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Effect of ethane and ethylene on catalytic non oxidative coupling of methane†

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The effect of addition of ethane and ethylene (C₂) on methane coupling at 1000 °C was investigated. A Fe/SiO₂ catalyst was used to determine the contributions of catalytic as well as C₂ initiated methane activation. The catalyst load as well as the residence times at 1000 °C downstream of the catalyst bed were varied. C₂ addition significantly increases methane conversion rates, similarly for both ethane and ethylene, although ethylene is more effective when operating with long residence times in the post-catalytic volume. Methane activation *via* C₂ addition proceeds dominantly in the gas-phase whereas catalytic C₂ activation is negligible. The catalyst has no effect on methane conversion when the feed contains more than 2 vol% C₂. Product selectivity distribution as well as total hydrocarbon yield at 10% conversion is not influenced by C₂ addition, but is influenced by the amount of catalyst as well as residence time in the post-catalytic volume at high temperature. It is proposed that C₂ impurities in natural gas change from a nuisance to an advantage by enhancing methane conversion and simplifying purification of the natural gas feed. A process is proposed in which ethylene is recycled back into the reactor to initiate methane coupling, leading to a process converting methane to aromatics.

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Introduction

Natural gas, consisting typically of 90 vol% CH₄ and 5 vol% C₂ hydrocarbons,¹ is considered as a highly interesting resource for the production of olefins and aromatics.^{2–4} Traditionally, these products are by-products of crude oil refining.^{5,6} It is, however, predicted that raw oil extraction will reach peak production in the coming decades,⁷ and hence the increase of global olefins and aromatics demand needs to be met *via* another process. Natural gas is already used on large industrial scale for the synthesis of base chemicals and fuels.^{8–10} These processes convert natural gas to specific hydrocarbons with high carbon efficiencies to olefins and aromatics over crude oil refining.^{11,12}

Current industrial methods for converting natural gas into base chemicals are multistep processes starting from methane steam-reforming to obtain syngas.¹³ The syngas can be converted into paraffins in the Fischer–Tropsch (FT) process⁸ or used for methanol synthesis, followed by methanol to gasoline (MTG)¹⁴ or methanol to olefins (MTO).¹² The large number of process steps at different

temperatures and pressures makes the processes energy intensive and only viable at large installed capacities.¹⁵

Direct conversion of methane to higher hydrocarbons is hence receiving a growing interest over the last decades, as a more efficient alternative to the indirect routes mentioned before. Three main research directions can be distinguished, *i.e.* methane dehydro-aromatization (MDA),¹⁶ oxidative coupling of methane (OCM)¹⁷ and non-oxidative coupling of methane (NOCM) at high temperature.¹⁸ Both MDA as well as OCM suffer from low single pass conversion as well as low product yields.^{16,17} Guo *et al.*¹⁹ reported in 2014 that a Fe/SiO₂ catalyst is able to couple methane non-oxidatively to olefins and aromatics at high conversion levels and without coke formation. Measurements were carried out at temperatures in excess of 950 °C. Follow-up research showed that methane conversion can be increased by *in situ* hydrogen removal,²⁰ by using a catalytic wall reactor²¹ and by increasing the residence time in the reactor at high temperature downstream the catalyst bed (post-catalytic volume).^{22,23} However, all these studies reported coke formation and lower catalytic activity^{19–25} compared to the original work of Guo *et al.*¹⁹ It is generally accepted that the catalyst initiates methane conversion, through formation of methyl radical, followed by free radical chain reactions and coupling reactions in gas phase, determining the product distribution.^{19,22,25} The publications concerning the Fe/SiO₂ catalyst have focused on a pure methane feed, even though natural gas contains a significant fraction of C₂ hydrocarbons.¹

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Ethane and ethylene readily form radicals that participate in the autocatalytic cycle of methane pyrolysis at reaction temperature, *i.e.* above 950 °C.^{26–29} Addition of ethane can significantly reduce the induction period during non-catalytic methane pyrolysis as reported in early work by Germain *et al.*³⁰ Methane conversion rates can thus be significantly increased by introduction of small amounts of C₂ hydrocarbons, up to 3%, into the reactant mixture as reported in early work by Schneider²⁹ and Rokstad *et al.*³¹ Ogihara *et al.*³² reported recently that methane is activated by ethane addition even at relatively low temperatures 700–800 °C in absence of any catalyst. Guo *et al.*¹⁹ reported a significant increase in methane conversion upon addition of 1–5% C₂H₆ at 900 °C using the Fe/SiO₂ catalyst, although ethane addition caused coke formation, which crucially was not observed in their experiments using pure methane. SABIC²³ patented a concept where post-catalytic ethane injection would quench the free radical coupling reaction, stabilizing the formed olefin products. The effect of addition of C₂ hydrocarbons in presence or absence of the Fe/SiO₂ catalyst on methane conversion and especially product distribution has not been reported, to the best of our knowledge.

This study reports for the first time on the interaction between catalytic methane activation and activation *via* the addition of free-radical initiators, *i.e.* ethane and methane conversion, product selectivity distribution and deposit formation.

Experimental

Catalyst synthesis

The Fe/SiO₂ catalyst is synthesized according to the method described in ref. 19 and details concerning the catalyst synthesis as well as characterization can be found in ref. 22. In short, inhouse synthesized Fe₂SiO₄ is mixed with quartz and ball-milled in N₂ atmosphere overnight using a zirconia milling jar. The resulting powder is fused for 6 h at 1700 °C in air. The resulting slab is crushed and sieved, the fraction between 250–500 μm is used in the experiments. The sieved catalyst particles are leached for 2 h in 0.5 M HNO₃, rinsed and dried to obtain the final catalyst.

Reactor setup

A modular three-zone oven is used for catalytic testing. Each zone is thermally insulated from the others and from the environment, allowing for steep temperature gradients, as presented in Fig. 1. The pre-heater is always operated at 400 °C and the reactor-zone, containing the catalyst, at 1000 °C. The post-heater is either operated at 400 °C, cooling down the gas-stream directly after the reactor-zone, or at 1000 °C to create an extended residence time at higher temperature, increasing the conversion as reported in our previous work.²² Both temperature profiles, for operating the post-heater at 400 °C or 1000 °C have been included in Fig. 1. Gas flowrates are controlled using digital mass flow controllers. Product gasses are analyzed using a three-channel Varian CP-3800 in-

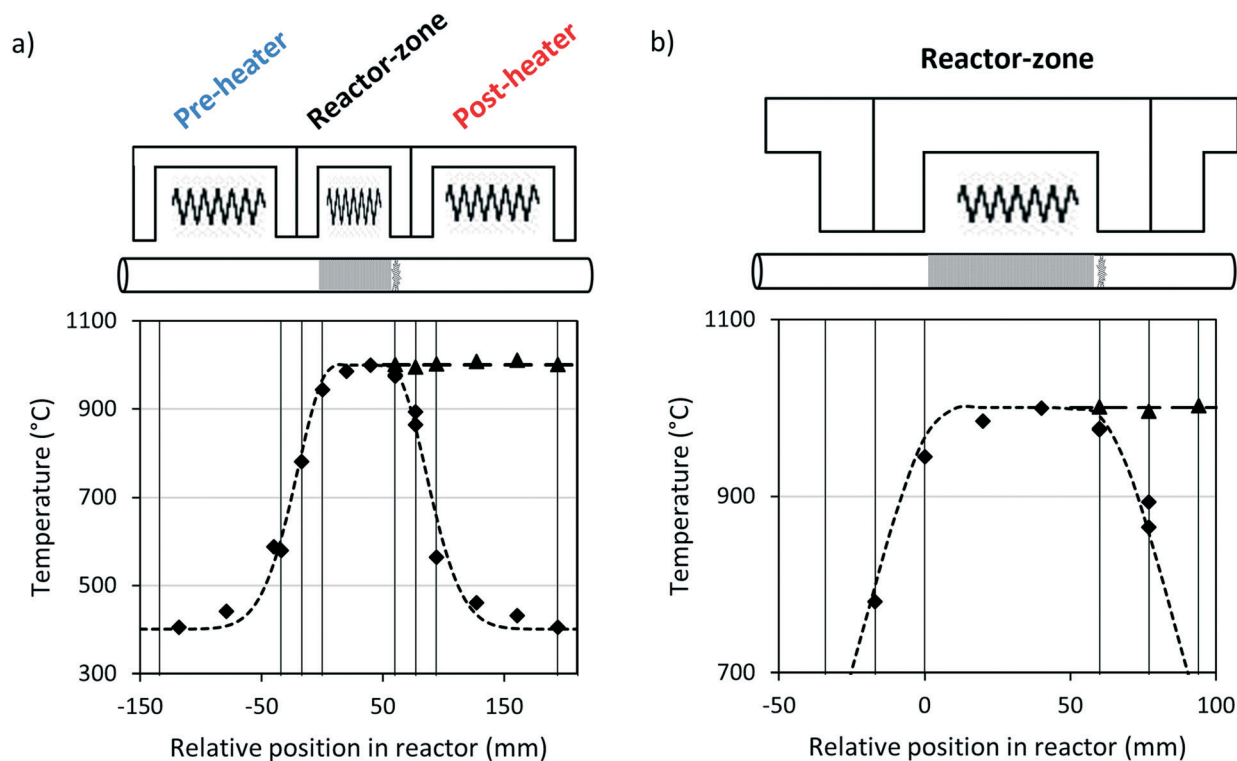


Fig. 1 Temperature profile inside the reactor measured with an empty reactor tube b) zoom in on Fig. 1a; gas-flow rates of 10 ml min⁻¹ N₂; vertical bars represent the insulating layers between the 3 different zones. ---◆--- Pre- and post-heater at 400 °C; reactor-zone at 1000 °C; ---▲--- post-heater at 1000 °C.²²



line gas chromatograph, the tubing between the reactor and the GC is heated to 200 °C to minimize hydrocarbon condensation. Further details concerning the reactor setup are presented in ref. 22.

Experimental procedure

The experimental procedure is adapted from ref. 22. The catalyst is placed at the desired position in the quartz reactor according to Fig. 2, held in position by a small quartz wool plug. The reactor is flushed for 10 minutes with 10 ml min⁻¹ N₂. The catalyst is heated with 20 °C min⁻¹ to 900 °C under 10 ml min⁻¹ N₂. The catalyst is then activated in 90 vol% CH₄ in N₂ at 1000 ml g_{cat}⁻¹ h⁻¹ during two hours, following the procedure of Bao c.s.³³ After activation, the reactor is flushed for 10 minutes at 900 °C with N₂ 10 ml min⁻¹. The temperature is increased to the desired reaction temperature, with a heating rate of 20 °C min⁻¹ and the experiment is started by feeding 90 vol% CH₄ balanced with N₂. Ethane or ethylene are added while reducing the amount of nitrogen in the feed, keeping the methane concentration as well as the total flowrate constant. Catalyst,¹⁹ gas-phase²⁷ and reactor-wall³⁴ all contribute to methane conversion so that space velocity is not useful as a descriptor; therefore the total gas flowrate was kept constant at 16.6 ml min⁻¹ to allow for a fair comparison. The product gas is analyzed using an online three channel gas chromatograph (Varian CP-3800), measuring CH₄ and N₂ on the first channel, H₂ on the second channel and all hydrocarbon products on a third FID channel. At the end of an experiment, the reactor is flushed for 10 minutes with N₂ at 10 ml min⁻¹ after which the reactor ovens are turned off and left to cool, maintaining the N₂ flow while cooling. The catalyst is removed from the reactor and analyzed for coke deposition using TGA (Mettler Toledo TGA/DSC 3+ Star System). Details on the temperature-programmed-oxidation measurement in the TGA can be

found in ref. 22. The online GC samples every 27 minutes and all data points presented are based on at least 3 measurements in steady state conditions.

Methane conversion is calculated according:

$$\zeta_{\text{CH}_4} = 1 - \frac{P_{\text{CH}_4\text{out}} \cdot P_{\text{N}_2\text{in}}}{P_{\text{CH}_4\text{in}} \cdot P_{\text{N}_2\text{out}}} \quad (1)$$

ζ_{CH_4} : conversion of methane (-).

P_X : partial pressure of compound X (bar).

Methane conversion is corrected for any change in the molar flow rate based on change in the nitrogen tracer concentration according eqn (1).^{19,35} In the same way, the conversion of the C₂ reactant is calculated, according eqn (2). Note that C₂ conversion as calculated is determined by both C₂ fed to the reactor and formation of the same C₂ species in the reactor, therefore this value can also become negative when formation dominates over conversion.

$$\zeta_{\text{C}_2} = 1 - \frac{P_{\text{C}_2\text{out}} \cdot P_{\text{N}_2\text{in}}}{P_{\text{C}_2\text{in}} \cdot P_{\text{N}_2\text{out}}} \quad (2)$$

ζ_{C_2} : overall conversion of the added C₂ hydrocarbon (-).

Product selectivity is calculated on carbon base, corrected for any change in the molar flow rates based on the concentration of the N₂ tracer, eqn (3).^{19,35} The selectivity calculation takes into account conversion of both methane and C₂, adjusting for stoichiometry.

$$S_{\text{C}_x\text{H}_y} = \frac{x \cdot P_{\text{C}_x\text{H}_y} \cdot P_{\text{N}_2\text{in}}}{P_{\text{N}_2\text{out}} \cdot (P_{\text{CH}_4\text{in}} + 2 \cdot P_{\text{C}_2\text{in}}) - P_{\text{N}_2\text{in}} \cdot (P_{\text{CH}_4\text{out}})} \quad (3)$$

$S_{\text{C}_x\text{H}_y}$: selectivity towards C_xH_y hydrocarbon (-).

The measured hydrogen signal was used to validate the calculated conversion and selectivity distribution and closes to within 5%.

Experiments were performed over a maximum period of 8 h, a stability test over 16 h showed a deactivation of 10% methane conversion in a test using pure methane at 1.8% conversion.

Results

Effects on methane and C₂ conversion

Fig. 3 shows the effect of adding small amounts of ethane and ethylene to the reactant mixture, varying the amount of catalyst and post-catalytic volume. Addition of ethane and ethylene significantly increases CH₄ conversion. Interestingly, there is no discernible difference between the methane conversion when adding ethane (Fig. 3a and b) or ethylene (Fig. 3c). The effect of C₂ addition on methane conversion is decreases with increasing catalyst amount, to the extent that catalyst has no effect when at least 2% ethane is added. (Fig. 3b) Fig. 3d shows a linear correlation, including the origin, between the free-volume in the reactor-zone and the linear acceleration in methane conversion rate by addition of C₂ molecules, as calculated based on Fig. 3a and c.

Fig. 4 shows the overall ethane and ethylene conversion in all experiments. Ethane is close to completely converted in

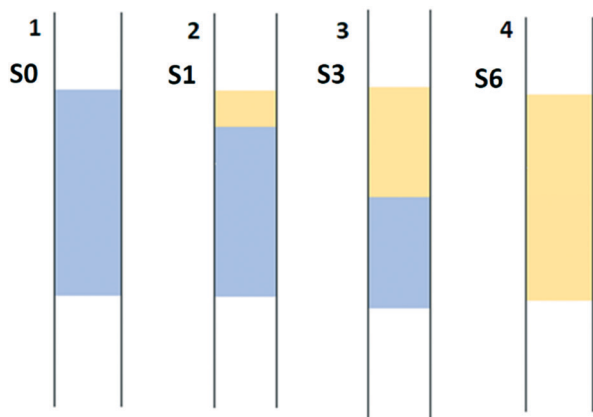


Fig. 2 3 ways to position the catalyst inside the 6 cm zone of oven segment 2. Situations 1 (S0) contains no catalyst, 2 (S1) contains a 1 cm high bed (160 mg), situation 3 (S3) contains a catalyst bed 3 cm height (420 mg), situation 4 (S6) a catalyst bed height of 6 cm (820 mg), leaving no free volume at reaction temperature.



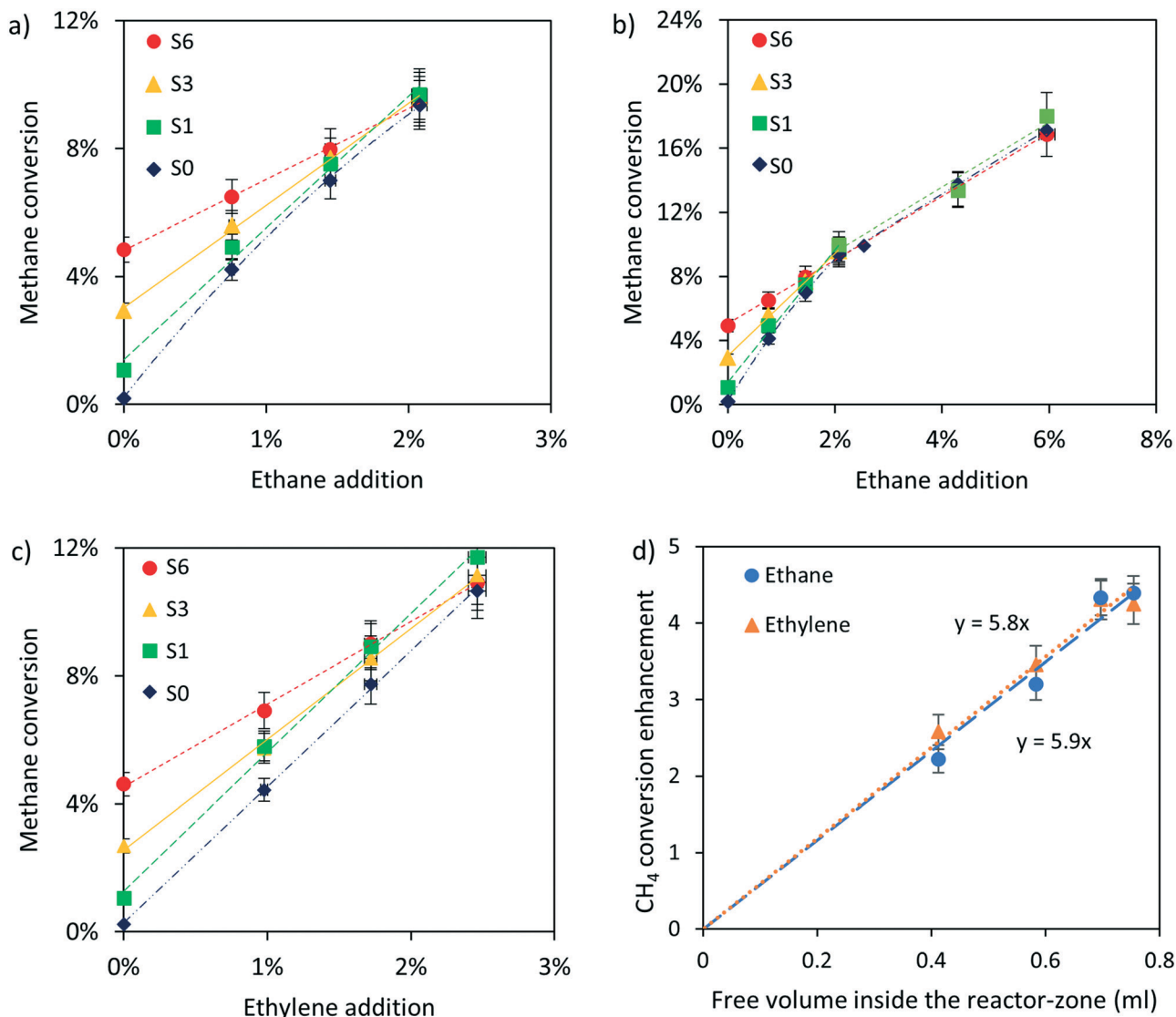


Fig. 3 Effect of C₂ addition on methane conversion, varying the amount of catalyst inside the reactor-zone: (a) ethane addition at lower concentration and b) higher ethane concentration; (c) ethylene addition; (d) the effect of free volume inside the reactor on the slope of increase in methane conversion when increasing C₂ addition, as shown figures (a) and (c) (free volume is estimated based on the free volume surrounding the catalyst particles as well as the free volume downstream of the catalyst bed at packing density of 0.45); conditions: 90% CH₄ N₂ balance; 16.6 ml min⁻¹ total flowrate. Reactor-zone at 1000 °C, pre-heater and post-heater at 400 °C.

all cases although conversion is somewhat lower when employing a large catalyst bed. Ethylene conversion on the other hand is always lower than 50%. Interestingly, ethylene production dominates over conversion when using the post-heater and when using a large catalyst bed (S6), *i.e.* conditions resulting in high methane conversion. Note that the C₂ conversions presented in Fig. 4 are the result of C₂ conversion and C₂ production in the reactor.

The post catalytic residence time at high temperature can also be increased by using the post-heater as shown in Fig. 1. Fig. 5 shows a significant increase in methane conversion, keeping the level of ethane addition constant, when operating the post-heater at the same temperature as the reactor, *i.e.* 1000 °C. Fig. 5 also shows that ethylene addition results in the highest methane conversion when compared

with ethane addition at the same concentration. Note that the difference in post-catalytic free volume between the S1 case with or without post-heater is a factor of 3.7, significantly larger than the increase in conversion observed in Fig. 5, showing a diminishing return of C₂ addition at longer residence time.

Fig. 6 shows the selectivity towards the 3 major product groups: C₂ hydrocarbons, C₃₋₅ hydrocarbons and aromatics, keeping the conversion constant at 10%. This conversion level was achieved by either addition of ethane, addition of ethylene, or by changing the flow rate and consequently space-velocity without any addition of C₂. The selectivity data at 10% conversion of hydrocarbon in the feed stream, assuming full conversion of the added C₂ species, are obtained by interpolation as presented in the ESI,† Fig. S1 and S2. It is



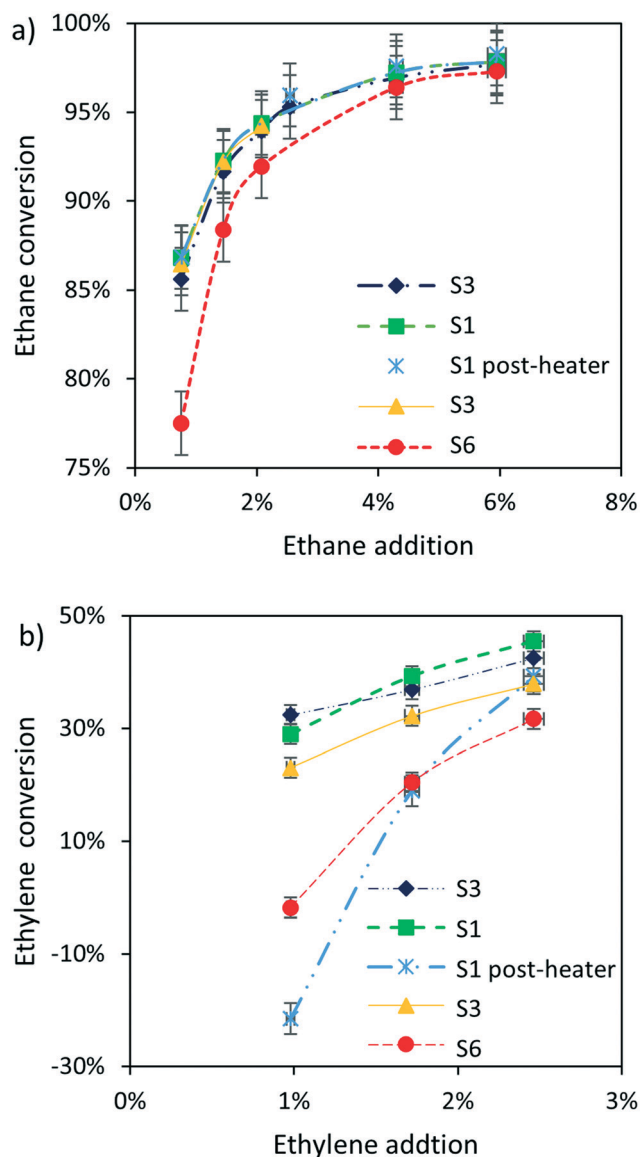


Fig. 4 C_2 conversion as function of C_2 addition, (a) ethane, (b) ethylene; 90% CH_4 N_2 balance; 16.6 ml min^{-1} total flowrate. Reactor-zone at $1000 \text{ }^\circ\text{C}$, pre-heater and post-heater at $400 \text{ }^\circ\text{C}$, except for S1 post-heater having $T_{\text{post-heater}}$ at $1000 \text{ }^\circ\text{C}$.

assumed that the added C_2 compounds are completely converted whereas any ethane and ethylene detected in the product is assigned to formation in the reactor. In Fig. 6, data are interpolated to 10% conversion of hydrocarbons in the feed stream including full conversion of the added C_2 compounds. Fig. S5† presents the same type of analysis at 10% methane conversion, disregarding the conversion of C_2 compounds, resulting in very similar trends. Fig. 6 shows that, within the error margin, the selectivity to the various product groups is constant, independent of the method to enhance CH_4 conversion to 10%, *i.e.* C_2 addition or decreasing space velocity. The formation of aromatic products is somewhat suppressed when ethylene is added, as well as when a full-catalyst bed is used (S6). The product distribution of all hydrocarbon products is given in ESI† Fig. S4.

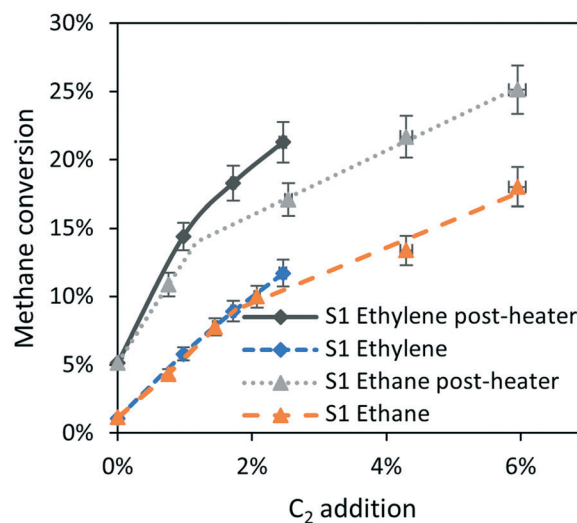


Fig. 5 Effect of the post-heater on methane conversion with ethane or ethylene added, compared the effect of ethane addition without the post-heater (identical to S1 case Fig. 3b). 90% CH_4 , N_2 balance; 16.6 ml min^{-1} total flowrate. Reactor-zone and post-heater at $1000 \text{ }^\circ\text{C}$ (except for S1 ethane case), pre-heater at $400 \text{ }^\circ\text{C}$.

Fig. 7 shows the product yield to the three main product groups as function of conversion obtained with the small catalysts bed (S1) with and without the post-heater at high temperature, as a result of adding ethane (Fig. 7a) and ethylene (Fig. 7b). C_2 yield and C_{3-5} yield are independent of the use of the post-heater, in contrast the aromatics yield which increased when using the post-heater.

Discussion

Addition of ethane

Methane conversion is significantly increased on ethane addition, apparent from Fig. 3a and b. The explanation for this significant enhancement in methane conversion is found in the studies of Roscoe and Thompson²⁷ as well as Dean²⁸ and schematically presented in Fig. 8. Formation of higher hydrocarbons from ethane, *via* ethyl radicals, releases hydrogen radicals which react with methane according to $CH_4 + H \rightarrow CH_3 + H_2$, which is the dominant reaction for activation of methane. The methyl radicals react to ethane, which maintains the cycle in Fig. 8, known as methane auto-catalysis.^{22,27,28} In the presence of catalyst, methyl radicals form on the Fe active sites while the hydrogen atoms stay on the active site, to combine to form di-hydrogen, according to DFT calculations.^{19,25}

A similar enhancement in methane conversion is also observed in non-catalytic operation in literature.²⁹⁻³¹ Germain *et al.*³⁰ and Rokstad *et al.*³¹ observed decreasing effectiveness of ethane addition on methane conversion when the ethane concentration is above 2% in absence of catalyst, very similar to the observation with catalyst in Fig. 3b, which Germain attributed to scavenging of free radical by formed aromatic species.



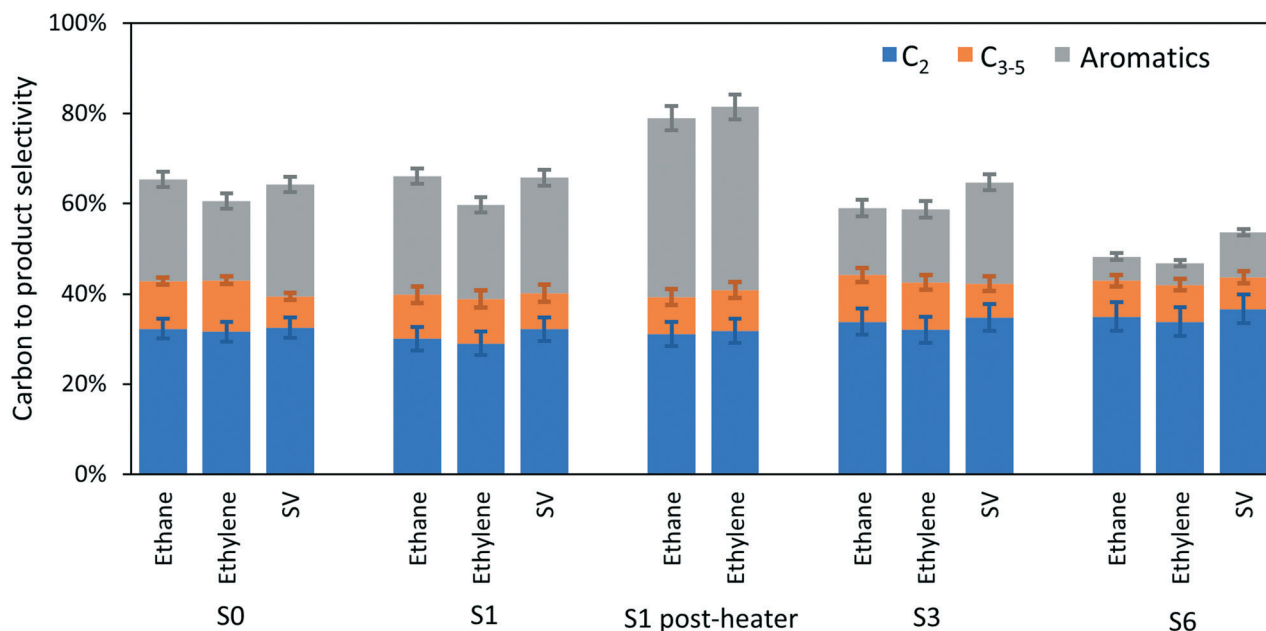


Fig. 6 Product selectivity distribution for different methods of increasing methane conversion, by ethane addition or ethylene addition to the reactant mixture or by decreasing space velocity. The results have been linearly interpolated at 10% total hydrocarbon conversion as explained in the experimental section. Similarly, selectivity is also calculated based on total hydrocarbon conversion. Reactor-zone at 1000 °C; pre-heater and post-heater are both at 400 °C, except for the S1-post-heater in that case $T_{\text{post-heater}} = 1000$ °C. The total flowrate for the cases with ethane or ethylene addition is 16.6 ml min^{-1} 90% CH_4 , N_2 as balance. The graphs used for the interpolation can be found in ESI† Fig. S1 and S2.

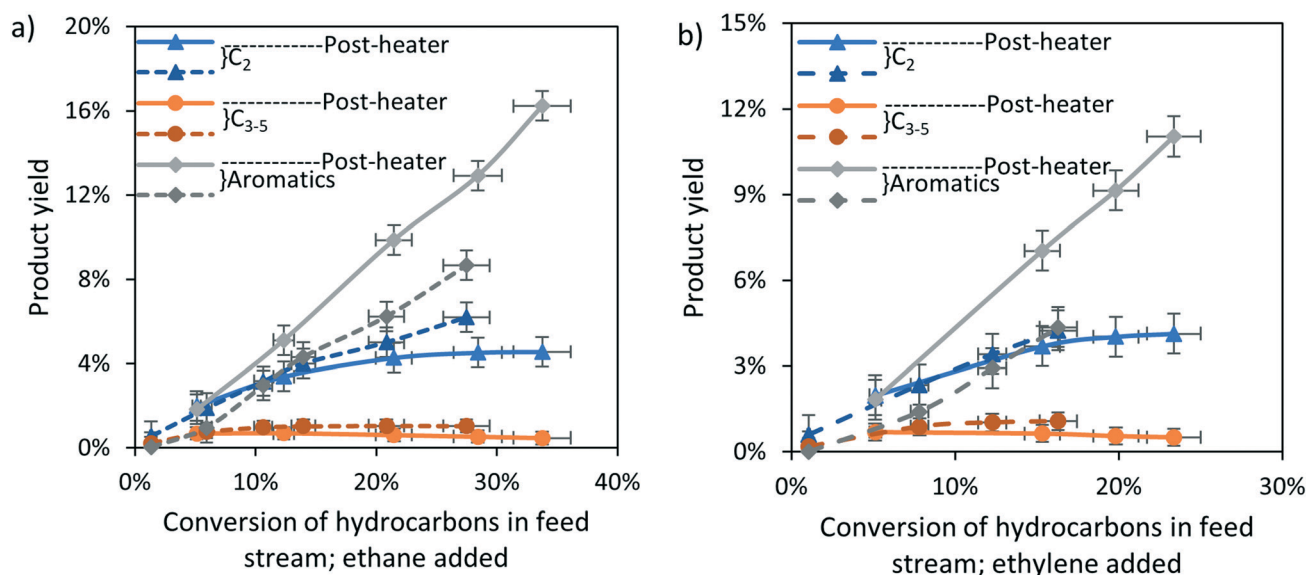


Fig. 7 Effect on product yield when using the post-heater in the S1 case, (a) ethane, (b) ethylene; 90% CH_4 N_2 balance; 16.6 ml min^{-1} total flowrate. Reactor-zone at 1000 °C, pre-heater and post-heater at 400 °C, except for S1 post-heater having $T_{\text{post-heater}}$ at 1000 °C.

The catalyst has no effect on methane conversion at ethane concentrations above 2%. This shows that ethyl radicals and consequently H radicals mainly form in gas phase whereas catalytic ethane activation is negligible. This is further supported by the linear correlation between the enhancement of methane conversion due to ethane addition and the free volume available in the reactor, as shown in Fig. 3d, as well as by the observation that a large amount of catalyst reduces the

ethane conversion, as shown in Fig. 4a. Methane conversion is not influenced by the presence of catalyst when at least 2% ethane is added, as shown in Fig. 3b. Apparently, enhancement of methane conversion on the catalyst is compensated by a decrease in methane conversion by decreasing the free volume when catalyst is introduced.

Fig. 5 shows that the increase in methane conversion due to an extended post-catalytic residence time²² is also



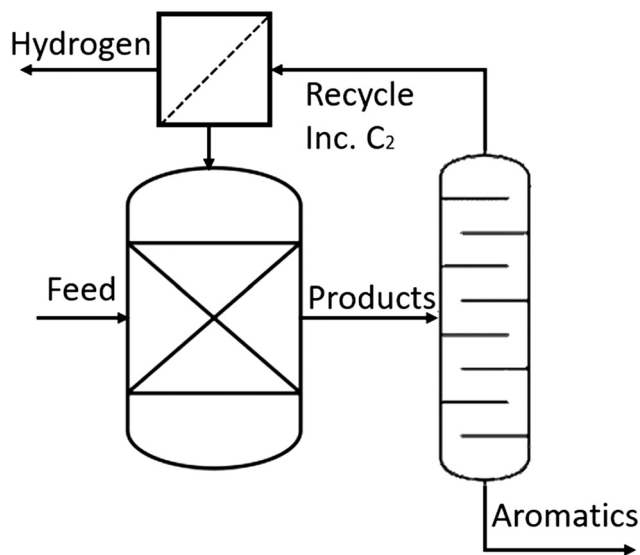


Fig. 9 Basic schematic concept process for natural gas to aromatics based on NOCM including C_2 recycle.

reactor volume. C_{2+} hydrocarbons are often present in natural gas with typical concentrations similar to the concentrations used in this study, for example Gulf-coast natural gas contains up to 4.9 vol% C_2 hydrocarbons¹ (details in Table S1†). Our study implies that removal of ethane from natural gas is not required, simplifying the purification of natural gas. The catalyst becomes obsolete at C_2 concentrations above 2 vol% as shown in Fig. 3. Thus, the reaction can be operated at high productivity without requiring a catalyst, simplifying the reactor design and operation. Deposit build-up on the catalyst can be prevented and catalyst purchasing, handling, disposal and plant-stops for catalyst replacement are all not required. Rapid control of reactor capacity would be possible by tuning the concentration of the C_2 initiator in the reactor feed.

Overall ethylene conversion is typically low or even negative, meaning net-production at high methane conversion levels as shown in Fig. 4b. Ethylene can be recycled back into the reactor, to generate a process with a net-zero consumption of ethylene, as presented in the schematic process diagram in Fig. 9. In this scheme, the aromatic products are separated from the olefins, which are recycled back to the reactor, after separation of hydrogen. Fig. S3† shows that the yields of ethane as well as C_{3-5} hydrocarbons becomes negligible at high methane conversion levels. The activity increase in the reactor can be achieved both by recycled ethylene as well as ethane in the feed. Fig. 4 shows that overall ethylene conversion is positive at higher added ethylene concentrations and negative, *i.e.* net-production, at lower ethylene concentrations, demonstrating an internal feedback-loop that will prevent both ethylene accumulation as well as depletion in the process. This results in a process converting methane to aromatics, also alleviating the energy intensive cryogenic recovery of ethylene. The highest single pass methane

conversion achieved in this paper, 25%, with 16% aromatics yield is already higher than benchmark performance of MDA, around 15% conversion and 11% aromatics yield.¹⁶ On the other hand, it must be noted that the main product is naphthalene compared to more valuable benzene produced in the MDA reaction and additional conversion might be required. Also, this scheme is obviously not producing any ethylene as originally targeted with methane pyrolysis.

Conclusion

The effect of C_2 addition on non-oxidative methane coupling at high temperature was investigated. Addition of ethane and ethylene (C_2) up to 6 vol% in methane significantly increases methane conversion during both catalytic as well as non-catalytic NOCM. The activation of methane through C_2 addition is dominated by homogenous gas-phase reaction and catalytic ethane or ethylene conversion is negligible. Product selectivity at constant conversion is unaffected by C_2 addition and is influenced by catalyst amount and residence time in the hot zone of the reactor, both within and downstream of the catalyst bed, in line with our previous work. The catalyst has no effect on methane conversion at C_2 concentrations above 2%, showing that an optimal reaction design involves either no catalyst at all or a minimal amount of catalyst. The catalyst is needed to initiate methane coupling *via* radical chain reactions only when less than 2 vol% of ethane or ethylene are present in the feed. C_2 addition during NOCM significantly increases productivity and allows simplification of purification of the reactor-feed. A process design is envisioned in which the produced ethylene is recycled back into the reactor to initiate the methane conversion, resulting in a process converting methane to aromatics.

Conflicts of interest

There are no conflicts to declare.

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