Herein we report the first example of the catalytic aerobic partial oxidation of allyl ether to its acrylate ester derivative. Many partial oxidations often need an expensive oxidant such as peroxides or other species to drive such reactions. In addition, selective generation of esters using porous catalysts has been elusive. This reaction is catalyzed by a Li ion promoted mesoporous manganese oxide (meso-Mn$_2$O$_3$) under mild conditions with no precious metals, a reusable heterogeneous catalyst, and easy isolation. This process is very attractive for the oxidation of allyl ethers. We report on the catalytic activity, selectivity, and scope of the reaction. In the best cases presented, almost complete conversion of allyl ether with near complete chemo-selectivity towards acrylate ester derivatives is observed. Based on results from controlled experiments, we propose a possible reaction mechanism for the case in which N-hydroxyphthalimide (NHPI) is used in combination with trichloroacetonitrile (CCl$_3$CN).
Acrylics and acrylates\textsuperscript{1–3} have become important building blocks for the chemical industry today. Acrylates are key ingredients for producing dyes, glues, papers, adhesives, binders, paints, and thickeners\textsuperscript{4,5}. The industrial source of acrylics and acrylates is propylene that is produced as either the by-product of fluid catalytic cracking (FCC)\textsuperscript{6}, steam cracking, or by propane dehydrogenation\textsuperscript{7}. Propene or propylene can be subsequently oxidized to acrylates via the Mitsubishi process\textsuperscript{8} or propene ammoxidation\textsuperscript{9–11} using mixed metal oxides. These processes involve the formation of acrylonitrile intermediates that are converted into acrylates, acrylamides, and acrylic acids depending on the reaction condition employed\textsuperscript{12}. Typically these oxidations are run around 450 °C, where formation of ethylene, HCN, acetonitrile, N\textsubscript{2}O, and CO\textsubscript{2} by-products are unavoidable.

Recently, several homogeneous processes have been recognized for efficiently oxidizing allylic C–H bonds\textsuperscript{13–15}. Peroxides and N-hydroxides were used for synthesizing various enones via allylic C–H oxidation\textsuperscript{16}. However, these methodologies primarily focus on cyclic unsaturated molecules. Crystalline manganese oxide based materials (e.g., K-OMS-2) are used for various catalytic transformations including the selective oxidation of alcohols to aldehydes\textsuperscript{16}, hydrocarbons to alcohols and ketones\textsuperscript{17}, styrenes to styrene oxides\textsuperscript{18}, alcohols to amides\textsuperscript{19}, amines to imines\textsuperscript{20}, some oxidative coupling reactions, such as alkynes to di-ynes\textsuperscript{21}, alkynylsilanes\textsuperscript{22}, anilines to azobenzenes\textsuperscript{23}, and dehydrogenative oxidation reaction of alkanes\textsuperscript{24}.

These mixed valent manganese oxide materials have numerous structural forms with high thermal stability\textsuperscript{25}. The reduction of Mn\textsuperscript{4+}/3+ to Mn\textsuperscript{3+}/2+ releases labile lattice oxygen atoms, which are responsible for their catalytic performance\textsuperscript{21,25}. These lattice oxygens act as basic sites for H abstraction during the reaction. This eliminates the requirement of using a base with mesoporous manganese oxides for oxidative dehydrogenations\textsuperscript{26}. Moreover, the presence of alkali and alkaline earth metal ions in the manganese oxide lattice enhances catalytic activity. Basicity increases due to the attraction of lattice oxygens toward these electropositive ions. We therefore introduced these types of ions in the allylic C–H oxidation reactions. These mesoporous manganese oxide materials, developed by an inverse-micelle method, were selected for their comparatively high surface areas compared with traditional mesoporous transition metal oxides\textsuperscript{27}.

In this paper, acrylates have been directly produced via an aerobic catalytic oxidation process. Allyl ethers were used in this low-temperature process with lithium ion impregnated mesoporous manganese oxides as heterogeneous catalysts. An advantage of these catalysts is that manganese oxides are abundant and inexpensive materials. Conversions of diallyl ether as high as 95% with selectivities to allyl acrylate greater than 99% were achieved via optimization of the reaction conditions. Such studies suggest that NHPI–air, hydrogen peroxides, and CCl\textsubscript{3}CN are essential in controlling activity, selectivity, and stability of these catalysts. Control experiments were done to determine the interrelationships of these critical factors in these oxidation reactions. A plausible mechanism is proposed based on mechanistic studies. Such studies suggest that radical intermediate formation, followed by successive hydroxylation and oxidation are significant for this reaction. Mobility of lattice oxygen in the meso-Mn\textsubscript{3}O\textsubscript{4} catalysts and CCl\textsubscript{3}CN promoted PINO radicals are the significant requirements in this catalytic process.

**Results**

**Reaction conditions.** The interdependence and synergistic effect of the reaction parameters spurred us to evaluate and compare reaction conditions (Supplementary Figs. 1 and 2). Reaction kinetics under different set of conditions were investigated. NHPI–air–CCl\textsubscript{3}CN emerged with the highest rate constant of 2.82 min\textsuperscript{−1}, while NHPI–N\textsubscript{2}–CCl\textsubscript{3}CN displayed the slowest rate constant of 0.3 min\textsuperscript{−1} (Supplementary Table 6, entry 1 and 4). Moreover, reaction under NHPI–air with no CCl\textsubscript{3}CN displayed a rate constant of 0.57 min\textsuperscript{−1}, which was slightly lower than 0.71 min\textsuperscript{−1} of the TBHP–CCl\textsubscript{3}CN system (Supplementary Table 6, entry 2 and 3). The four fold increase in activity for NHPI–air–CCl\textsubscript{3}CN than the TBHP–CCl\textsubscript{3}CN system proved its superiority. A ninefold decrease in activity upon replacing air with nitrogen and a fivefold decrease in the absence of CCl\textsubscript{3}CN, proved their significance. These studies helped evaluate the importance of reaction parameters. According to the results obtained, reaction conditions which increase the reaction rate are ranked as follows: NHPI–O\textsubscript{2}–CCl\textsubscript{3}CN > TBHP–CCl\textsubscript{3}CN > NHPI–O\textsubscript{2}– no CCl\textsubscript{3}CN > NHPI–N\textsubscript{2}–CCl\textsubscript{3}CN. In all cases, these kinetic experiments revealed a first-order rate dependence with respect to the starting material allyl ether. Further characterization was done with X-ray powder diffraction and BET surface area measurements (Supplementary Fig. 6). Various reactants and products were analyzed with nuclear magnetic resonance (NMR), See Supplementary Note 1.

**Substrate scope.** The substrate scope and limitations were then explored using structurally different ethers, with each possessing at least one allylic CH\textsubscript{2} group. Under optimized reaction conditions, diallyl ether and cyclic dihydrofuran (Table 1, entry 1 and 3) were transformed to their corresponding oxidized products exclusively with 92 and 99% yields, respectively. On the other hand, allyl acetate and allyl glycidyl ether (Table 1, entry 2 and 4) were transformed to their oxidized counterpart with 60 and 66% yields, respectively. However, a poor yield of 12% was obtained for allyl phenyl ether (Table 1, entry 5). Hyperconjugation and the loss of a proton from the α-carbon was easier in the case of allyl ether than that of phenyl allyl ether (B. E. = 67.2 ± 0.5, and 81.84 ± 0.5 kcal/mol, respectively, Table 1). This resulted in better stability and poorer reactivity of the phenyl allyl ether than the allyl ether. To investigate the reason for selective formation of monoesters over the anhydride, density functional theory (DFT) calculations were conducted, using the CBS–QB3 basis set. DFT results show that C–H bond dissociation energy of allylic C–H of diallyl ether is 80.8 kcal/mol, which significantly increases to 82.7 kcal/mol for the allyl acrylate ester C–H. A smaller increase in bond dissociation energy was observed when comparing reactants and products in Table 1, entry 3. Perhaps the cyclic nature of the substrate binding to the catalyst surface further limited the oxidation of the lactone to the corresponding anhydride. In Table 1, entry 4 the glycidyl C–H bond in the reactant is a competing site for oxidation with a calculated BDE of 91.2 kcal/mol. Upon oxidation of the allylic C–H bond the glycidyl C–H bond becomes stronger at 95.7 kcal/mol. Probably for similar reasons, no activation of α–C–H bonds was noted in 1-butene and ethyl substituted allyl ethers (Table 1, entry 6 and 7).

**Stability and reusability.** To determine if the observed catalysis is a result of leached active sites, the solid catalyst was separated from the reaction mixture by hot filtration after 45 min (about 25% conversion). The resulting filtrate was then subjected to reaction conditions for an additional 315 min. Aliquots were collected successively after each hour and were analyzed with GC-MS to track the progress of the reaction. No further improvement in GC yield was observed after the first 2 h (Supplementary Fig. 3), indicating no further conversion of substrate in the absence of a catalyst. The same filtrate was also analyzed by inductively coupled plasma mass spectrometry (ICP-MS), which
indicated the presence of a very low amount (165 ppm) of Manganese (Mn).

To verify reusability, the catalyst was retrieved (> 90%) after the reaction by simple filtration and washed with excess acetonitrile and methanol. The catalyst was reactivated prior to reuse at 250 °C for 30 min under air to remove any adsorbed substrates. As evident from Supplementary Fig. 4, the catalyst was successfully reused for three more cycles with less than a 50% drop in activity. The decrease in activity is probably due to metal poisoning (MnIII), caused by the strong coordination of different radicals with the metal centers. Therefore, our catalyst can be considered as truly heterogeneous, stable, and moderately reusable.

Discussion
Based on the above-mentioned observations, we propose mechanisms by both TBHP–CCl3CN and NHPI–air–CCl3CN catalysts. The role of manganese species during the reaction can be correlated with the work reported by Ishii et al. using electron paramagnetic resonance (EPR) measurements. To further understand our systems, EPR data for the acetonitrile solution of NHPI with and without meso-Mn2O3 under atmospheric condition were compared. There was a decrease in the EPR signal, which could be due to the formation of an extra radical (possibly PINO) in the former reaction at g = 2.0073 (Supplementary Fig. 5). This suggest the formation of a superoxomanganese (IV) or μ-peroxomanganese (IV) complex between MnIII and O2. In both cases, a series of steps contributed to the formation of monoesters from the allyl ethers. Manganese oxides (MnIV–oxide, 1) have been reported to react with molecular oxygen to form MnIV–peroxide radicals (2) (Fig. 1). These are considered to be very reactive and are expected to deprotonate the NHPI molecule to form a PINO radical and a Mn IV–hydrogen peroxide (3) species. A competition between PINO radical and MnIV–peroxide radical (2, is expected during the deprotonation of allylic hydrogens (5). The corresponding radical (6) is believed to couple with an oxygen molecule and MnII species to form an intermediate 7 that upon reacting with a molecule of NHPI (4) produced a MnIII–PINO (11) and allyl ether-hydrogen peroxide (8) species. The latter went through further rearrangements with the loss of an oxygen moiety to generate the desired allyl acrylate (10), whereas the MnIII–PINO (11) regenerated the TBHP and MnIV species.
The mechanism in the presence of TBHP is expected to follow a similar mechanism to that of the NHPI catalyzed pathway (Fig. 2). A fourfold decrease in the rate constant (i.e., from 2.82 min\(^{-1}\) for NHPI–O\(_2\)–CCl\(_3\)CN to the 0.71 min\(^{-1}\) by TBHP–CCl\(_3\)CN-based reaction) discouraged us from using this catalytic system. Moreover, we have independently determined that Mn\(_2\)O\(_3\) is an extremely potent material for decomposing TBHP into t-BuOH and O\(_2\). Since O\(_2\) is highly abundant and cheap, we focused our work using O\(_2\) as an oxidant, not only due to the relatively poor rates of TBHP–CCl\(_3\)CN system.

In both cases, probably an intermediate 6 type radical was oxidized to form an α-hydroxy ether (9). The transformation of intermediate 8 to 9, probably resulted in the release of oxygen, which are the terminal oxidants of the catalytic process (Fig. 1). Though the fate of this oxygen is known, their method of interaction in the oxidation process is still unknown. An α-hydroperoxide intermediate is likely to be formed, which, however, is very difficult to trap in rapid oxidations over manganese oxide.  

In summary, we have developed a low-temperature, heterogeneous catalytic aerobic oxidation procedure to form acrylates directly from allyl ethers using lithium ion impregnated mesoporous manganese oxide material. Mn being an earth-abundant element and O\(_2\) being the most abundant oxidant makes this catalytic reaction attractive. With diallyl ether we achieved a conversion of 95% to allyl acrylate with a selectivity of > 99% after extensive optimization of the reaction conditions. This revealed the crucial role of NHPI–air, hydrogen peroxides, and CCl\(_3\)CN in promoting the activity of the catalyst. The interdependence of these promoting agents along with the catalyst in driving the catalytic process was interpreted by control experiments that led to a plausible mechanism. Mechanistic studies invoked the possibility of radical intermediate formation, followed by successive hydroxylation and oxidation. The presence of the CCl\(_3\)CN promoted PINO radical and labile lattice oxygen of the meso-Mn\(_2\)O\(_3\) material are the most important features of this catalytic process.

**Methods**

**Catalysis.** In this paper, we have developed a new catalytic approach to the mild aerobic partial oxidations of diallyl ether, see Fig. 2. Diallyl ether was chosen as a model substrate that could potentially suffer from (i) a double-bond shift and (ii) oxidation of the olefin to aldehyde or epoxide prior to the activation of allylic C–H bond. We demonstrate the aerobic oxidation of allyl ethers to corresponding acrylate esters using various cation doped mesoporous manganese oxide materials. The absence of any precious metal, easy isolation of the products, and the use of a reusable heterogeneous catalyst in the absence of acidic or alkaline media makes this process attractive for the oxidation of allyl ethers.

We initiated the oxidation of allyl ether in air as a model reaction to determine the optimal conditions. In our early screening attempts, we evaluated the impact of solvent on this oxidation. The results of our screening studies are presented in Supplementary Table 1 of the Supplementary information. Interestingly, no trace of the oxidized product was detected until 1.2 eqv. NHPI or 2 eqv. meso-Li-Mn\(_2\)O\(_3\) was introduced (Supplementary Table 1, entry 1–4). Unsatisfactory conversion of allyl ether, even after using TBHP led us to employ a radical promoter. In 2010, Kamijo et al. successfully discovered the promotional activity of trichloroacetonitrile (CCl\(_3\)CN) for m-CPBA (metachloroperbenzoic acid) in either oxidations. The high electrophilicity of the CCl\(_3\)CN most likely promoted the homolytic cleavage of O–O bonds of mCPBA by forming a highly unstable peroxysimate adduct. The utilization of this hydroperoxide promoter (CCl\(_3\)CN) as a solvent increased the...
product formation in our system to as high as 82% (Supplementary Table 1, entry 5). Further optimization of the amount of this promoter (CCl3CN) displayed similar results (81% yield, Supplementary Table 1, entry 6), indicating no dependence of reactivity on the amount of CCl3CN used. The role of CCl3CN was further validated by performing a solvent-free reaction with only 15% of the allyl ether converted (Supplementary Table 1, entry 7). The promising results of solvent optimization, coupled with the other mechanisms that support the higher conversions than those for the Mn2+-rich and catalyst-free systems (Supplementary Table 2, entry 6–9) with at most 20% yield observed for the other supports. Better accessibility of the labile lattice oxygens and comparatively lower binding energy of metal–oxygen bonds resulted in higher yields. Reactions using Mn2+-enriched species (such as Mn2O3 and Mn(OAc)3) showed better conversions and no other oxygen supports. None were found to be as active as meso-MnO2 (81% yield) (Supplementary Table 2, entry 1–4) with the solvent, were excluded (Supplementary Table 4, compare entry 7). This confirmed that lattice oxygens of the catalyst were the terminal oxidants.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

References

1. Zhang, Z., Qu, Y., Wang, S. & Wang, J. Catalytic performance and characterization of silica supported sodium phosphates for the dehydration of methyl lactate to methyl acrylate and acetic acid. *Ind. Eng. Chem. Res.* 48, 9083–9089 (2009).
2. Lee, J. M. et al. Efficient dehydration of methyl lactate to acetic acid using Ca 3(PO4)2-SiO2 catalyst. *Catal. Commun.* 11, 1176–1180 (2010).
3. Zhang, J. et al. Efficient acrylic acid production through bio lactic acid dehydration over Na2Zelite modified by alkali phosphates. *ACS Catal.* 1, 32–41 (2011).
4. Jamróz, M. H., Jamróz, M. E., Rode, J. E., Bednarek, E. & Dobrowolski, J. Interpretation of vibrational and NMR spectra of allyl acrylate: An evidence for several conformers. *Vib. Spectros.* 50, 231–244 (2009).
5. Agarwal, Y. K., Kaushik, S. D. & Kumar, P. C. Synthesis and rheological studies of methacrylic acid-ethyl acrylate-ally methacrylate terpolymers. *J. Macromol. Sci. Part A* 44, 877–880 (2007).
6. Collins, J. P. et al. Catalytic dehydrogenation of propane in hydrogen permselective membrane reactors. *Ind. Eng. Chem. Res.* 35, 4398–4405 (1996).
7. Clarke, A. F. Process of making allyl acrylates. *U. S. Patent 2,464,364, 1–3 (1943).
8. Korovchenko, P. et al. M1 to M2 phase transformation and phase cooperation in bulk mixed metal Mo-V-M-O (M = Te, Nb) catalysts for selective ammoxidation of propane. *Top. Catal.* 50, 43–51 (2008).
9. Guliants, V. V. et al. Role of surface Te, Nb, and Sb oxides in propane ammoxidation to acrylonitrile with an alumina-supported Sb-V-O catalyst. *Ind. Eng. Chem. Res.* 45, 4537–4543 (2006).
10. Amakawa, K. et al. Multifunctionality of crystalline Mo(VTeNb) M1 oxide catalysts in selective oxidation of propane and benzyl alcohol. *ACS Catal.* 3, 1103–1113 (2013).
11. Brzädl, J. F. in *Ullmann’s Encyclopedia of Industrial Chemistry* (Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, 2000).
12. Ishii, Y., Sakaguchi, S. & Iwawama, T. Innovation of hydrocarbon oxidation with molecular oxygen and related reactions. *Adv. Synth. Catal.* 343, 393–427 (2001).
13. Horn, E. J. et al. Scalable and sustainable electrochemical allylic C–H oxidation. *Nature* 533, 77–81 (2016).
14. Hruszkewycz, D. P., Miles, K. C., Thiel, O. R. & Stahl, S. S. Co/NHPI-mediated aerobic oxygenation of benzylic C–H bonds in pharmaceutically relevant molecules. *Chem. Sci.* 8, 1282–1287 (2017).
15. Son, Y., Makwana, V. D., Howell, A. R. & Subh, S. L. Efficient, catalytic, aerobic oxidation of alcohols with octahedral molecular sieves. *Angew. Chem.* 113, 4110–4113 (2001).
16. Ophembe, N. N., Son, Y.-C., Srisukdakumar, T. & Suib, S. L. Kinetics and mechanism of 9H-fluorene oxidation catalyzed by manganese oxide octahedral molecular sieves. *ChemSusChem* 1, 182–185 (2008).
17. Ghosh, R. & Taneja, S. Role of manganese oxide octahedral molecular sieves in styrene epoxidation. *J. Phys. Chem. B* 110, 7592–7599 (2006).
18. Yamaguchi, K., Kobayashi, H., Oishi, T. & Mizuno, N. Catalytic amide synthesis heterogeneously catalyzed synthesis of primary amines directly from primary alcohols and aqueous ammonia. *Angew. Chem.* 124, 559–562 (2012).
19. Biswas, S. et al. Aerobic oxidation of amines to imines by cesium-promoted mesoporous manganese oxide. *ACS Catal.* 5, 4394–4403 (2015).
20. Biswas, S. et al. Mesoporous copper / manganese oxide-catalyzed coupling of alkynes: evidence for synergistic cooperative catalysis. *ACS Catal.* 6, 5069–5080 (2016).
21. Yamaguchi, K., Wang, Y., Oishi, T., Kuroda, Y. & Mizuno, N. Heterogeneously catalyzed aerobic cross-dehydrogenative coupling of
terminal alkynes and monohydrosilanes by gold supported on oms-2. Angew. Chem. Ed. 52, 5627–5630 (2013).

23. Dutta, B. et al. Mesoporous manganese oxide catalyzed aerobic oxidative coupling of anilines to aromatic azo compounds. Angew. Chem. 55, 2171–2175 (2016).

24. Jin, L. et al. Studies on dehydrogenation of ethane in the presence of co2 over octahedral molecular sieve (OMS-2) catalysts. Chem. Cat. Chem. 1, 441–444 (2009).

25. Sui, S. L. Porous manganese oxide octahedral molecular sieves and octahedral layered materials. Acc. Chem. Res. 41, 479–487 (2008).

26. Biswas, S. et al. Ion induced promotion of activity enhancement of mesoporous manganese oxides for aerobic oxidation reactions. Appl. Catal. B Environ. 165, 731–741 (2015).

27. Poyraz, A. S., Kuo, C.-H., Biswas, S., King’omodu, C. K. & Sui, S. L. A general approach to crystalline and monomodal pore size mesoporous materials. Nat. Commun. 4, 1–10 (2013).

28. Ishii, Y. & Sakaguchi, S. A new strategy for alkane oxidation with O2 using N-hydroxyphthalimide (NHPI) as a radical catalyst. Catal. Surv. Jpn. 3, 27–35 (1999).

29. Biswas, S. et al. Heterogeneous catalytic oxidation of amides to imides by manganese oxides. Sci. Rep. 8, 13649 (2018).

30. Lin, S.-S. & Gurol, M. D. Catalytic decomposition of hydrogen peroxide on iron oxide: kinetics, mechanism, and implications. Environ. Sci. Technol. 32, 1417–1423 (1998).

31. Kamijo, S., Matsumura, S. & Inoue, M. CCISCN: a crucial promoter of MCPBA-mediated direct ether oxidation. Org. Lett. 12, 4195–4197 (2010).

32. Helwani, Z., Othman, M. R., Aziz, N., Kim, J. & Fernando, W. J. N. Solid heterogeneous catalysts for transesterification of triglycerides with methanol: A review. Appl. Catal. A: Gen. 363, 1–10 (2009).

33. Sheldon, R. A. & Arends, I. W. C. E. Organocatalytic oxidations mediated by nitronyl radicals. Adv. Synth. Catal. 346, 1051–1071 (2004).

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Author contributions
B.D., P.N., and S.L.S. designed the experiments. B.D. synthesized the materials and conducted the catalytic reactions. R.C. performed some initial experiments and isolated the products. S.R. and T.D.S. performed computational studies and provided valuable insights. L.A. performed the electron paramagnetic resonance (EPR) experiments. S.L.S. and P.N. were the project coordinators. B.D., P.N., and S.L.S. prepared the paper. All authors have read and approved the final paper.

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