in recent years. A methane upgrading unit at relatively mild conditions of 400 °C and 3 MPa for successful partial upgrading of bitumen in an autoclave was proposed [22–24]. Moreover, a low temperature viscosity reduction scheme to achieve upgrading in situ during production has been attempted to some success [25]. A hydro-conversion process was applied to achieve high sulfur and residue conversion in a fixed-bed reactor configuration [26]. Ideally, a smaller, simpler carbon rejection upgrading unit could be the most desirable option for achieving pipeline requirement on-site, as the potential high complexity and logistical problems of high-pressure hydrocrackers could be avoided [4,27]. Moreover, the product needs only to be stable during the duration of the pumping process. Therefore, a thermal cracking process, or if potentially better liquid quality is desired, a catalytic thermal cracking process would be an ideal choice. To ensure a low process cost and minimal complexity, drill cuttings, a waste by-product from drilling industry, could be used for catalytic upgrading [28]. Drill cuttings from a sandstone or shale formation are typically composed of alumina/silica, which serve as the most common base for a catalytic cracking catalyst. We have shown in our previous work that drill cuttings are effective thermal cracking catalyst [29,30]. They could also be used as effective hydrocracking catalysts, when promoted with hydrogenation elements [31]. As the drill cuttings are a waste product, their re-usability is not a must. They could simply be discarded upon settling after their use in the cracker [32].

In this study, thermal non-catalytic and catalytic cracking in the presence of drill cuttings of Athabasca bitumen was carried out in an autoclave. The objective of the study was to explore a simple, inexpensive upgrading process that could feasibly be used on-site to reduce the viscosity and density of bitumen to acceptable pipeline standards. Effort was made to ensure a one or two-step treatment process, i.e., a reactor unit and a settling unit, that would completely eliminate the need for a diluent. The accompanied reduction in the overall cost could prove very significant and lead to higher profit margin for bitumen in the current competitive oil market. The liquid product quality and yield were the main deciding factors on the efficacy of the proposed upgrading unit. Nevertheless, other factors such as liquid product stability were also considered.

2. Results and Discussion

2.1. Catalyst Characterization

Figure 1 presents FT-IR, N\(_2\) adsorption–desorption, SEM/EDX and particle size characterization of the drill cuttings. The FT-IR spectra in Figure 1a confirm the presence of LAS and BAS on the
drill cuttings surface, per the visible peaks in the presence of pyridine at 1455 cm\(^{-1}\) and 1545 cm\(^{-1}\), respectively [33,34]. A slightly higher peak for LAS is found, suggesting that the ionic chain mechanism largely propagates through initial carbenium ion formation, potentially resulting in additional olefin formation, compared to catalytic cracking over BAS sites [35–38]. Bands at 1000 cm\(^{-1}\) and 600 cm\(^{-1}\) are associated with the natural silica framework and alumina bond stretching, respectively [39,40]. The \(\text{N}_2\) adsorption–desorption curve in Figure 1b is of H3 type hysteresis, which suggests that the drill cuttings have little to no pore volume [41]. Additionally, the surface area estimated by BET is very small, especially compared to traditional zeolite catalysts which surface area typically falls in the 200–600 m\(^2\) g\(^{-1}\) range [35,42,43].

![Figure 1](image)

**Figure 1.** Characterization of drill cuttings. (a) FT-IR spectra of raw drill cuttings (black), raw drill cuttings + pyridine (blue) and pyridine (orange), (b) \(\text{N}_2\) adsorption–desorption curve with reported BET surface area, (c) SEM/EDX imaging of drill cuttings surface (blue = silicon, yellow = aluminum), (d) particle size distribution of raw drill cuttings used in upgrading runs.

However, despite lacking such traditional catalytic properties, the drill cuttings have been shown to possess not only surface catalytic activity, but an additional dispersing effect which is effective at reducing coke formation by exposing the precursor radicals to potential hydrogen donors in the reacting media [29,44–46]. To make up for the lack of catalytic activity, a large concentration, 10 wt.%,
of drill cuttings was confirmed effective in our previous work [30]. Even in a continuous reactor configuration partial upgrading, regeneration or recycling of the cuttings may not be necessary or even desired, as they are a waste product with no value to begin with. SEM/EDX imaging in Figure 1c further confirms the catalytic nature of the cuttings, as silicon and aluminum are present and well distributed on the surface of the cuttings, forming the SiO$_2$/Al$_2$O$_3$ cracking base. Lastly, the particle size distribution in Figure 1d confirms that no cuttings with size > 355 µm are present, and that the majority of the cuttings are < 106 µm. A smaller size catalyst is typically beneficial for a slurry cracking catalyst as it improves the mass and heat transfer effects in the reacting medium [6,7,47].

2.2. Temperature, Pressure and Energy Measurements

Temperature and energy profiles formed during the upgrading runs can be found in Figure 2. Clearly, more severe runs required additional energy input throughout the duration of the upgrading. This suggests that the rate of endothermic reactions increased, and therefore additional heat input was required. It should be noted that we previously showed that the heat loss from the reactor arrangement may be important, but can never overshadow the heat of reactions [48]. Hence, the accelerated rates of cracking, which is known to increase with severity, increased the energy requirement by the reactor [49–51]. Catalytic runs required 10% more energy on average, perhaps in part due to differences in the reaction paths in the presence of the drill cuttings. It has been previously confirmed that through a dispersing effect the drill cuttings promote the formation of volatile components at the expense of decreased maltene yield [29,30].

![Figure 2. Temperature (circle) and energy (triangle) profiles for upgrading of Athabasca bitumen (solid = control; empty = catalytic). $t_i$ is the time that the temperature inside the autoclave reached 300 °C.](image)

This result is further reflected in Figure 3, which presents the pressure variation during upgrading runs. In general, and especially so for the 440 °C runs, the drill cuttings increased pressure evolution during upgrading, mostly in the form of condensable gases and volatiles which condensed to the liquid upon cooling to ambient conditions. The energetic requirement for the catalytic and non-catalytic thermal cracking fell in the range 5000–6500 MJ per barrel of feed. Pressure evolution was significant at higher temperatures, with pressures reaching upwards of 1000–1200 psi as a result of cracking. The difference between low and high temperature runs again stems from the fact that cracking reactions are much more favored at more severe temperatures [49–51].
viscosity is significantly reduced from the feed for both runs, 1100 ± 230 cSt and 720 ± 100 cSt, respectively, it is well over the 350 cSt requirement for pipelines. Additionally, °API gravity improvement is only 1–2 points from the feed for both control and catalytic runs. Thus, while the yields are encouraging, either the residence time or the severity of the reaction must be increased to further reduce the viscosity. A higher residence time could be employed, however, for the sake of partial upgrading, where the main objective is to obtain transportable crude, residence times should ideally be kept short in order to minimize reactor volumes.

At 420 °C the toluene insoluble (TI) yield becomes relatively significant with 4.2 ± 0.8 wt.% and 5.3 ± 0.7 wt.% coke being formed as a result of non-catalytic and catalytic thermal cracking, respectively. Likewise, the gas yield slightly increased in comparison to the runs at 400 °C. This is also reflected in Figure 3, where the pressure evolution was much more significant at 420 °C compared with 400 °C. Therefore, lower liquid yield, 87.9 ± 4.6 wt.% and 85.7 ± 5.9 wt.%, was attained for both the control and the catalytic runs, respectively. It is worth noting that the volatile yield became significant at this temperature. On the other hand, the quality of liquid product improved by a large margin. The viscosity and °API gravity was 64 ± 22 cSt and 22.8 ± 3.1 for the control runs, and 42.3 ± 5.7 cSt and 24.9 ± 1.6 for the catalytic runs. This product quality is well under the 350 cSt and 940 kg m⁻³ viscosity and density requirement for pipeline transport. Thus, only ~5 wt.% of the liquid product was sacrificed in the form of coke to further push the quality of the liquid product to an acceptable standard.

2.3. Product Yield and Quality

Figure 4 presents the yield of the different products obtained from the catalytic and non-catalytic cracking of Athabasca bitumen, and Figure 5 presents quality measurements of the centrifuged liquid product at 400 °C, 420 °C and 440 °C. At 400 °C, almost no coke is formed for either control or catalytic runs resulting in 95.0 ± 4.7 wt.% and 93.5 ± 5.5 wt.% liquid product, respectively. However, the volatile yield became significant at this temperature. On the other hand, the quality of liquid product improved by a large margin. The viscosity and °API gravity was 64 ± 22 cSt and 22.8 ± 3.1 for the control runs, and 42.3 ± 5.7 cSt and 24.9 ± 1.6 for the catalytic runs. This product quality is well under the 350 cSt and 940 kg m⁻³ viscosity and density requirement for pipeline transport. Thus, only ~5 wt.% of the liquid product was sacrificed in the form of coke to further push the quality of the liquid product to an acceptable standard.

Figure 3. Pressure profiles for upgrading of Athabasca bitumen (solid = control; empty = catalytic). $t_i$ is the time that the temperature inside the autoclave reached 300 °C.
The liquid yield further degraded to 75.3 ± 5.7 wt.% for the control runs and 65.5 ± 3.2 wt.% for the catalytic runs when the severity of the reaction was pushed to 440 °C. The TI and gas yields increased to 13.1 ± 0.8 wt.% and 11.6 ± 1.4 wt.% for the control runs, and 16.6 ± 1.3 wt.% and 17.8 ± 1.5 wt.% for the catalytic runs. On the other hand, the viscosity and °API gravity of the liquid further changed to 16.3 ± 3.4 cSt and 23.5 ± 3.6 for the control runs, and 12 ± 3.6 cSt and 22.2 ± 2.6 for the catalytic runs. It is concluded that this further reduction in viscosity is not worth the substantially increased TI and reduced liquid yield, as the viscosity and density pipeline requirements were already met through upgrading at 420 °C. It should be noted that the depreciation of the °API gravity of the liquid product relative to the 420 °C runs could result from entrained coke into the liquid product at the high coke yield encountered in these runs.
Based on the conditions investigated, it would be recommended to run the process at 420 °C as it resulted in the best balance of yields and liquid quality. The upgrading could potentially be run with a shorter residence time, while still achieving acceptable viscosity and density. It is well-known that coke forms following an induction time [52,53]. Therefore, significant coke formation could be avoided if the residence time was limited, however the liquid quality must be monitored carefully. It is interesting to note that throughout the conditions, catalytic upgrading resulted in lower liquid yield than the control runs in exchange for slightly higher liquid quality. At 440 °C the difference in liquid yields between the control and the catalytic runs was upwards of 10 wt.% in return for not very significant quality improvement. This is most likely due to the increased yield of gaseous and volatile products, which reduced the liquid yield slightly, but in terms of the volatiles, contributed to increased liquid quality. On the other hand, at 400 °C the liquid yield difference was only 1 wt.%, yet the viscosity difference was ~400 cSt. Clearly, a balance must be struck at the chosen operating conditions. Even though the drill cuttings are a waste material, it is entirely possible that the proposed process could be sufficiently run under thermal cracking, control conditions, if proved significantly advantageous.

Table 1 presents the Conradson carbon residue (CCR) and the sulfur content of the product oil of the non-catalytic and catalytic thermal cracking of Athabasca bitumen. At 400 °C, CCR increased from the feed for both the control and the catalytic runs, however much less so for the catalytic runs. The increase in CCR most likely results from condensation and dehydrogenation reactions, and therefore formation of aromatic species, especially since no coke formed at 400 °C, as a result of thermal cracking. In comparison, for the catalytic runs, lower CCR yield suggests that the drill cuttings promoted formation of reactions that led to paraffins and olefins, rather than aromatics, perhaps through the aforementioned dispersion effect and the abundance of LAS. At 420 °C, the CCR yields were 11.6 wt.% and 10.4 wt.% for the control and catalytic runs, respectively, both showing a decrease from the feed. This is reflected by the apparent TI yield, as seen in Figure 4.

**Table 1.** Conradson carbon residue (CCR) and total sulfur content of upgraded bitumen samples.

| Feed Temperature | Control CCR (wt.%) | Catalytic CCR (wt.%) | Control Sulfur (wt.%) | Catalytic Sulfur (wt.%) |
|------------------|--------------------|----------------------|-----------------------|------------------------|
| 400 °C control   | 20.9               | 14.2                 | -                     | -                      |
| 400 °C catalytic | 11.6               | 10.4                 | 4.1                   | 4.2                    |
| 440 °C control   | 14.8               | -                    | -                     | -                      |
| 440 °C catalytic | 8.4                | -                    | -                     | -                      |

The majority of the aromatic species formed coalesced during upgrading and ended up in the coke fraction, resulting in a net decrease compared to the feed. Under carbon rejection conditions, ~25% of the sulfur in the feed was removed from the liquid through either formation of gaseous species such as H2S or ejection into the coke fraction. The drill cuttings did not improve sulfur reduction in comparison to the control runs, which suggests that the catalyst lacked affinity to the C–S bond. To further test the stability of the liquid product of the catalytic and the non-catalytic thermal cracking at 420 °C, viscosity and °API gravity was re-measured after a period of 3 weeks, as shown in Figure 6. It is clear that the quality of the liquid has experienced some degradation possibly due to olefin instability and polymerization, but to no more than 10% deterioration in the viscosity and the °API gravity. The resulting liquid product is still in the acceptable range for pipeline transport (viscosity < 350 cSt, °API gravity > 19). It should be noted that a three-week timeframe is much more than expected to transport oil through pipelines [18].
2.4. Kinetic Modelling

The results of the predictive capability for catalytic and non-catalytic thermal cracking of Athabasca bitumen at 400 °C and 420 °C for 1 h are presented in Figure 7. Fitted parameters are summarized in Table 2. The model predicted the overall liquid, coke and gas yields at both conditions to an excellent extent. For example, at 400 °C, a 95.8 wt.% liquid yield was calculated through the model and 95.5 ± 2.9 wt.% was experimentally obtained. Similarly, at 420 °C, a liquid yield of 87.6 wt.% was calculated through the model versus 85.7 ± 5.9 wt.% experimentally measured. The discrepancy between the model prediction and the experiments is only limited to predicting the asphaltene and the volatile yields for the non-catalytic experiment. At 400 °C the model overpredicted an asphaltene yield of 14.1 wt.% compared with the experimental yield of 5.5 ± 2.0 wt.%, while underpredicted the distillate yield of 3.3 wt.% compared with the experimental yield of 9.9 ± 0.6 wt.%. This could be due to a difference in asphaltenes molecular structure between Athabasca bitumen and Arabian atmospheric residue, which in principle should contribute different products from their cracking [54–56]. Moreover, the model equations do not account for direct conversion of asphaltenes to volatiles, since this reaction was deemed relatively slow when fitting Arabian atmospheric residue data. On the other hand, the same observation was not reported for the catalytic cracking run, and except for predicting slightly less asphaltene conversion to gas and coke, the model captured the catalytic conversion of Athabasca bitumen to the different cuts much more accurately. The difference between the model prediction of the catalytic experimental results can still be attributed to the difference in the molecular structure of asphaltenes between the Arabian atmospheric residue and the Athabasca bitumen, whereas the enhanced predictability relative to the non-catalytic run are attribute the effective role of the catalyst at quenching the radicals and exposing the adsorbed asphaltenes to hydrogen donors from the solvent [57]. These reactions contribute to light and medium hydrocarbon distillate components, per our previous findings [57].
2.5. Energy Analysis

To fully comprehend the efficacy of the onsite upgrading process proposed here compared with diluent-based transport, energetic analysis was carried out. Sample calculations as well as a summary of the energy analysis can be found in the supporting material and Table S1. Our calculations show that the energy needed for pumping partially upgraded bitumen is approximately 30% of the energy needed to pump the diluted bitumen. It should be noted that pumping the diluent back to the producer only accounts for ~3% of the total energy needed to pump the diluted bitumen, given its low viscosity and density. If one takes into account the energy needed to evaporate the diluent, however, the energy requirement for pumping partially upgraded bitumen falls down to only 0.02%. However, our measurements and calculations showed that the energy needed for pumping 1 bbl of bitumen, without heating, might be three orders of magnitude lower than the energy required for its upgrading.
In this context, refinery upgrading might be more advantageous since it provides the option of only subjecting the vacuum residue stream to the upgrading process. Onsite upgrading, on the other hand, and in order to minimize the number of required units on-site, subjects the whole bitumen stream to thermal cracking. Assuming the same value of energy consumption between refinery and onsite upgrading, pumping diluted bitumen with refinery upgrading option consume ~45% of the energy needed for the onsite upgrading of bitumen. It should be noted, however, that a high temperature and pressure hydrotreating process in a refinery may require much more energy than the simple thermal cracking process proposed in this work, especially when methane reforming step to supply the required hydrogen is taken into consideration. If, on the other hand, the diluted bitumen is pumped to an upgrader, where the whole bitumen stream is subjected to upgrading, then onsite upgrading would still make more energetic sense with energy savings of ~30 MJ/bbl.

It is worthwhile noting that the above analysis is made purely from an energetic standpoint. Economic consideration may prove that onsite upgrading of bitumen is probably much more profitable, especially given the specification/quality of the end product relative to the raw bitumen. Assuming same performance as the autoclave of this study, the volume of a continuous flow mixed tank reactor needed to upgrade 100 bbl of bitumen product per day on-site would only be 1 m$^3$. The energy needed for onsite upgrading may be supplied from the gas produced from the same well or a nearby gas well, as well as supplemented through the gas produced during upgrading.

3. Materials and Methods

3.1. Materials

Toluene (BDH 99.8%, VWR, Canada) and n-heptane (BDH technical, VWR, Canada) were used for separating toluene insolubles (coke, catalyst, etc.) and asphaltenes from the product oil. Thermal and catalytic cracking runs were carried out under a N$_2$ (99% purity, Praxair Specialty Gas and Equipment, Calgary, AB, Canada) atmosphere. Athabasca bitumen (Cenovus, Calgary, AB, Canada) was used as the feedstock to the upgrading experiments. Properties of the feedstock are presented in Table 3.

| Property          | Value          |
|-------------------|----------------|
| °API gravity      | 13.1 ± 2.2     |
| Viscosity (cSt)   | >200,000       |
| Asphaltenes (wt.%)| 15.4 ± 1.2     |
| Maltenes (wt.%)   | 82.2 ± 0.3     |
| Sulfur (wt.%)     | 5.4            |
| CCR (wt.%)        | 12.9           |

Dry sandstone drill cuttings were received from Executive Mat Service (EMS, Vancouver, BC, Canada). The as-received drill cuttings were purified by thoroughly washing with water followed by drying in an oven at 40 °C for 12 h. To reduce the drill cuttings to an acceptable size, they were first ground using a mortar and pestle and then sieved to filter out >355 µm cuttings. The particle size distribution of the resultant drill cuttings was determined using a tower of sieves with various mesh size. Scanning electron microscopy (SEM) (Quanta FEG 250, FEI Inc., Canada) and energy-dispersive X-ray spectroscopy (EDX) (Quantax 5030, Bruker, Montreal, QC, Canada) were performed to investigate the morphology and surface elemental analysis of the drill cuttings, respectively. N$_2$ adsorption/desorption experiments were carried out at −196 °C in order to evaluate the surface area of the drill cuttings. First, the drill cuttings were degassed at 150 °C under a N$_2$ atmosphere for 24 h, and then fed to a Micromeritics surface area analyzer (TriStar 2000, Micromeritics Instrument Corporation, Norcross, GA, USA). Surface area measurements were calculated using the Brunauer–Emmet–Teller (BET) method and total pore volume was estimated using the Barrett–Joyner–Halenda (BJH) method at a relative pressure (P/P$_0$) of 0.99. Fourier-transform infrared spectroscopy (FT-IR) analysis was carried out on an Agilent Cary 630 (Agilent, Santa Clara, CA, USA) to determine Lewis and Brønsted acid site strength.
and concentration. As a control sample, the dry drill cuttings were introduced into the instrument and analyzed. Then, a drop of pyridine was introduced onto the drill cuttings and the sample was re-introduced into the instrument. Thus, both dry and pyridine drill cuttings spectra were collected and compared in order to determine the presence of Lewis and Brønsted acid sites.

3.2. Upgrading

The bitumen feed was first prepared by heating in a gravity convection oven (DX300, Yamato Scientific America Inc., Santa Clara, CA, USA) at 90 °C until it became pourable. The source sample was then mixed well with a metal rod to ensure homogeneity, and then a 50 g and 45 g samples were taken for the control and catalytic runs, respectively. A mass of 5 g drill cuttings was added to 45 g of oil for the catalytic runs to achieve a 10 wt.% (100,000 ppm) concentration. This concentration was previously found to be suitable for catalytic cracking of Athabasca vacuum residue [29,30]. Upgrading runs were performed in a 100 mL Parr autoclave (1.3 in. i.d. and 4.6 in. length, 4590 Micro Bench Top Reactor, Parr Instrument Company, Moline, IL, USA). When placed in the main apparatus, the reactor was surrounded by a heating jacket, which was electrically connected to a digital control unit which could monitor the temperature and pressure. A DI-50E meter (Electro-Meters Company Ltd., Pickering, ON, Canada) was placed in series with the apparatus for monitoring heater usage. Once secured in the apparatus, the reactor was purged with N₂ for 3 min to introduce a N₂ atmosphere. Then, a pressure leak check was performed by holding the pressure within the autoclave at 100 psi for 5 min, which upon completion, was released to achieve an initial atmospheric pressure in the reactor. The inlet and outlet were sealed tightly, and heating commenced at a rate of 25 °C min⁻¹. Upon reaching a temperature of 120 °C inside the autoclave, mixing at a rate of 500 rpm commenced. Our previous work showed that a lack of mixing resulted in additional formation of TI during upgrading, as well as improper mass and heat transfer caused by settled drill cuttings. The start point of the cracking reaction was considered from 300 °C, as cracking is well known to commence in the range of 300–350 °C [58]. After reaching this point, the upgrading proceeded for 1 h. Upon completion, the autoclave was quickly quenched several times using a cold-water bath, and then left to cool to ambient conditions (20 °C). The final pressure of the reactor was then recorded and any final, uncondensed, gases were slowly vented out of the system. The reactor was then removed from the main holding apparatus, liquid was scraped off the mixer shaft, and weighed. The overall gas yield of the reaction was taken to be the missing mass found by comparing the initial and final weights, as a result of the upgrading.

3.3. Analysis

After weighing, a liquid product sample was taken for characterization. A volume of 30 mL of the liquid product was poured into a centrifuge tube and centrifuged for 30 min at 5000 rpm. The supernatant was collected, and the bottoms of the tubes were weighed and added to the rest of the liquid still present in the reactor. Thus, two separate fractions, the supernatant and bottom fractions, were created out of the liquid product. The viscosity of the supernatant was measured using a cylindrical Brookfield viscometer (Model: LVDV-1 PRIME, Brookfield Engineering Laboratories Inc., Middleborough, MA, USA), and °API gravity using a specific gravity bottle (Thomas Scientific, Swedesboro, NJ, USA) at 16 °C. °API gravity was calculated as follows:

\[ °\text{API gravity} = 141.5/((m_{\text{oil}} - m_{\text{pycnoc}})/((m_{\text{water}} - m_{\text{pycno}}) - (m_{\text{sol}} - m_{\text{sol}}))) - 131.5 \]  

(1)

The variables are measured as follows: \( m_{\text{oil}} \) – pycnometer + few drops of oil, \( m_{\text{pycno}} \) – empty pycnometer, \( m_{\text{water}} \) – pycnometer + water, \( m_{\text{sol}} \) – pycnometer + few drops of oil + water. To ensure accuracy of results, the viscosity measurement was carried out at 5 different mixing speeds, and °API gravity was repeated 5 times. The supernatant was then washed with heptane at a 1:40 v/v ratio and filtered using 25 µm VWR filter paper. The filter paper was allowed to dry, and then placed into a double layer cellulose VWR thimble. The thimble was then introduced into a Soxhlet and washed with
heptane for 24 h. After washing, the thimble was dried and the mass of the extracted asphaltenes was determined by weighing the thimble and the filter paper. The heptane/maltene filtrate was introduced into a rotary evaporator (model: Hei-Vap value digital HL/G3, Heidolph Instruments GmbH & Co., KG, Schwabach, Germany) operating at a temperature of 93 °C and pressure of 2 psi, where the heptane was evaporated out of the solution. After the final drop of condensate appeared, the evaporation was allowed to continue for 5 min, in order to dry the leftover maltene fraction. The difference between the mass before and after was taken to be as the volatile yield of the supernatant fraction of the overall liquid product. The leftover maltene fraction was weighed and collected.

The bottoms fraction of the liquid product was thoroughly washed with toluene at a 1:40 v/v ratio and filtered using 25 µm VWR filter paper, ejecting the TI out of the liquid. The filter papers were allowed to dry in ambient conditions for 24 h, weighed, and then the dried TI solids collected. The toluene was then evaporated out of the system using a rotary evaporator operating at 93 °C and 2 psi. Losses in the form of volatiles were considered by comparing weights of the feed to the rotary evaporator before and after evaporation. After toluene evaporation, similar to the supernatant, the asphaltenes were separated out using heptane at a 1:40 v/v ratio, and 25 µm VWR filter paper. As before, the dried filter paper was wrapped into a thimble and introduced into a Soxhlet and thoroughly washed with heptane for 24 h. Finally, as with the supernatant, the heptane was evaporated, any remaining volatiles accounted for, and the remaining maltene fraction weighed and collected. To obtain 95% confidence intervals, three replicates were carried out for every upgrading run. The total sulfur concentration of the feedstock and the liquid product was measured using a Trace Sulfur Analyzer Model TS-100 (Mitsubishi Chemical Analytech Co., LTD, Shimotsuruma, Japan). Conradson carbon residue (CCR) content of similar oil samples was measured using thermogravimetric analysis (TGA), following a procedure outlined elsewhere [59].

3.4. Kinetic Model

A 5-lump, 6-reaction, kinetic model developed earlier by our group to describe the thermal catalytic and non-catalytic cracking of Arabian atmospheric residue was employed to predict the thermal cracking of bitumen [57]. Figure 8 shows the model reactions between the major lumps. The model parameters were optimized for thermal cracking of Arabian atmospheric residue and were directly employed here without data fitting [29,57].

According to Figure 8, the reaction network is governed by the following equations:

\[ \text{coke} \rightarrow \frac{dx_j}{dt} = k_1 x_2 u(t - t_{\text{induction}}) \]  

Figure 8. Five-lump, 6-reaction, thermal cracking kinetic model.
asphaltene → dx₂/dt = -(k₁ + k₂ + k₃)x₂ + k₄x₄ (3)
gas → dx₃/dt = -k₂x₂ + k₅x₃ (4)
maltene → dx₄/dt = -(k₄ + k₅ + k₆)x₄ + k₃x₂ (5)
distillate → dx₅/dt = k₆x₄ (6)

A step function was introduced into (2) in order to account for the induction period of coke formation. A value of 30 min was used for the induction time in this work, based on previous observations, albeit various induction times are reported for different feeds in literature [60–62]. The fitted parameters were optimized by reducing root mean squared deviations (RMSE) between the model and the experimental Arabian atmospheric residue data using a Monte Carlo type procedure [57].

4. Conclusions

Thermal cracking of Athabasca bitumen was carried out in a batch reactor for a range of temperature between 400 °C and 440 °C and 1 h duration. At 400 °C, for both catalytic and non-catalytic runs, minimal amounts of coke were produced, however the viscosity of the product oil was >350 cSt, and therefore insufficient for pipeline standards. At 420 °C for both catalytic and non-catalytic runs, a good balance between coke yield and liquid yield and quality is found. Only ~5 wt.% coke formed in return for a viscosity of ~50 cSt and °API gravity of >25. A sulfur and CCR reduction of 25% and 10%, respectively, were achieved from the feed at 420 °C. The liquid product was shown to be stable with only 10% depreciation in viscosity and °API gravity over a period of 3 weeks, with the liquid still remaining well within acceptable pipeline standards. At 440 °C, the non-catalytic run produced ~75 wt.% liquid, and catalytic run produced ~65 wt.% liquid with only slight improvement in quality in comparison with the runs at 420 °C. The required thermal energy input into the process was measured to fall between 5500 and 5900 MJ per barrel of feed, accounting for potential heat losses. Overall energetic analysis showed that the pumping requirement for the partially upgraded product was 3 times lower than that for diluted bitumen, without accounting for diluent recovery through evaporation.

Supplementary Materials: The following are available online at http://www.mdpi.com/2073-4344/9/5/431/s1, Table S1: Energetic comparison between diluted bitumen and partial upgrading pathways.

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Abbreviations

The following abbreviations are used in the article:

BAS Brønsted acid sites
CCR Conradson carbon residue
LAS Lewis acid sites
TI Toluene insoluble

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