Catalytic transformation of ethanol into 1,3-butadiene

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Abstract
1,3-Butadiene is an important constituent of many products that we rely upon. It is currently prepared from non-sustainably derived sources. However, in the early part of the 20th Century the use of ethanol as a source of 1,3-butadiene has been reported. With the arrival of a cheap and bountiful supply of crude-oil derived sources the need for the sustainable route was deemed unnecessary. In recent years the conversion of ethanol to 1,3-butadiene has undergone somewhat of a mini resurgence as the chemical industry looks to try to find a sustainable and secure route to this important building block. This review will emphasise some of the most recent work in the field and look ahead to what needs to be achieved to make this research a reality.

Keywords: Ethanol, 1,3-butadiene, Heterogeneous, Catalysis

Introduction
1,3-Butadiene (1,3-BD) is an important building block in many chemical processes. Its main use is as a monomer for the production of synthetic rubbers [1]. The most significant use of 1,3-BD is in the manufacturing of styrene-butadiene (SBR) rubbers which are primarily used in the production of tyres. Currently, 1,3-BD is mainly formed as a by-product of the naphtha steam cracking process – a coproduct of ethene manufacturing – with 1,3-BD being isolated after costly extractive distillation steps. 1,3-BD can also be produced by the dehydrogenation of butane or butene (Houdry process) [1,2]. In recent years the cost of 1,3-BD has fluctuated massively and the price has increased, for example the cost of a tonne of 1,3-BD was ca. $1,500 USD in March 2014 compared to $850 USD in August 2013. The cost (and fluctuation) is not sustainable in the long term for the major users of this important building block. The increase in cost can be attributed to several factors i) the increasing rise in the price of crude oil; ii) the move to lighter feedstocks from the cracking process and iii) the “shale-gas” surge in Europe and the US [3]. Shale gas contains ethane which can be dehydrogenated to ethene which is a 1,3-BD free route and consequently causing a reduction in ethene production from the steam cracking route and, hence, 1,3-BD. This necessitates the need to produce 1,3-BD via a bio-based route. Whilst “shale-gas” has its sceptics it may well necessitate the increased development of 1,3-BD from renewable sources. Increased dependence on “shale-gas” as a source of natural gas may lead to acute shortfalls in the supply of key starting materials for industry that are currently sourced from more traditional feedstocks. Paradoxically, the chemical industry needs to look back in time to find a solution to this problem. The former USSR and the US developed a one-step and two-step process for the conversion of ethanol into 1,3-BD, Scheme 1:

The mechanism is still open to much debate, but it is generally accepted that the following are key to its production. Initially, ethanol is dehydrogenated to afford acetaldehyde then an aldol process can occur between two acetaldehydes to form acetaldol. A Meerwein-Ponndorf-Verley (MPV) reduction step can then follow (also generating more acetaldehyde that can be used in the aldol step) to form 1,3-butadiol, subsequent dehydration forms butadiene. It is obviously possible that if the final two steps are reversed (Scheme 2), and acetaldol is dehydrated to form crotonaldehyde this can then undergo a MPV reduction to form crotylalcohol, which can be dehydrated giving 1,3-BD. Clearly, there are many undesired processes that can be occurring alongside the desired path forming a plethora of side products. By-products of this process include (but are not limited to) – acetone, propane/propane, butenes, diethyl ether, pentenes, hexenes, ethyl acetate, butanol and...
Acetaldehyde can also be thought of as a by-product, but this can be recycled into the feed and reused. The financial viability of the process will depend upon minimising the formation of by-products and thus maximising the overall process yield of 1,3-BD.

The success of this technology will depend on a cheap and readily available source of ethanol [4]. As with all processes relying on sustainable and renewable resources the new route must not impart undesired consequences on existing supply chains or impact on food provisions and security. With the arrival of algal biomass for bioethanol production then the possible environmental concerns can be circumvented [5]. The sustainable assessment (techno-economic) of the ethanol to 1,3-BD process has also been investigated by Patel and co-workers, who found that the bio-based route compares favourably with the traditional naphtha-based route [6,7].

**Review**

The conversion of ethanol to 1,3-BD is not a new process – with much work being carried out in the early part of the 20th Century. However, with the advent of a cheap and seemingly plentiful supply of crude-oil this research fell out of favour. In the 1980s it became more popular and in the 21st Century it is essential that we fully make use of this chemical technology. Thus, this short review will highlight the most recent examples of catalytic investigations into this process, the seminal initial work of Lebedev, Corson, Bhattacharyya and Ostromislenskiy has been the subject of review articles [8,9] and the reader is directed towards those and the primary literature for further details [10-21]. However, the scope of this review is to look at the most current literature in the area – the review focusses on sepiolite, MgO-SiO₂ and pure SiO₂ supported catalysts.
Post 1980’s Catalytic conversion of Ethanol to 1,3-BD

In 1981 Kitayama and co-workers investigated the catalytic activity of sepiolite \( [(\text{H}_2\text{O})_4(\text{OH})_4\text{Mg}_8\text{Si}_{12}\text{O}_{30}\cdot6\text{H}_2\text{O})] \) for the production of 1,3-BD [22]. Sepiolite was chosen as it is relatively easy to exchange the Mg(II) centres in the material with transition metals. In their study Mn(II) exchanged materials were investigated. The most promising results were observed at a temperature of 300°C with a 33.4% selectivity towards 1,3-BD, with 41.4% ethene. This comparison to just 2.4% 1,3-BD selectivity for the pure sepiolite material. The high quantity of ethene is a problem for this catalyst system and ethene should be avoided. Fripiat further investigated the substitution of sepiolite with either vanadyl or silver cations [23,24]. In the vanadyl case they observed only a modest selectivity to 1,3-BD when pure ethanol was used as the feed [23]. However, when the feed was rich in acetaldehyde a high selectivity, ca. 80%, to 1,3-BD was achieved. This was observed with and without the presence of the vandyl cation on the aluminated sepiolite. They proposed that the mechanism for this process involves the Prins reaction, Scheme 3:

For the silver aluminated sepiolite they observed that both the ethene and 1,3-BD selectivities increase linearly with total conversion [24]. They attributed this observation as support for the Prins mechanism. The Prins mechanism is an attractive approach as this can utilise the ethene by-product. However, since these initial studies the role of the Prins reaction has not been further investigated. The use of zeolites for this process is also not widely employed, this is presumably related to the high quantity of low-value ethene produced due to the acidic properties of most zeolites.

Magnesia-silica materials have been shown to be very effective catalysts (either pure supports or with added metal centres) for this process [25-27]. The magnesia offers basic sites and the silica acidic sites. It is hypothesised that the magnesia enhances the aldol and dehydrogenation steps of the mechanism, whereas the silica assists the dehydration steps on the mechanism. The preparation method and molar ratio of MgO-to-SiO2 have proved to be of critical importance in this field. For example, Ohnishi prepared MgO:SiO2 (with a 1:1 molar ratio) in three different ways and observed yields of 1,3-BD ranging from 2–42% \((T = 350°C, \text{Flow EtOH} = 6.5 \times 10^{-4} \text{ mol h}^{-1})\) [27]. It was observed that the most effective catalyst was prepared by the wet kneading of Mg(OH)\(_2\) [prepared by hydrolysis of Mg(NO\(_3\))\(_2\) with NH\(_3\)] with SiO\(_2\) [prepared from hydrolysis of Si(OEt)\(_4\) with HNO\(_3\), EtOH and NH\(_3\)]. Using MgCl\(_2\) as the source of MgO resulted in significantly poorer yields of 1,3-BD. Interestingly, with the addition of 0.1 wt% of either Na\(_2\)O or K\(_2\)O then the selectivity to 1,3-BD significantly increases. This maybe due to the reduction in the Brønsted acidity of the support, highly Brønsted acidic materials are well known to catalyse the conversion of EtOH to ethene and diethyl ether. MgO:SiO\(_2\) catalysts were also studied by Kvisle et al., the preparation of which was analogous to that of Ohnishi. Their catalytic tested were performed at 350°C, flow EtOH = 8 \(\mu\text{l h}^{-1}\), mass of catalyst 200 mg [26]. They observed that as the flow of EtOH increases then the conversion decreases and the selectivity to 1,3-BD also decreases. The selectivity to 1,3-BD could be increased by the addition of acetaldehyde to the feed or, interestingly, adding oxygen to the carrier gas. The addition of a second alcohol (methanol or isopropanol) did not alter the yield of 1,3-BD. It was thus hypothesised that the rate limiting step is not the hydrogen transfer step, which would be enhanced with the addition of other alcohols. Thus, the rate determining step occurs prior to the MPV step. It was also noted that the wet-kneaded catalyst significantly out performs (in terms of selectivity and longevity) a mechanical mixture of MgO and SiO\(_2\).
illustrating that there is a synergistic effect between the two oxides. Kitayama also prepared large surface area (up to 784 m²g⁻¹) nickel magnesium silicate materials for this process [28]. The catalysts were prepared with a 10 wt% NiO content, it was found that the optimum ratio of Si/Mg was 1.5, which afforded a 31% yield to 1,3-BD and a very low selectivity to ethene of 0.5%. It is believed that this ratio gives the optimum ratio of acid-to-base sites to enhance the selectivity to 1,3-BD. This was attributed to the decrease in the acid sites. Recently, Sels added transition metal centres to MgO:SiO₂ materials [25]. In their study the optimal ratio of Mg/Si was 2. Adding CuO, ZnO and Ag to this support (T = 350°C, EtOH concentration = 1.5 × 10⁴ppm) was seen to have a positive effect on selectivity. The preparation of these ternary systems was also investigated and it was found that the MgO:SiO₂ must be prepared first followed by addition of the transition metal. Recently, Jones and co-workers have prepared a series of bimetallic (ZnO/ZrO₂) supported MgO:SiO₂ catalysts [29]. For the undoped materials an optimum ratio of Mg/Si was 2:1 (T = 325°C, WHSV = 0.3 gEtOH/gcat·h, mass of catalyst = 1.0 g). However, this was not the case for the case for the bimetallic catalyst and the optimum ratio was 95:5 respectively. Interestingly, a small amount of SiO₂ is essential for this process, with 100% MgO the conversion was 5% with a 19% selectivity to 1,3-BD compared to 30% and 68% for the 95:5 material. There are earlier examples of the exploitation of MgO-SiO₂ by Niiyama who showed that an 85:15 ratio was optimal for the process [25]. They observed that the rate determining step was acetaldehyde formation which was catalysed by the basic sites in the material. They argue that it is important to control the acidity and basicity of the catalyst. [30] Takezawa have also studied the mechanism for the conversion of ethanol to acetaldehyde on MgO supports, and the formation of an ethoxde species on the MgO was observed via IR spectroscopy [31]. Supports such as SiO₂ and Al₂O₃ have also been utilised in the last three years for this process [32-35]. Jones has developed a series of bi- and tri-metallic catalysts for this reaction. They observed that with EtOH as the feed (LHSV = 1.0 h⁻¹, T = 375°C) a ZrO₂:ZnO:CuO:SiO₂ (1 wt% of each metal) catalyst was the most efficient giving a 67.4% selectivity to 1,3-BD [34]. Interestingly, as the pore diameter (40 – 60 – 150 Å) of the porous SiO₂ increased then the selectivity to 1,3-BD increased. This was attributed to the acidity of the support decreasing, thus reducing the quantity of ethene formed. It was hypothesised that the ZnO was active for the dehydrogenation and the ZrO₂ the aldol coupling step. Ordomskiy and co-workers have patented a series of gold, silver or copper and a metal oxide from magnesium, titanium, zirconium or tantalum deposited on SiO₂ [35]. They achieved up to an 82% yield of 1,3-BD (T = 325°C, WHSV = 0.3 gEtOH/gcat·h, 1:10 acetaldehyde to ethanol) with a Au-ZrO₂-SiO₂ material. They also observed high yields with Ag and CeO₂ containing catalysts. Carbon (coking) formation is also possible for this process and will block access to active sites and thus reduce activity. This will require regeneration of the catalyst and periodic production downtime. Ezinkwo has shown that if H₂O₂ is added as a “process initiator” then this is a potential solution to coking issues [33]. Without the addition of the process initiator the catalyst was only lasted for 48 hours on stream. However, with the addition of H₂O₂ the catalyst activity persisted for 120 hours without reduction, indicating the possibility of running this catalysis in a continuous operation. The catalyst used in this study was a ZnO/γ-Al₂O₃ system. In 2014, inspired by work in the early 20th Century, Chae and co-workers produced Ta₂O₅ supported on ordered mesoporous silica (SBA-15, MMS and KIT-6) [32]. They achieved conversions up to 47% and selectivity to 1,3-BD of 80% (T = 350°C, LHSV = 1 h⁻¹). Importantly, it was observed that catalysts based on these ordered materials showed higher tolerances towards coke and higher activity.

Conclusions
The conversion of ethanol to 1,3-butadiene is becoming increasingly important in the 21st Century. However, for the opportunities that this process offers to be fully realised the following problems still remain:

1) Increasing the selectivity, one of the most significant costs associated with the industrial scale-up of this process will be the separation of the 1,-BD from the by-products. Thus, the higher the selectivity then potentially the lower the cost of separation. The current literature clearly demonstrates that there needs to be the “goldilocks” condition of “just the right” balance of acid and base sites in the catalyst. It is anticipated that selectivities in excess of 70% will be required. This is a very challenging target given the plethora of possible side reactions.

2) The mechanism has been much debated in the literature. Future studies are required to fully ascertain the respective rates of the Prins mechanism compared to the aldol/hydrogen transfer process. These could entail isotopic labelling studies and/or computational analysis to fully probe which steps of the mechanism are occurring on which active site of the catalyst.

3) Reactor engineering, once the optimum catalyst has been developed studies are urgently required to develop reactors for this process – the relative merits of a fixed bed or fluidised bed reacts still needs to be investigated further [10]. Furthermore,
work needs to be directed towards complications with coking [33].

Competing interests
The author declares that they have no competing interests.

Acknowledgments
MDJ wishes to thank the University of Bath, MDJ would also like to thank the collaborators of references 29 and 34.

Received: 21 May 2014 Accepted: 8 August 2014
Published online: 10 September 2014

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Cite this article as: Jones: Catalytic transformation of ethanol into 1,3-butadiene. Chemistry Central Journal 2014; 8:53.