The Direct Dimethyl Ether (DME) Synthesis Process from Carbon-Based Feed Stocks: Current Status and Future Prospects II. Kinetic Studies and Catalyst Deactivation

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Abstract
In Part I of this series, it was seen that the favorable thermodynamic and kinetic coupling in the LPDME™ process—of methanol dehydration reaction (very rapid and at/near thermodynamic equilibrium) with the methanol synthesis reaction (slower kinetics and highly thermodynamic) —leads the beneficial "chemical synergy". This synergy helps to overcome the limitation on thermodynamic equilibrium conversion and increases the per-pass syngas conversion and reactor productivity. This increase in the per-pass syngas conversion can be as high as 50-100% and depends primarily on the feed \( \text{H}_2: \text{CO} \) ratio.

In this part II of Series, we briefly discern the intrinsic kinetics of the LPMeOH™ and LPDME™ systems, and also shed light of the catalyst deactivation phenomena in these processes. Among the many reports on intrinsic kinetics of the one-step LPMeOH™ and LPDME™ processes, two illustrative kinetic studies, from the groups of University of Akron and Air Products and Chemicals, Inc. are highlighted and discussed further. These are mainly based on the independent, component kinetic models of methanol synthesis (Vanden Bussche and Froment) and methanol dehydration (Bercic & Levec). From an overarching analysis of the deactivation of supported copper catalysts for methanol synthesis and other reactions (methanol decomposition and methanol steam reforming), we propose that thermal sintering, i.e., increase in Cu particle site and loss of metal surface area, is the only cause of catalyst deactivation in methanol synthesis reactions over Cu/ZnO/Al_2O_3 industrial-type methanol catalysts. In closing, we outline the reactor design/scale-up and plant operational experience of the 3 commercial technologies, as currently practiced by JFE holdings, BP-AMOCO, and Halder-Topsoe.

Keywords: Natural gas; Steam reforming; Coal; Syngas; Methanol; DME; Bi-functional catalysts; Cu/ZnO/Al_2O_3; γ-Al_2O_3; Slurry reactors; Bubble column reactors; Chemical synergy; Methyl equivalent productivity (MEP); Intrinsic kinetics; Phase equilibrium; Chemical reaction equilibrium; Catalyst deactivation

Introduction
In Part I of this series, we illustrated the process feasibility of the LPMeOH™ and LPDME™ processes and shed an illuminating light on the chemical synergy and role of the water gas shift reaction in the overall process chemistry. These 2 processes have been illustrative examples of how a mature technology on a commercial scale (ICI low temperature methanol synthesis process) can be successfully adapted to a liquid phase operation. It is possible that this adaptability and flexibility of a cost-effective liquid phase alternative can be extended to other parallel reactions; involving multiple components; especially if the overall reaction is highly exothermic and reversible. The LPMeOH™ process is a highly flexible process that is well-suited to syngas feeds of all \( \text{H}_2: \text{CO} \) ratios (from \( \text{H}_2 \)-rich to \( \text{CO} \)-rich); uses milder reactor design conditions (of temperature and pressure); and a simpler one from a process engineering standpoint. Some of the key advantages of the LPMeOH™ process include better heat transfer characteristics and isothermal operation; use of the \( \text{CO} \)-rich syngas (from low-cost coal sources); and a very high chemical selectivity to methanol.

The process chemistry of LPDME™ system is covered in detail in Part I of this Series. The process is based on application of dual catalysis in a single reactor stage; and based on a combination of an equilibrium-limited reaction (methanol synthesis) and an equilibrium unlimited reaction (methanol dehydration). The selective chemical removal of product methanol; from Reaction (1); via its dehydration; via Reaction (3); overcomes the chemical equilibrium barrier on methanol synthesis alone; and improves the per-pass syngas conversion and reactor productivity.
Kinetic Studies on LPMeOHtm and LPDMEtm Processes

In this Part II; we focus on the kinetic studies of the liquid phase methanol; LPMeOHtm and direct one-step DME synthesis from syngas; the LPDMEtm process. The kinetic studies often have important implications for process design; scale-up; and commercial operation. The literature studies on the intrinsic kinetics can be classified into 3 distinct categories:

a) Kinetics of vapor phase synthesis of methanol from CO/CO2 and H2 (ICI low pressure methanol synthesis; over Cu/ZnO/Al2O3 catalyst)

b) Kinetics of liquid phase methanol from CO/CO2 and H2 over Cu/ZnO/Al2O3 (LPMeOHtm process; by Chem Systems; Inc.)

c) Intrinsic kinetics of the direct; one-step DME process (LPDMEtm process; which was introduced by UA researchers; in 1989)

Here we will only consider the state-of-art literature on kinetic studies on the kinetic phase methanol and liquid phase DME (LPDMEtm) processes and provide a critical assessment of proposed kinetic rate expressions; and discern the benefits and drawbacks of our current understanding of the literature on intrinsic and in-situ kinetics. It is interesting to note that the vapor phase mechanisms and proposals for kinetic rate expressions are more difficult to discern than the liquid phase systems because of the differences in reactor designs/operational conditions in differential or pseudo-differential modes (which can confound data analysis and comparison to integral modes of operation); a problem not encountered in the liquid phase mode of operation.

For development of intrinsic kinetic models of LPMeOHtm and LPDMEtm systems; it is first necessary to develop a detailed thermodynamic framework of this multi-component; multiphase system; where at least 8 independent components participate in 3 parallel reactions (methanol synthesis from CO/CO2; water-gas shift; and methanol dehydration; over solid catalysts; well-dispersed in Witco-40 inert mineral oil). We illustrate briefly the computational procedure to develop the multi-component phase and chemical reaction equilibria models for LPMeOHtm and LPDMEtm processes. We refer to the interested reader a few seminal reports from the researchers at University of Akron [1-3]. Regardless of the kinetic model and the type of formalism chosen to describe it (L-H; power law; etc.); it is first necessary to compute the concentrations (and fugacities) of all species; i.e.; reactants and products; in the liquid phase. To this end; extensive experimental studies on the binary and multi-component phase equilibria of syngas components at a range of pressures and temperatures; to cover the range of practical interest; were carried out by UA researchers’. The data from the binary experiments was incorporated into a multi-component phase equilibria model-the Margule’s equations were used to model the binary vapor-liquid equilibria and to generate the Margule’s parameters; using Lewis-Randall standard state for vapors/liquids (CH3OH; H2O; DME; Witco-40); and the Soave-Redlich-Kwong equation of state (EOS) was used to model the multi-component phase equilibria; with Henry’s law constants as standard states for permanent gases (H2; CO; CO2; CH4). A complete computer program package (in FORTRAN-90) was developed by the UA research group; that calculates concentrations of all reactive species in the liquid phase; based on the simultaneous multi-component phase and multi-reaction chemical equilibria [1-3]. As input data; only the vapor phase compositions of all species (from experimental data); pressure; temperature; and volume of oil are required. The system of non-linear algebraic equations with 8 unknowns (coupled) is solved using the Newton-Raphson iteration method; with Gauss-Jordan reduction algorithm for computation of inverse Jacobians; for rapid convergence. The detailed calculations of thermodynamic equilibrium constants for the 3 principal reactions; KH2; KCO and KCH3OH; as functions of T; are readily available in the literature [4-9].

The first results on the kinetic studies on the LPMeOHtm process were presented by the APCI group at the International Symposium of Chemical Reaction Engineering (popularly given as I.S.C.R.E.); in Edinburgh; Scotland; in 1984 [10] and subsequently presented in a more refined form at 2 other Symposia/Conferences [11,12]. Later; researchers’ from APCI presented their kinetic models in a Langmuir-Hinshelwood form. The APCI work was also presented in another form-a modified L.H. form-also including an activity factor; A; to account for catalyst deactivation; i.e.; to allow for the gradual loss in activity as a function of time on stream [13-17]. An overall analysis of APCI kinetic studies indicates that the apparent activation energy is in the range of 75,000-90,000J/mol; and the pre-exponential factors are of the order of k ij = 2.43x106 mol CH3OH/s/kg cat. MPa; and for the kinetic model based on fugacity terms; kij = 3.68x106 mol CH3OH/kg cat.atm. As an anecdotal note; Sherwin & Blum of Chem Systems; Inc; proposed the very first kinetic rate expression of LPMeOHtm process; as a power law form; but incorrectly used the CO-CO2 term as a variable for the functional form (and not CH3OH-CO2). It is interesting to note that almost all kinetic rate expressions on liquid phase methanol seem to overwhelmingly agree that CO exhibits a very weak dependence on methanol rates-the reaction order with respect to CO is often found to be close to zero. At the same time; three independent research groups; UA (Akron; Ohio); University of Aachen/Groningen (Germany/Netherlands); and Technical University of Lodz (Poland) also proposed the kinetic rate expressions for the LPMeOHtm process. For example; the Lodz research group [18] proposed a kinetic rate expression written as the difference between a forward kinetic term and a backward kinetic term (for the reverse reaction):

\[ r_{\text{CH3OH}} = k_{\text{F}} \exp(-E_{\text{F}}/RT) - k_{\text{B}} \exp(-E_{\text{B}}/RT) \text{MPa}^{-1} \text{CH3OH} \]  

(1)

The values of E F and E B; according to the kinetic model; were given to be 56000J/mol and 85000J/mol. The forward and backward kinetic terms were given as 5.5x106 and 6x105 mol CH3OH/sec. kg cat. MPa-1 [The negative exponent on the MPa unit; of -2; indicates the value of Δn; or; change in number of moles]. The low estimates of reaction orders; 0.4 for H2 and 0.18 for CO; indicate that they are likely not true reaction orders; but apparent ones; likely influenced by external mass transfer (due to high slurry ratios); or even pore diffusional issues. The particle size; of 60μm; is rather high. The
work of Von Wedel et al. [18] was extended by Ledakowicz et al. [19]; who proposed a kind of hybrid kinetic model-a power law combined with elements of L-H formalism. In contrast with the works above and the extensive kinetic analysis of Graef et al. [4,5] that follows; based on a detailed micro-kinetic model of methanol synthesis reaction; a relatively simple kinetic analysis was presented by the UA research group [1]; for liquid phase methanol process; over commercial Cu/Zn/Al catalyst of United Catalysts; Inc. The rate of methanol synthesis was found to be independent of partial pressures of CO; CO\textsubscript{2} and H\textsubscript{2}O. Consequently; the methanol rate was modeled only as a function of the concentration driving force (approach to equilibrium) of H\textsubscript{2} as follows:

\[ r_\text{CH}_3\text{OH} = k_{eqm3}(\text{CH}_3\text{OH})^{n} \]  

(2)

A statistical regression analysis of the rate data as a function of the H\textsubscript{2} driving force showed that the reaction order; n; ranged from 0.80 to 1.15; and could be approximated well by the value of 1; over the entire range. This model; although simple; is in good agreement with all literature results; which suggest that methanol synthesis rates exhibit a very weak dependence on concentrations of CO in the liquid phase. An extended kinetic model for the liquid phase methanol process was developed by [4,5]. Their experimental component was carried out in a 300cm\textsuperscript{3} stirred autoclave; with a very low catalyst loading; to ensure absence of external gas-to-liquid mass transfer limitation. For the three principal reactions; CO hydrogenation; water-gas shift (CO\textsubscript{2}+H\textsubscript{2}); and CO\textsubscript{2} hydrogenation; a detailed micro-kinetic model based on series of elementary surface reactions was proposed. Based on a series of 4 elementary reactions for (A); 2 reactions for (B); and 6 reactions for (C); a total of 48 (4x2x6) kinetic models could be constructed and were considered; for further analysis. Based on experimental data; statistical analysis (\chi\textsuperscript{2}-test; with 95% C.I.s; and analysis of error residuals); and physico-chemical constraints; a model termed as A3-B2-C3 was proposed to be the best fit for the kinetics of liquid phase methanol process. The typical L-H forms of kinetic rate expressions are given in [4,5]. For the APCI portfolio; we begin by considering “seminal” works from the APCI research; both from 1999 [20,21]. The work of Peng et al. [21] is also the most specific in regards the intrinsic kinetics of direct; one-step DME synthesis in slurry mode of operation. For all kinetic studies; the reactor was considered to behave as a CST Rand was free of external mass transfer (external to liquid) limitations and pore diffusional limitations. All three expressions are power law form multiplied by an approach-to-equilibrium term as shown below:

a. Methanol synthesis reaction:
\[ R_a = k_a \exp\left(\frac{\Delta H_a}{RT}\right) \]  

(3)

b. Forward water-gas shift:
\[ R_p = k_p \exp\left(\frac{\Delta H_p}{RT}\right) \]  

(4)

c. Methanol dehydration reaction:
\[ R_r = k_r \exp\left(\frac{\Delta H_r}{RT}\right) \]  

(5)

The models have excellent predictive capability and the methyl equivalent productivities (MEPs) as predicted from the kinetic models are in good agreement with experimental results from the data gathered at conditions as above (Figure 1) [21]. In fact; the models have excellent predictive power across the range of syngas types; or H\textsubscript{2}:CO inlet ratios; for the 3 typical syngas feeds; viz.; Texaco (Koppers-Totzek); the most typical composition of CO-rich syngas; Shell (similar to Texaco); and Dow (close to H\textsubscript{2}-rich type). The second report from APCI research by Peng et al. [21] is on intrinsic kinetics and modeling of the direct; one-step DME process; in the vapor phase. Ng et al. investigated the kinetics of the one-step DME synthesis process over a commercial CuO/ZnO/Al\textsubscript{2}O\textsubscript{3} methanol catalyst and a γ-alumina catalyst at 250 °C and 5MPa; using a gradient-less; internal-recycle-type reactor (Berty-type). A kinetic model for the combined synthesis methanol and DME (co-production of methanol and DME); based on a methanol synthesis kinetic model proposed by Vanden Bussche and Froment (1996) and a methanol dehydration kinetic model by Bercic and Levec (1992) was tested using experimental results obtained in the Berty-type reactor from a wide range of syngas compositions and CO/CO\textsubscript{2} ratios.

The kinetic parameters of the work of peng et al. [21] are summarized in Figure 1. The proposed kinetic models of peng et al. [21] and the kinetic parameters that are derived from the best fits to the experimental data are given in (Figure 1-3); Table 1; respectively. The illustration of chemical synergy in the LPDME\textsuperscript{™} process system (methanol+DME) vs. LPMeOH\textsuperscript{™} alone is given in Figure 2. The chemical synergy is a very strong function of the H\textsubscript{2}: CO ratio and the fraction of CO\textsubscript{2} in the feed syngas. Finally; it is worth noting that most kinetic models proposed in the literature are based on direct CO\textsubscript{2} hydrogenation via formate as the intermediate [22,23]. The 3 customarily proposed reaction mechanisms for direct CO\textsubscript{2} hydrogenation comprise of formate; formyl (via RWGS reaction); and carbonyl. The three mechanisms are illustrated in Figure 2.

**Table 1:** Kinetic rate parameters for the rate models given in Figure 1.

| Parameters | A(i) | B(i) |
|------------|------|------|
| k\textsubscript{1} | 1.65 | 36,696 |
| k\textsubscript{2} | 3.61 × 10\textsuperscript{-5} | 0 |
| k\textsubscript{3} | 0.37 | 17,197 |
| k\textsubscript{4} | 7.14 × 10\textsuperscript{-11} | 124,119 |
| k\textsubscript{5} | 1.09 × 10\textsuperscript{-10} | -94,765 |
| K\textsubscript{APCI1} | Twigg (1986) | |
| K\textsubscript{APCI2} | Twigg (1986) | |
| K\textsubscript{hydr} | 7.9 × 10\textsuperscript{-4} | 70,500 |
| k\textsubscript{K} | 3.7 × 10\textsuperscript{-10} | -105,000 |
| K\textsubscript{hydr} | 0.84 × 10\textsuperscript{-1} | 41,100 |
| K\textsubscript{hydr} | Stull et al 1969 | |
Figure 1: Kinetic rate models for methanol synthesis reaction via CO₂ hydrogenation, water gas shift, and methanol dehydration reaction.

Figure 2: The extent of chemical synergy in LPDME™ vs. LPMeOH™ alone in terms of the effect of GHSV (h⁻¹) on the methanol equivalent yield (MEY), at 250 °C and 50 bar and with a gas composition of 18% CO, 72% H₂, and 10% He.

Figure 3: Surface reaction mechanisms and reactive intermediates for direct CO₂ hydrogenation to methanol over Cu catalysts.
Catalyst Deactivation in LPMeOH<sub>tm</sub> and LPDME<sub>tm</sub> Process

An important distinction needs to be made at the very outset here; between vapor and liquid phase modes of operation. For methanol synthesis; this distinction is more pertinent; as the one of the primary reasons for the liquid phase methanol synthesis process (LPMeOH<sup>tm</sup>) was introduced by Chem Systems (now Necant) in 1975 was to control the very highly exothermal nature of the commercial IC<sub>i</sub> vapor phase methanol process; and thus reduce the thermal sintering phenomena. In this section; we will make only a brief but important comment on the long-term activity patterns (vs. time) and catalyst stability and will focus only the catalyst deactivation phenomena for LPMeOH<sup>tm</sup> and LPDME<sup>tm</sup> processes and elucidate how the presence of the liquid phase is able to moderate the exothermal nature of methanol synthesis reactions; and; in fact; extend on-stream catalyst lifetimes. For sake of completeness; we refer the interested reader to the following papers on the catalyst deactivation of bi-functional catalysts for direct; one-step DME vapor phase process and the catalytic dehydration of methanol to DME [24-33].

An interesting question that has always been confounding to researchers is the nature of the catalyst deactivation--of individual micron-sized Cu/Zn/Al methanol component and the most common γ-Al<sub>2</sub>O<sub>3</sub> or H-ZSM-5 DME component in the LPDME<sup>tm</sup> process; when used in the form of a physical mixture (also called an admixture) and dispersed in an inert mineral oil (like Witco-40; Witco-70; or Freezeene-100); at typical operating conditions of 250 °C and 70atm; as a function of on-stream time. It is intuitively obvious; however; that the thermal sintering of the copper component--most often proposed to be cause of catalyst deactivation in the vapor phase and validated by many studies; including most notably those of [34-36]; Kung [24] will be greatly alleviated in the liquid phase operation; as the presence of the liquid phase (with its high heat capacity)serves to moderate the strongly exothermal nature of methanol synthesis reactions; and the presence of “hot spots”; in particular. As the finely dispersed copper component is completely wetted/filled (external and internal surface of pores) by the liquid phase; the reaction is essentially “isothermal”; with no local heat gradients or “hot spots”. The mechanism of thermal sintering that leads to an increase in size of copper crystallites with the increase in surface area and a concomitant decrease in methanol rates and TOFs—is thus vastly “suppressed” in the liquid phase.

The presence of in-situ produced CH<sub>3</sub>OH and H<sub>2</sub>O within the pores of Cu/Zn/Al and γ-Al<sub>2</sub>O<sub>3</sub> is often intuitively considered to be the cause of hydrothermal sintering (i.e.; contribute to the “thermal” sintering effects) and even leaching of the copper and zinc components. Obviously; it is more difficult to remove H<sub>2</sub>O from the surface of Cu-based catalyst in a slurry reactor configuration; as the surface is within the pore structure of the Cu/ZnO catalyst; completely wetted by the inert liquid phase. This liquid phase poses an additional barrier for a facile back-diffusion of products; CH<sub>3</sub>OH and H<sub>2</sub>O; back into bulk liquid and into vapor phase. However; we do not agree with the hypothesis of hydrothermal leaching or loss of crystallinity as primary causes of deactivation of Cu and Zn crystalline phases.

Our seminal works at University of Akron in Prof. Lee’s group with A. Sawant and B. Lee as doctoral candidates [37-44] and other works conclusively prove that the hydrothermal sintering that leads to a crystal size growth of Cu and the concomitant loss of Cu surface area are the only plausible and possible causes of catalyst deactivation in this phenomena; and the physicochemical interaction between the 2 components is not just benign but can be synergistic. In fact; it is quite apropos to state here that the in-situ processes developed by the UA researchers’ for regeneration of aged copper catalysts during LPMeOH<sup>tm</sup> processes are based on the principle of “crystallite re-dispersion”; i.e; size-reduction accompanied re-dispersion of crystallite sizes. These processes have been demonstrated under bench and pilot scale conditions and even commercial scale operation; at the commercial-scale LPMeOH<sup>tm</sup> and LPDME<sup>tm</sup> processes at Eastman Chemical Co’s plant site in Kingsport; TN; and can restore completely the original particle size distribution of Cu crystallites and the initial activity. The applicability of catalyst regeneration process for aged (deactivated) catalysts based on the principle of crystallite re-dispersion underscores our assertion of the primary governing mechanism for deactivation and aging; i.e.; thermal sintering and crystal growth that leads to a loss in surface area of active copper.

It is interesting to note that the fact that thermal sintering is the primary mechanism of catalyst deactivation in Cu catalysts in the LPMeOH<sup>tm</sup> and LPDME<sup>tm</sup> processes has been asserted and corroborated by other research reports [45,46]. However; other causes including presence of trace carbonyls such as Fe(CO)<sub>5</sub> and Ni(CO)<sub>4</sub>; carbon deposition by reverse boudouard reaction; and hydrothermal leaching by the high water concentrations in the slurry; can also be present to a smaller extent and can be important. The poisoning by carbonyl sulfide-a compound of CO and sulfur-can also be more significant in the liquid phase processes. These causes can be considered to be minor and can be eliminated to a great extent by good plant design/engineering practice.An interesting mechanism of catalyst deactivation in dual catalysis; or LPDME<sup>tm</sup> process; was proposed by Peng et al. [21] as detrimental physical interaction of methanol and DME catalyst components. The data as reproduced in Figure 4 is considered to be questionable and thus unreliable.

Commercial Technologies and Future Markets

Di-methyl ether (DME) is being touted for its potential as a very versatile chemical intermediate as well as a clean-burning alternative fuel and a “green” substitute for diesel; as it does not emit any soot or particulate matter (PM) upon combustion in diesel engines [47-50]. DME is a colorless; odorless gas with physical and chemical properties similar to that of “liquefied petroleum gas (LPG)” which is popular in Southeast Asia (India; China; Pakistan; Bangladesh; etc.) and Pacific including Japan; for household cooking and even heating. Like LPG; DME can be sold in standard bottles/cylinders as a pressurized liquid at about 5atm; and holds a higher
The direct; one-step DME process as described in this paper and the various elements that comprise the current research areas on bench-scale and especially at pilot scale (principally at UA and APCI research) has now formally made the transitional debut at the commercial scale; in Hokkaido; Japan.JFE company (previously NKK Nippon Corporation); in collaboration with Taiheiyo Coal Mining Co.; Sumitomo Metal Industry; Ltd.; and CCUJ (Center for Coal Utilization; Japan); have already demonstrated this technology on a 5ton DME/day (5TPD) pilot scale in 2002 [40-42]. Based on this achievement; JFE Co. successfully translated this pilot scale demonstration into a commercial-scale 100 ton DME/day (100 TPD) DME plant at its site in Hokkaido; Japan; starting 2005.

Korea Gas Corporation (KGC); Halder-Topsoe (based in Denmark); Air Products (Allentown; PA; U.S.A.); and BP-AMOCO (Des Plaines; IL; U.S.A.) are also on the commercialization track for the direct; one-step DME process/technology. A Halder-Topsoe patent (WIPO International Patent WO96/23755) outlines their research efforts towards development of the direct; one-step DME process. The status of DME technology development efforts by BP-AMOCO for their licensees in China and other players in Asia are summarized in a review [46]. A totally new and highly innovative “application” of the direct; one-step DME technology; which is inherently simpler and cost-effective (when compared to the conventional process with 2 separate steps) can be thought of in the traditional gas-to-liquids (GTL) market segments. The GTL technologies offer the owners/operators of stranded gas reserves a cheap alternative to ship stranded gas in the form of liquids; which are easier and more economical to transport. However; the technologies are complex; costly; and tightly held by a few major companies; which present significant impediments for new (and independent) large-scale plants; in terms of capital costs and long-term technology transfer decisions. In this regard; a small-scale; direct DME technology modular application; can be ideal for such stranded gas reserves or even for associated gas (typically sent to flares); for local end uses.

Summary

In Part I of this Series; we illustrated the chemical synergy in the LPDME™ process; and saw how the combination of an equilibrium-limited reaction; methanol synthesis; and an equilibrium-unlimited one; methanol dehydration to form dimethyl ether; can help lift the chemical equilibrium limitation on methanol synthesis alone.
This thermodynamic-kinetic coupling is the basis of the chemical synergy observed in both LPMeOH™ and LPDME™ processes. In this part II of Series; we have discerned the intrinsic kinetics of the LPMeOH™ and LPDME™ systems; and also shed light of the catalyst deactivation phenomena in these processes. From an overarching analysis of the deactivation of supported copper catalysts for methanol synthesis and other reactions (methanol decomposition and methanol steam reforming); we have demonstrated that thermal sintering; i.e.; increase in Cu particle size and loss of metal surface area; is the only cause of catalyst deactivation in methanol synthesis reactions over Cu/ZnO/Al₂O₃ industrial-type methanol catalysts.

The future of DME as an alternative fuel and a chemical intermediate/commodity for targeted end use is very bright. The direct; one-step DME process as described in this paper and the various elements that comprise the current research areas on bench-scale and pilot scale (principally at UA and APCI research in the United States and Halder Topsoe in Denmark) has now formally made the transitional debut at the commercial scale; in Hokkaido; Japan. The JFE Holdings owned commercial-scale DME plant has an on-stream capacity of 300 TPD and has been in operation since 2007. Korea Gas Corporation (KGC); Halder-Topsoe (based in Denmark); Air Products (Allentown; PA; U.S.A.); and BP-AMOCO (Des Plaines; IL; U.S.A.) are also on the commercialization track for the direct; one-step DME process/technology.

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