Mechanochemical Nitration of Organic Compounds

Statement of Need WP-19-2748-F1

SEMS Project Number: WP19-1383

Final Outbrief

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18. NUMBER

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Mechanochemical nitration of aromatic precursors

 Previous SERDP-funded work established possibility of mechanochemical nitration:

$$C_7H_8 + NaNO_3 \xrightarrow{MoO_3} C_7H_7(NO_2) + NaOH$$

Lagoviyer, O.S., Krishtopa, L., Schoenitz, M., Trivedi, N.J., Dreizin, E.L. *Journal of Energetic Materials* pp. 1-11 (2017) Lagoviyer, O.S., Schoenitz, M., Dreizin, E.L. *Journal of Materials Science* 53, pp. 13690-13700 (2018)

- What processes govern this reaction?
- → Conduct experiments with systematically varied reactants:
 - Different catalysts, nitrating toluene
 - Nitrating different precursors
 - Different nitrates as nitronium sources

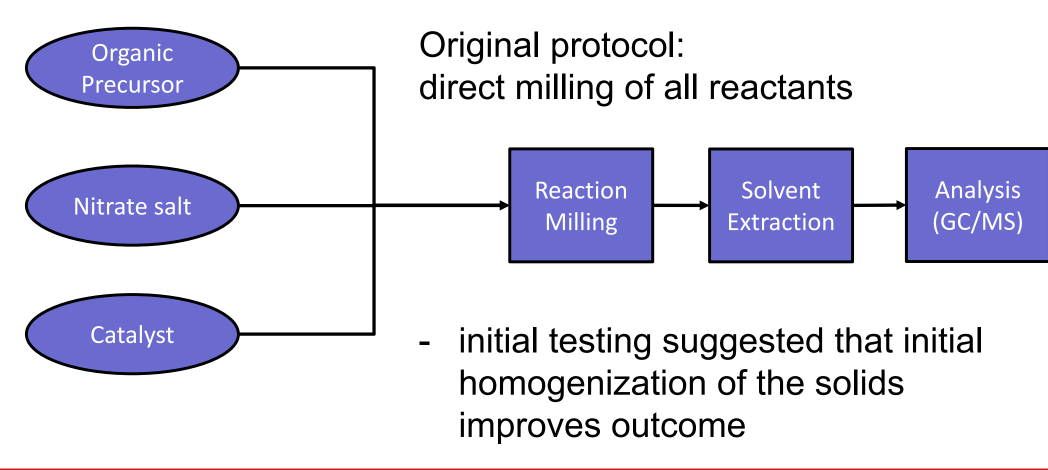


Equipment

- Retsch PM400-MA planetary mill
 - -4×500 mL milling jars
 - Custom temperature control
- General reactant amounts
 - aromatic precursor: 0.5 mL (0.25 mL 8 mL)
 - **nitrate salt**: 1.67 g (0.4 g 16 g)
 - metal oxide catalyst: 41.63 g
- Exploratory experiments used batch reactor
- Scale up expected to rely on attrition milling enabling semi-batch or continuous synthesis

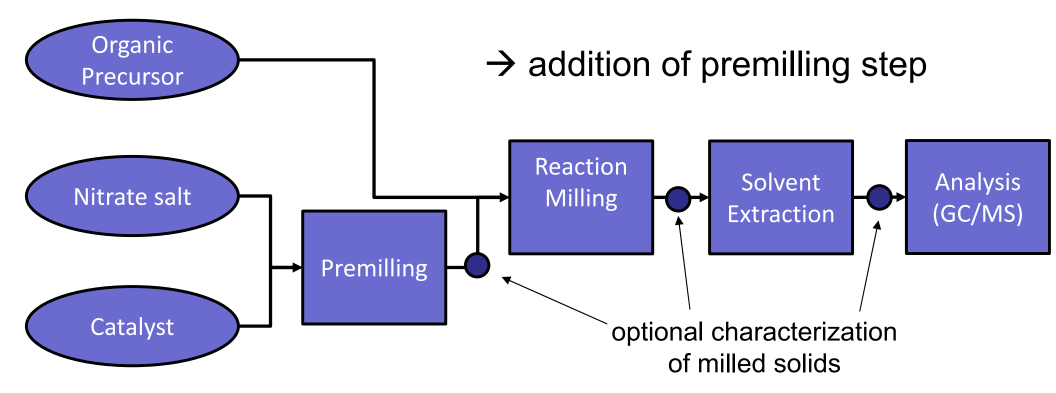


Experimental Protocol





Experimental Protocol





Part 1. Effect of <u>catalyst</u> on nitration of toluene

- Different inorganic oxide catalysts: Mo, W, V, Mn, Ti, Fe, Al, Bi
 - range of acid strengths (Lewis and Brønsted) of surface sites
- Toluene precursor
- NaNO₃ nitrate salt/nitronium source

Table 1
Materials tested as catalysts for mechanochemical nitration of toluene.

Material	Supplier and nominal purity	Relative Acidity, [38]	Type of acidity [37]			
Molybdenum oxide, MoO ₃	Alfa Aesar, 99.95 %	5.35	Medium Bronsted + Strong Lewis sites			
Tungsten oxide, WO ₃	Alfa Aesar, 99.95 %	5.05				
Vanadium oxide, V ₂ O ₅	Acros organics, 99.6 %	4.54				
Manganese oxide, MnO ₂	Alfa Aesar, 99.9 %	3.65	Not specified			
γ- alumina, Al ₂ O ₃	Alfa Aesar, 99.9 %	2.28	Strong Lewis sites ^a			
Titanium oxide, TiO ₂	Alfa Aesar, 99.9 %	3.05	Medium Lewis sites			
Iron oxide, Fe ₂ O ₃	Alfa Aesar, 99.5 %	2.5				
Bismuth oxide, Bi ₂ O ₃	Alfa Aesar, 99 %		Medium Lewis + Basic sites ^b			
Zeolite HZSM-5, Si/Al 38	ACS chemicals		Bronsted + Lewis sites ^b			
Zeolite HZSM-5, Si/Al 360						

^a Alumina phase was not specified in ref. [38].

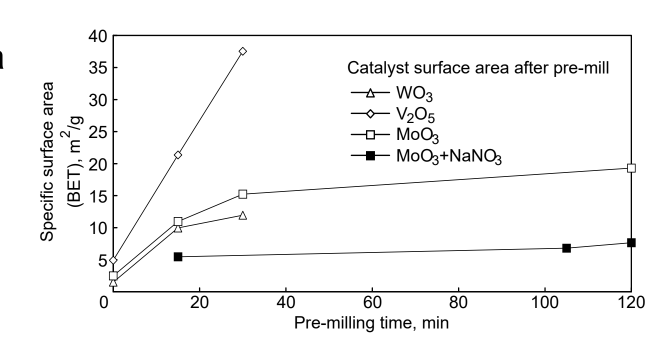
b Acidity values not stated in literature.



Evaluation of premilling: Development of catalyst & nitrate

BET specific surface area

- increases with time
- saturated after 2 hours
- absolute surface area determined by oxide

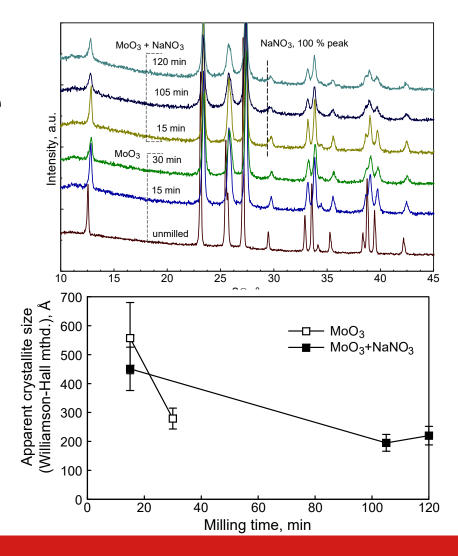




Evaluation of premilling: Development of catalyst & nitrate

XRD: <u>nitrate</u> no longer crystalline enough to be detected after premilling

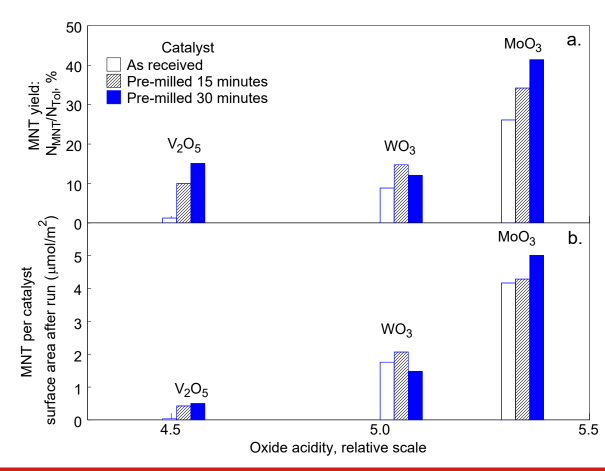
Catalyst also shows signs of decreased crystallinity





Overall results: Mononitrotoluene yields

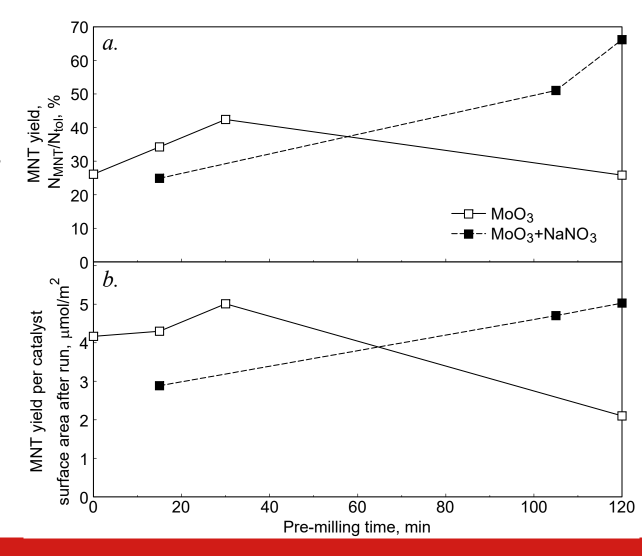
- Only Mo, W and V oxides showed quantifiable yields
- Rough correlation with oxide acidity, apparent when normalized by catalyst surface area
- Premilling catalyst with nitrate affects all catalysts
 - Highest yield with MoO₃





Pre-milling time and distribution of NaNO₃

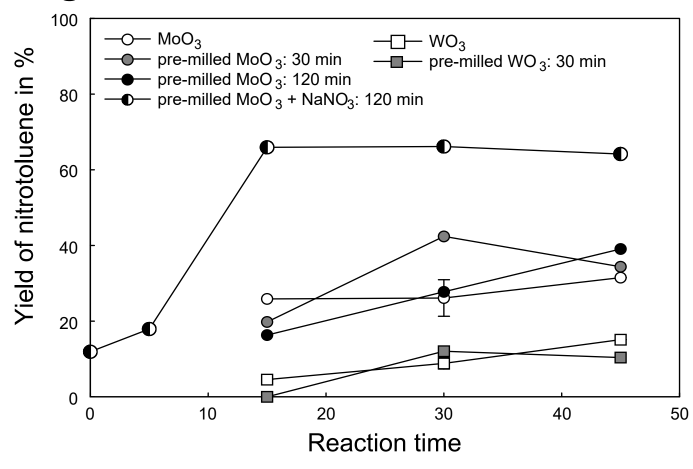
- Effect of MoO₃ pre-milling time is minor
 - Longer times may be undesired
- Distribution of NaNO₃ is important
 - Longer pre-milling MoO₃ with
 NaNO₃ leads to a better
 distribution
 - Higher yield





Effect of reaction milling time

- \square MoO₃, WO₃: premilling the catalyst is not important
 - A weak trend of yield increasing with the reaction time
- ■MoO₃+NaNO₃: yield reaches a **plateau after 15 minutes**
 - Nearly complete conversion





Findings guiding further work:

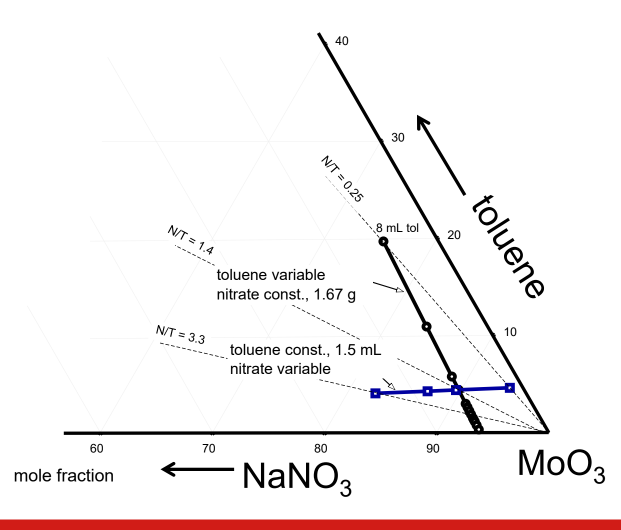
- MoO₃ most suitable catalyst, to be used for further work
- Premilling for 2 h gives most consistently high yields
 - becomes part of standard protocol



Systematic variation of reactant proportions

2 experimental series, varying nitrate/toluene reactant ratios

- toluene = const.
- nitrate = const.
 - □ premilled/not premilled

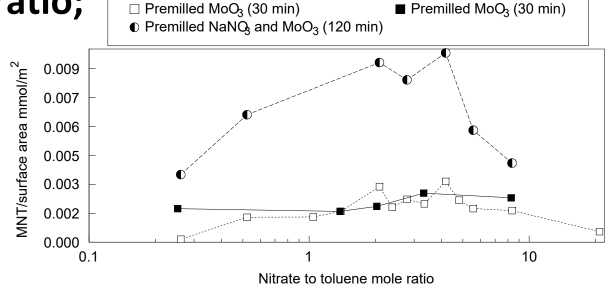




Effect of NaNO₃/toluene ratio;

MNT yield

- Varied toluene:
 - A clear peak in yield for a range of NaNO₃/toluene ratios
 - Better seen for pre-milled NaNO₃+MoO₃
 - Corresponds to a toluene spread over catalyst in a very thin layer
 - Close to monolayer
- Varied NaNO₃
 - No effect
- Toluene amount (layer thickness) matters



Toluene varied; 1.67 g NaNO₃



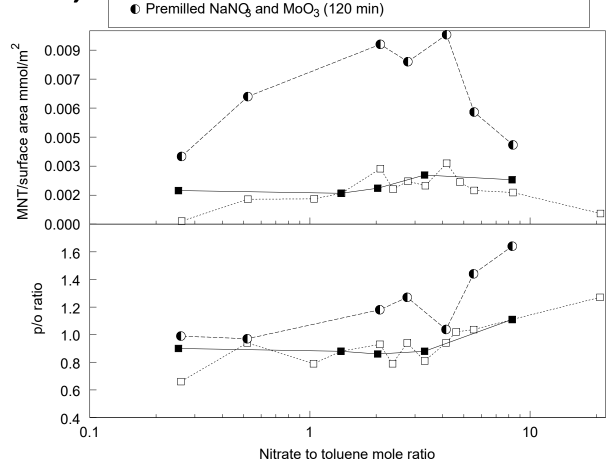
NaNO₃ varied; 1.5 ml toluene

Effect of NaNO₃/toluene ratio;

MNT isomer ratio

 Higher p/o ratios for greater yield

Unusually high p/o
ratio for runs limited
by toluene amount



Toluene varied; 1.67 g NaNO₃

☐ Premilled MoO₃ (30 min)



Part 1: Catalyst

NaNO₃ varied; 1.5 ml toluene

■ Premilled MoO₃ (30 min)

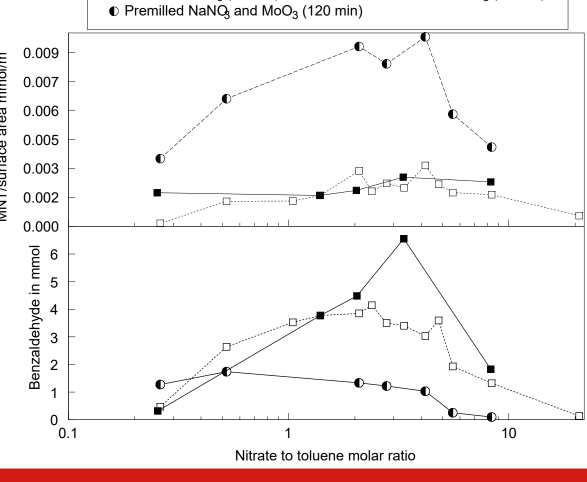
Effect of NaNO₃/toluene ratio;

Oxidized byproducts

 Benzaldehyde: main byproduct

 Production does not directly correlate with MNT yield

 Lowest benzaldehyde production for greatest yield



Toluene varied; 1.67 g NaNO₃

☐ Premilled MoO₃ (30 min)



Part 1: Catalyst

NaNO₃ varied; 1.5 ml toluene

■ Premilled MoO₃ (30 min)

Discussion: nitration occurs on surface of catalyst

- Reaction was not detected
 - for catalysts with low acidity
 - for zeolites: interior of the pores where the reaction might be taking place is poorly accessible
- For all three successful catalysts, MNT yield scales
 - with acidity
 - With surface area of the catalyst
- Pre-milling the catalyst powder has a limited effect on nitration
 - The catalyst is milled again during the reaction
- The effect of pre-milling is stronger when both catalyst (MoO₃) and the solid reactant (NaNO₃) are pre-milled together



Complete toluene conversion to MNT observed

- High yield with short 15-min reaction time
 - 70% conversion measured
 - Close to 20% of toluene could have evaporated
 - Close to 10 % could be unaccounted for
- High yield is only observed when NaNO₃ was pre-milled with the catalyst for 120 min.
 - Homogenizing NaNO₃ and the catalyst was the rate-limiting process when the powders were not pre-milled together
- For pre-milled MoO₃ + NaNO₃, mechanochemical reaction occurs fast, but some time is still required
 - Toluene must be distributed over the catalyst surface
 - New catalyst surface becomes continuously available during milling



Mechanism of mechanochemical nitration

- Toluene is nitrated reacting with nitronium complexes attached to acid sites
 - para MNT isomer is preferentially produced
 - Consistent with the present high p/o ratios
- Both yield of MNT and selectivity are reduced simultaneously when the amounts of toluene are increased
 - Excess of toluene may be blocking acid sites from nitronium
 - Formation of nitronium involves interaction of dissolved NaNO₃ with Bronsted acid sites
 - blocked acid sites may lead to a less effective production of nitronium slowing down nitration



Part 1: Summary

- Mechanochemical nitration of toluene occurs on surface of catalyst
 - High acidity, high surface area lead to high yield
- Complete conversion of toluene achieved in short 15-min reaction time
 - Pre-milling catalyst and nitrate prior to reaction accelerates mechanochemical nitration
- Toluene is nitrated reacting with nitronium complexes attached to acid sites
 - High p/o ratio
 - Low byproducts
- Both yield of MNT and selectivity are high for small amounts of toluene (close to monolayer on catalyst surface)



Part 2. Nitrating different aromatic precursors

- Various organic precursors
 with wide range of aromatic
 activation
 and other properties
- MoO₃ as oxide catalyst
- NaNO₃ nitrate salt/nitronium source

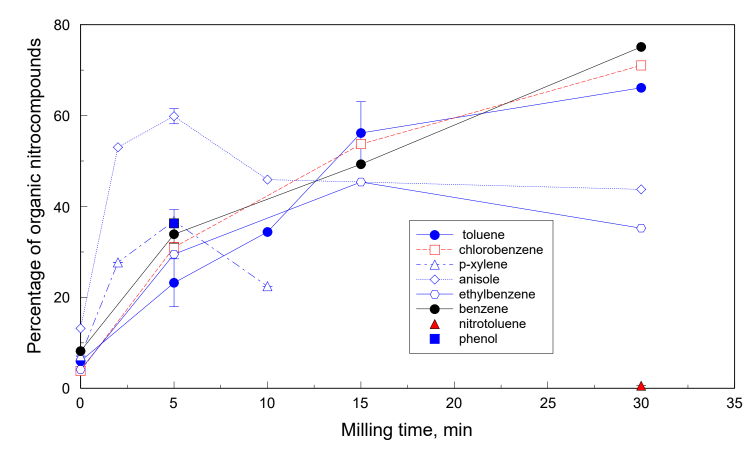
Precursor	Functional group	Relative rate log(k/k ₀) [1]
4-Nitrotoluene	NO ₂	-3.72
Chlorobenzene	Cl	-0.66
Benzene	Н	0
Ethyl benzene	C ₂ H ₅	0.9
Toluene	CH ₃	1.02
p-Xylene	(CH ₃) ₂	2.04
Phenol	ОН	2.22
Anisole	OCH ₃	4.68

Observed relative rates in liquid phase nitration using H_2SO_4 and HNO_3 at 25°C [1] C. Hansch, A. Leo, R.W. Taft, Chemical Reviews, 91 (1991) 165-195



Reaction rate – substituent effects

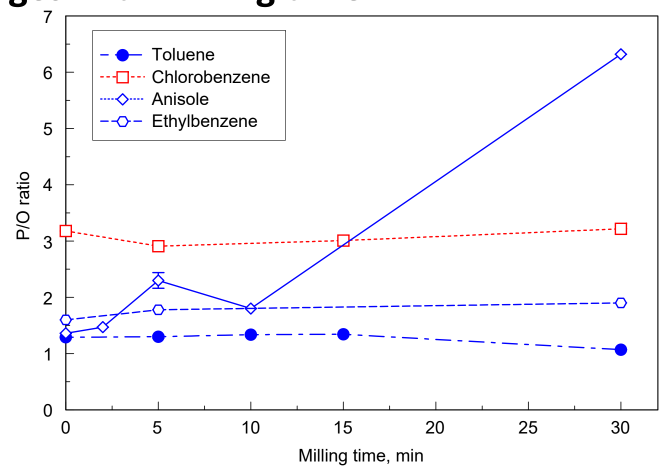
- High yield of nitro products observed for most precursors in short time
- Initial nitration rate roughly scales with the activation
- Nitration slows down in time or even reversed for some precursors





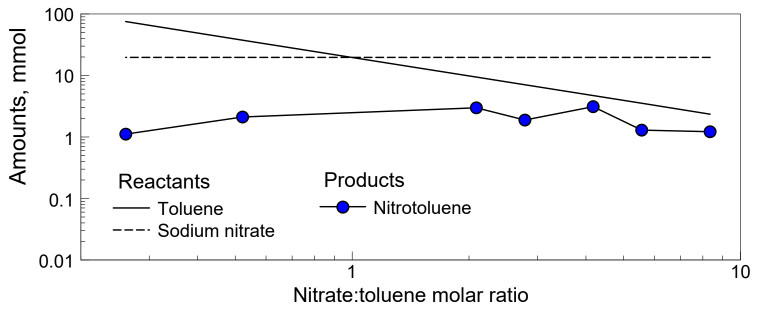
Regioselectivity changes with milling time

- ☐ High p/o ratio at 30 min for anisole is not associated with increased yield
 - may suggest selective decomposition of ortho isomer
- □P/O ratios for other precursors consistent with literature
 - Reactions on surface of catalyst





Yield vs. molar reactant ratio: toluene

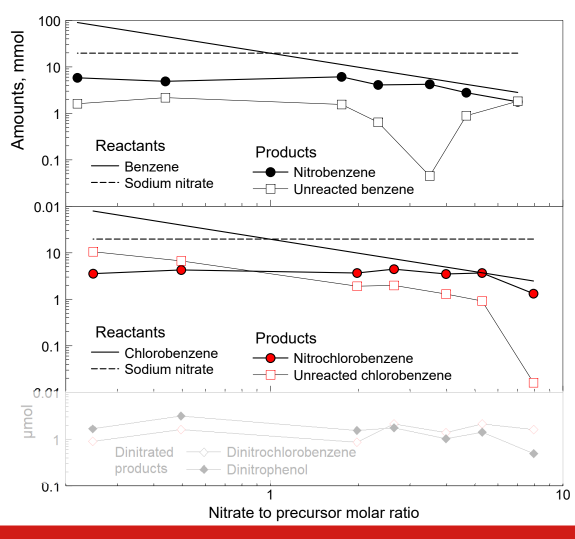


- ☐ Peak yield observed near reactant ratio of 4
- □Toluene-limited reactions show highest yields
- □ Excess of nitrate limits nitroproducts, possibly through blockage of active sites



Yield vs. mol. react. ratio: benzene, chlorobenzene

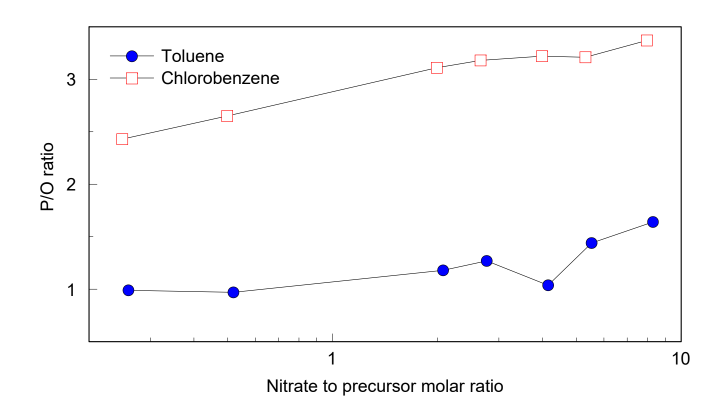
- □ Near complete conversion observed when reaction is precursor-limited
- ☐ Nitration attains a plateau, does not approach the limit when the reactants are nitrate-limited
- ☐ Small traces of **dinitrated products** formed with chlorobenzene
 - Also observed for other precursors





Regioselectivity

- □ Selectivity increases for both toluene and chlorobenzene when the reaction becomes limited by the available precursor
- ☐ Implication: reaction is favored when it occurs in a thin layer of precursor on surface of the catalyst





Which precursor properties control mechanochemical nitration?

Model milling as a simplified nitration reaction

 $Precursor(P) \rightarrow Nitrated\ product(NP)$

– 1st order reaction:

- Test correlation of rate constant, k, with precursor properties
- Statistical approach, i properties considered
- Use linear models:
 - $ln(k) = ln(k_0) + \sum_i a_i \cdot \zeta_i$

 ζ_i : precursor properties, a_i : linear coefficients

- Coefficients a_i determined using least-squares fitting



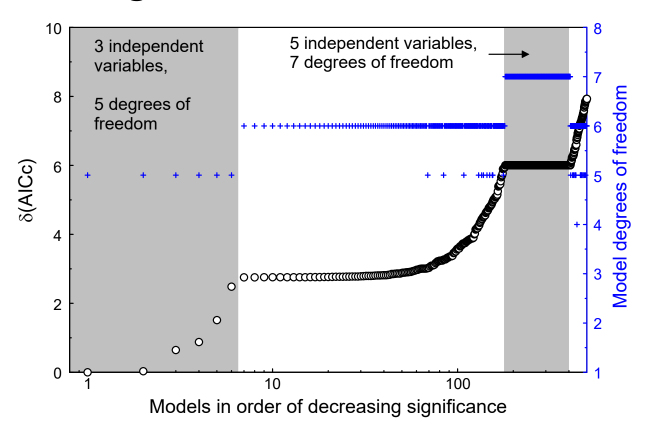
Reactant	Active group	Relative rate log k/k _H	Heats of formation		Viscosities			Heat of		Steric factors			Gas	
			reactant	product	Kinematic	Dynamic	Density	vapor, reactant	Dipole moment	A- factor	Ligand Repulsive energies	Proton affinity	basicity	Ioniz. energy
			kJ/mol	kJ/mol	mm²/s	сР	g/mL	kJ/mol	Debye	kcal/m ol	kcal/mol	kJ/mol	kJ/mol	eV
Nitro- toluene	NO ₂	-3.72	-48.2	-66.4	solid	solid	1.1	74.8	1.35	1.10	18	815.2	782.7	9.46
Chloro- benzene	CI	-0.66	11.5	-48.74	0.92	1.021	1.11	41	1.69	0.53	3.3	753.1	724.6	9.07
Benzene	Н	0	49	12.5	0.647	0.567	0.876	33.9	0	0	0	750	725.4	9.24
Ethyl benzene	C ₂ H ₅	0.90	27	-13.22	0.669	0.579	0.866	41	0.59	1.79	34	788	760.3	8.77
Toluene	CH ₃	1.02	12	-48.2	0.62	0.538	0.867	37	0.332	1.74	17	784	756.3	8.83
p-Xylene	(CH ₃) ₂	2.04	-24.4	-5.62	0.93	0.800	0.86	42	0	3.48	34	812.1	766.8	8.44
Phenol	ОН	2.22	-165	-207	solid	solid	1.07	69.7	1.224	0.60	10	817.3	786.3	8.49
Anisole	OCH ₃	4.68	-120	-197	0.99	0.985	0.995	44	1.262	0.75	31	839.6	807.2	8.20

Data Analysis

- 13 solvent properties considered
- experimental results for 6 different precursors
- → set of equations is underdetermined
- selection:
 - fit at most 5 = (6-1) coefficients/solvent properties at a time
 - do not simultaneously fit coefficients/solvent properties that are highly correlated
 - gas basicity & proton affinity
 - kinematic & dynamic viscosity
 - different measures of steric hindrance
- 1120 parameter combinations remain
 - ranking of statistical significance using logarithmic likelihood of the model fit (highest is best), and the Akaike's information criterion [$\delta(AIC_c)$, lowest is most prudent = "best"]



Ranking models



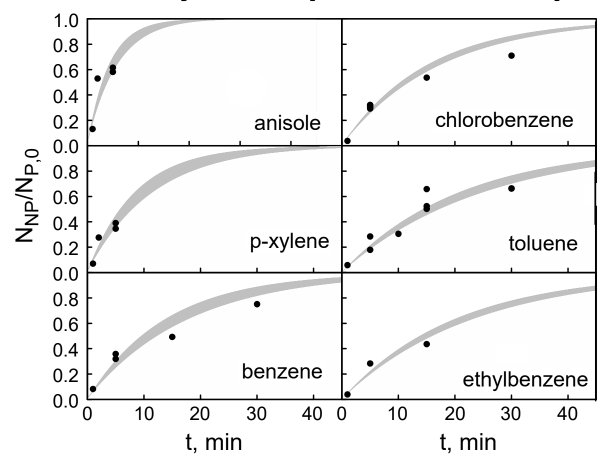
- Major differences in how models fit according to how many properties considered (3, 4, 5)
- Practically, only first handful of models need to be considered
 - differences between those are minor

Coefficients for top models

					<u> </u>					<u> </u>					
$\ln(k_0)$	Dipole moment(Nelson et al., 1967)	Dynamic viscosity, η (Kroenlein et al., 2012; Van Velzen et al., 1972)	Ionization energy (Hunter and Lias, 1998)	Gas basicity (Hunter and Lias, 1998)	ΔΗ _{νap} (P) (Acree and Chickos, 2010)	ΔH _f (P) (Cox, 1961; Geiseler, 1970; Roux et al., 2008)	ΔH _f (NP) (Lebedeva et al., 1971; Lenchitz et al., 1971; Sabbah and Gouali, 1994)	Relative rate, $\log(k/k_{\rm H})$ (Hansch et al., 1991)	Kinematic viscosity, v (Kroenlein et al., 2012; Van Velzen et al., 1972)	Proton affinity (Hunter and Lias, 1998)	Density, ρ (Lide, 2004)	Steric A-value (White et al., 1999)	Ligand repulsive energies (White et al., 1999)	$\delta ext{-AIC}_{ ext{c}}$	model weight, % (Bartoń, 2020)
					1	coe	efficient	$s a_i$							
31.8				-0.051	0.074			0.979						0	2.01
-1.79						-0.023	0.012					-0.286		0.030	1.98
-161			8.762						4.233	0.099				0.646	1.45
19.2				-0.031				0.697	1.371					0.880	1.29
20.8		0.921		-0.033				0.754						1.515	0.94
-18.1			1.494					0.446	2.280					2.481	0.58
-1.43						-0.023	0.012				-0.358	-0.304		2.755	0.51
	-0.498			0.010					3.243			-0.267		2.755	0.51
-12.5	-0.452								2.875	0.011		-0.302		2.755	0.51
-1.60		-0.183				-0.024	0.012					-0.306		2.755	0.51
69.4				-0.099			-0.008	1.279					0.052	2.756	0.51
5.36	-0.494		-1.061						2.667			-0.340		2.756	0.51



Predictions by the top model vs. experiments



Yields of nitroproducts are predicted reasonably well

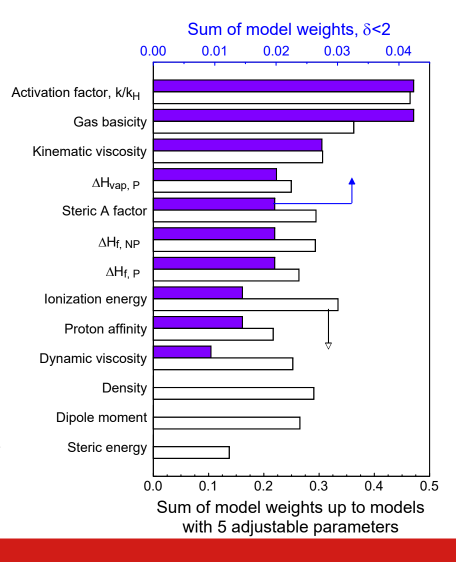


Influential individual factors

"Parameter Importance" calculated as sum of model weights of the individual parameters for all models with δ_{AICc}<2
 (5 models)

Importance for mechanochemical nitration:

- □ Activation factor has the strongest effect
- ☐Gas basicity, kinematic viscosity are likely important
- ☐ Effects of density, dipole moment, steric energy on nitration rate are insignificant



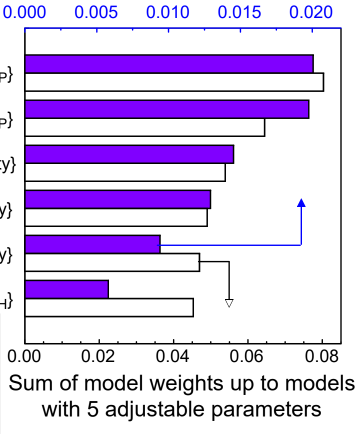


Influential factor combinations

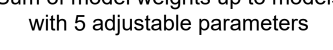
□ Groups of factors identified in the best ranked models rank the same as top models

{Activation factor, k/k_H ; Gas basicity; $\Delta H_{vap. P}$ } {Steric A factor; $\Delta H_{f NP}$; $\Delta H_{f P}$ } {Kinematic viscosity; Ionization energy; Proton affinity} {Activation factor, k/k_H; Gas basicity; Kinematic viscosity} {Activation factor, k/k_H; Gas basicity; Dynamic viscosity} {Kinematic viscosity; Ionization energy; Activation factor, k/k_H}

□ Activation, gas basicity, heat of vaporization, steric A factor, heat of formation of nitro product and heat of formation of precursor likely need to be considered in a mechanistic model



Model weights





Part 2. Summary

- □ Near complete mechanochemical nitration achieved with a broad range of aromatic precursors using MoO₃ as catalyst
 - Small amounts of dinitrated products observed in all cases when mononitrated products formed
- □ **Selectivity** observed is mostly consistent with earlier work using MoO₃ as catalyst; abnormally high selectivity is observed in certain experiments
- ☐ Extended milling may reduce yield of nitro products
- ☐ Processing experimental data suggests that mechanochemical nitration rate depends on properties of aromatic precursor: activation, gas basicity and kinematic viscosity are most influential
- ☐ Results of data processing indicate **potential avenues for process optimization**, e.g., by varying temperature of mechanochemical nitration



Part 3. Using different <u>nitrate salts</u> as nitronium sources

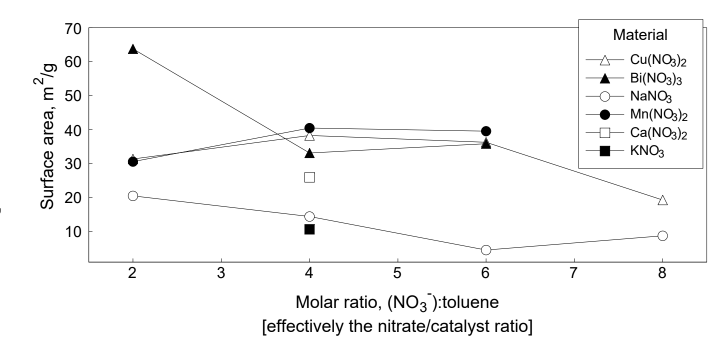
- Nitrates varied
 - different cation charge
 - different cation electronegativity
 - different hydration states
- Reactant ratio varied
 - expressed as (NO₃)/toluene
 - implies equiv. values of (NO₃)/catalyst
- Toluene precursor, constant amount
- MoO₃ oxide catalyst, constant amount

Nitrate	Mass in g for reactant ratio,				
	(moles of NO ₃ ⁻)/(moles of C ₇ H ₈)				
	2	4	6	8	
NaNO ₃	0.80	1.60	2.40	3.20	
KNO_3	0.95	1.91	2.86	3.82	
$Ca(NO_3)_2 \cdot 4H_2O$	1.11	2.22	3.33	4.44	
$Cu(NO_3)_2 \cdot 2.5H_2O$	1.30	2.60	3.90	5.20	
$Bi(NO_3)_3 \cdot 5H_2O$	1.52	3.04	4.58	6.16	
$\mathbf{Mn}(\mathrm{NO}_3)_2 \cdot 6\mathrm{H}_2\mathrm{O}$	1.18	2.36	3.54	4.72	



Effect of nitrate on premilling step

- Gradual changes only
- Anhydrous (Na, K)
 nitrates show opposite
 trend compared to
 hydrated nitrates (Cu, Bi,
 Mn, Ca)



Specific surface areas of MoO₃ catalyst after premilling with different nitrates for 120 min



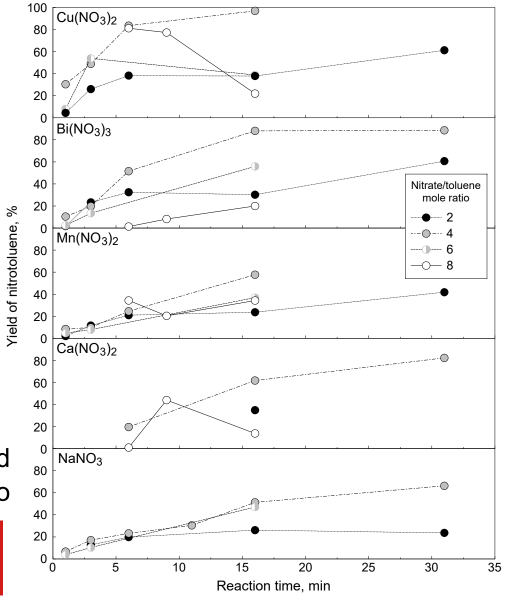
Effect of nitrate on mechanochemical nitration

Reaction time trends

- Increase for all, as expected
- Rates different for different nitrates
- Reactant ratio shows inconsistent trends

MNT yield vs. time, grouped by nitrate and reactant ratio





Effect of nitrate on mechanochemical nitration

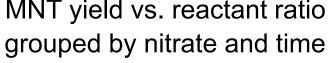
Reactant ratio trends

Highest yield observed for molar reactant ratio of (NO_3) /toluene = 4

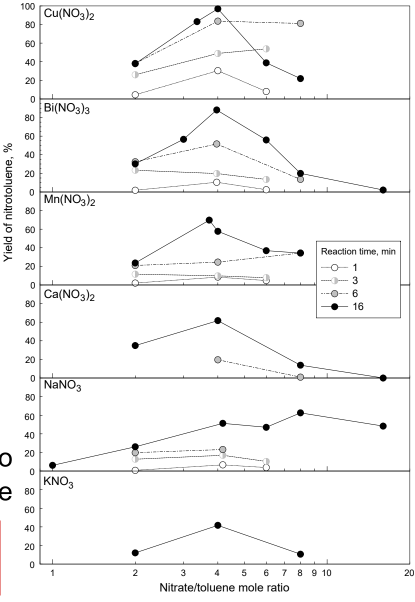
Part 3: nitrate

Except for NaNO₃

MNT yield vs. reactant ratio grouped by nitrate and time







Oxidation byproduct: benzaldehyde

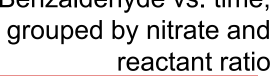
Reaction time trends

Amount of oxidation byproducts, spec. benzaldehyde generally increases with increasing reactant ratios

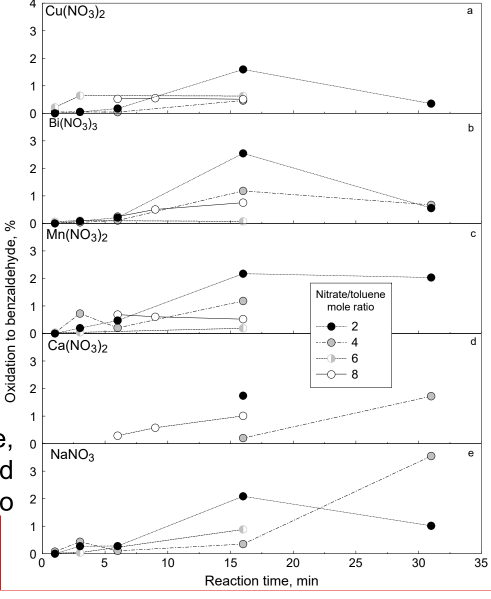
Part 3: nitrate

Nitrates are oxidizers

Benzaldehyde vs. time,







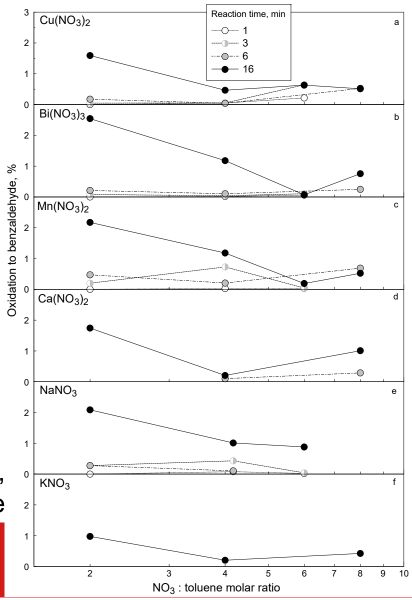
Oxidation byproduct: benzaldehyde

Reactant ratio trends

- No comparable highest formation rate observed at reactant ratio of 4
 - Benzaldehyde formation does not correlate with nitration

Benzaldehyde vs. reactant ratio, grouped by nitrate and time





Double nitration

Reaction time trends

- Higher nitrate amounts also result in formation of dinitrotoluene (DNT)
- Amounts low, but well detectable

DNT yield vs. time, grouped by nitrate and reactant ratio



0.2 Yield of dinitrotoluene, % $Mn(NO_3)_2$ 0.4 0.3 0.2 0.1 0.0 $Ca(NO_3)_2$ 1.5 1.0 0.5 0.0 NaNO₃ 0.4 0.3 0.2 10 15 20 25 30 35 Reaction time, min

Nitrate/toluene

mole ratio

Cu(NO₃)₂

0.4

0.2

0.1

0.0 0.4 0.3

Which nitrate property governs nitration rate?

- Attempts to correlate rates with cation properties showed no trends
 - electronegativity
 - ☐ charge
 - □ hydration state of the nitrate salt

Consider energy of **global reactions**:

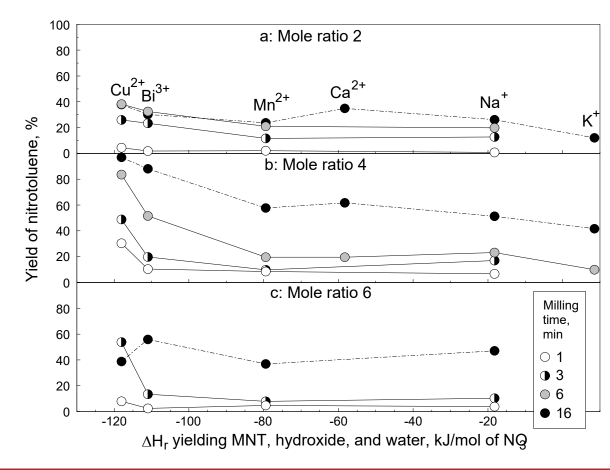
- Reactions leading to corresponding **hydroxides** (instead of oxides)

Global reaction	ΔH _r , kJ/mol
	of NO ₃ -
$KNO_3 + C_7H_8 \rightarrow KOH + C_7H_7NO_2$	8.5
$NaNO_3 + C_7H_8 \rightarrow NaOH + C_7H_7NO_2$	-18.23
$Ca(NO_3)_2 \cdot 4H_2O + 2C_7H_8 \rightarrow Ca(OH)_2 + 2C_7H_7NO_2 + 4H_2O$	-58.31
$Bi(NO_3)_3 \cdot 5H_2O + 3C_7H_8 \rightarrow Bi(OH)_3 + 3C_7H_7NO_2 + 5H_2O$	-110.99
$2Cu(NO_3)_2 \cdot 5H_2O + 4C_7H_8 \rightarrow 2Cu(OH)_2 + 4C_7H_7NO_2 + 5H_2O$	-118.06



Yield vs. energy of global nitration reaction

- Correlations observed
- Particularly consistently, for reactant ratio 4





Reaction rate constant vs. energy of global nitration

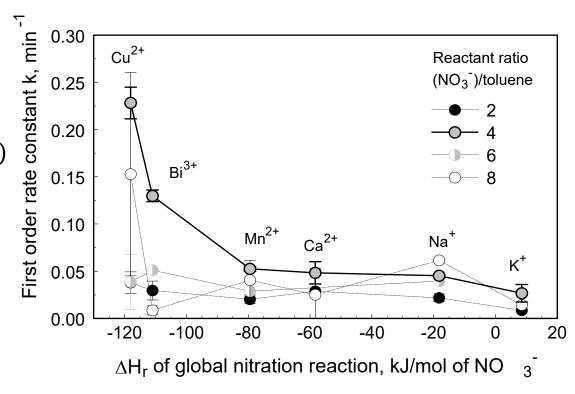
reaction

Model milling as simplified nitration reaction

Precursor $(P) \rightarrow Nitrated \ product \ (NP)$ 1st order reaction:

$$\frac{dC_P}{dt} = -k \cdot C_P \rightarrow k = -\ln\left(1 - \frac{C_{NP}}{C_{P,0}}\right) \cdot t^{-1}$$

 Correlate rate constant, k, with reaction enhalpies

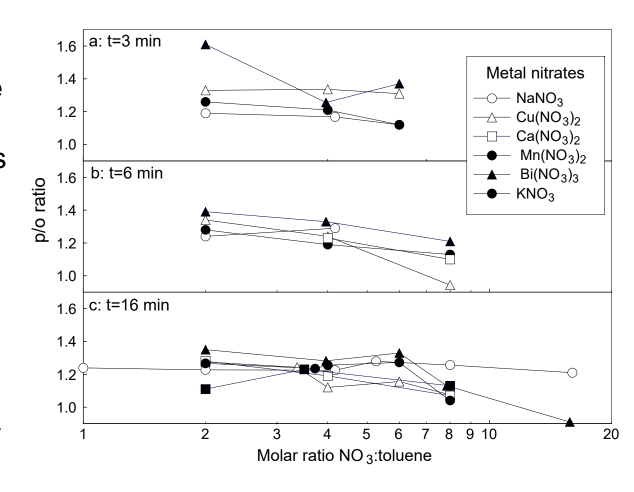


Consistent correlation for reactant ratio 4



Regioselectivity

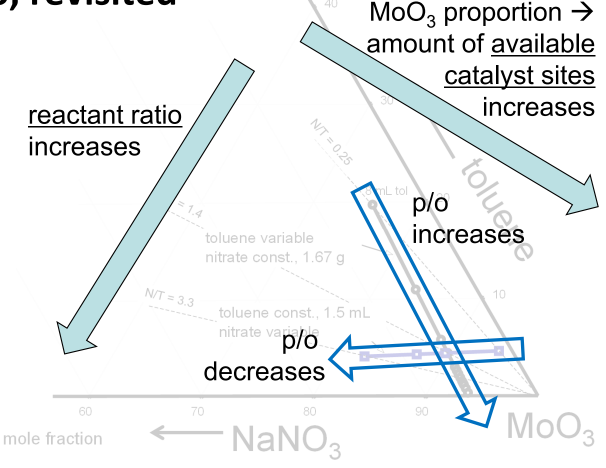
- para/ortho MNT isomer
 ratios consistently decrease
 with increasing
 (NO₃)/toluene reactant ratios
- No distinct dependence on type of the nitrate, or milling time
- This is an opposite trend from earlier observations where p/o ratio increased w/ increasing reaction ratio





Varying the reactant ratio, revisited

- Experimental series keeping one reactant constant:
 - nitrate variable (precursor const.)
 - precursor variable (nitrate const.)
- This implies changes in the proportion of catalyst in either case
- p/o ratio is determined by available cat, surface



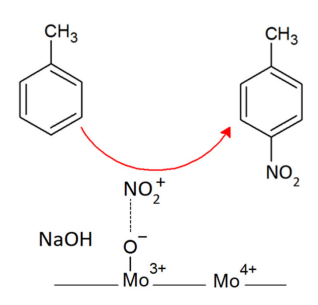


Reaction mechanism

Toluene is nitrated reacting with nitronium complexes

- attached to catalyst surface acid sites
- in the bulk fluid, away from these sites

Step 1: Sodium nitrate forms nitronium complexes on acid sites

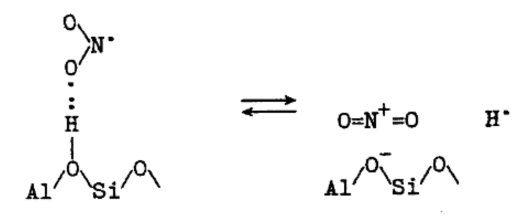


Step 2: aromatic substitution reaction

Reaction mechanism

- NO₂, nitronium, can exist
 - at the catalyst surface, or
 - in the fluid

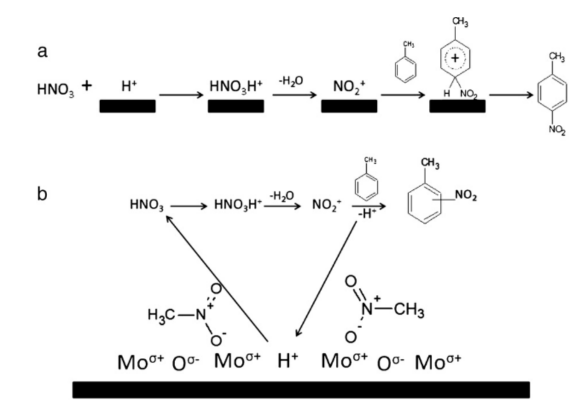
demonstrated in literature for other oxide surfaces



V. MALYSHEVA LUDMILA, A. PAUKSHTIS EUGENE & G. IONE KAZIMIRA (1995) Nitration of Aromatics by Nitrogen Oxides on Zeolite Catalysts: Comparison of Reaction in the Gas Phase and Solutions, Catalysis Reviews, 37:2, 179-226

Reaction mechanism

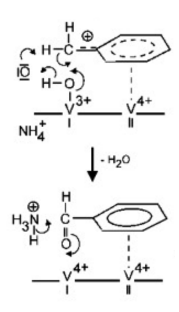
- Reaction at the catalyst surface preferentially produces the para isomer
- Reaction in the bulk fluid does not show this preference
- Increased p/o ratio indicates more reaction occurring at the catalyst surface



Joanna Adamiak, Waldemar Tomaszewski, Wincenty Skupiński, Interaction of nitromethane with MoO₃/SiO₂ and its influence on toluene nitration, Catalysis Communications, Volume 29, 2012, Pages 92-95

Reaction mechanism and oxidized byproducts

- Oxidation of aromatics on catalyst surface described in literature
- May necessitate multiple surface sites
- Consistent with observation that benzaldehyde amount decreases when higher amounts of nitrates block more and more catalyst surface sites (cf. slide 41)



Angelika Brückner (2003) Looking on Heterogeneous Catalytic Systems from Different Perspectives: Multitechnique Approaches as a New Challenge for In Situ Studies, Catalysis Reviews, 45:1, 97-150

Part 3. Summary

- Nitronium source appears to affect nitration only via energy of global reaction
- Optimum reaction rate consistently observed for (NO_3) /precursor molar ratio of 4
- Isomer ratios can be controlled by choice of reactant ratios



Project Summary

- Mechanochemical nitration of organic precursors requires solid catalysts with high acidity and both, Bronsted and Lewis sites
 - MoO₃ was the preferred catalyst
- Nearly complete mechanochemical nitration was achieved in many experiments
- Homogenizing catalyst with nitrate by premilling accelerates ensuing mechanochemical nitration
- Mechanochemical nitration with MoO₃ catalyst was successful for multiple aromatic compounds
- The selectivity was enhanced and the yield of the nitroproduct was increased when the volume of the aromatic
 precursor was reduced while the mass of metal oxide catalyst was fixed
 - The reaction preferably involves nitronium complexes attached on the catalyst surface
- Reaction depends on the aromatic activation by the functional group, gas basicity and enthalpy of vaporization of the aromatic precursor. Additional effect by reaction enthalpies and kinematic viscosity
- The highest mechanochemical nitration rate observed for Cu(NO₃)₂ serving as the source of nitronium.
- The **yield** of MNT as well as the **reaction rate** *correlated* with the **enthalpy of the global nitration reaction** with the corresponding metal hydroxide as a product
- A second nitration of the aromatic ring was observed for all the precursors used



Future work

- A path is outlined to further scale-up and optimization of mechanochemical nitration of organic compounds.
- First practical goal: mechanochemical synthesis of single-nitrated products of aromatic precursors
 - Use identified relationships between the process parameters, materials, and yield and reaction rate for the nitroproducts
- Challenges:
 - determine conditions necessary to transfer the reaction parameters for the attritor mill configuration
 - develop methods for in-situ separation of the reaction products from the milling tools and catalyst
- Other research:
 - Reusing or recycling the catalyst
 - A detailed mechanism of mechanochemical nitration



Supplement: ARL Effort

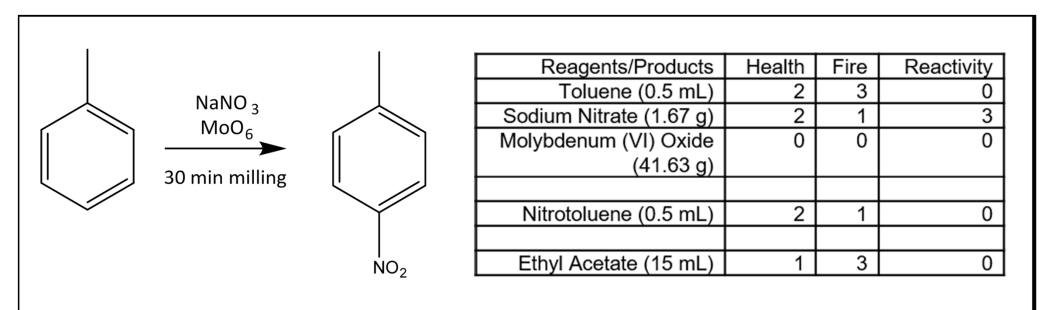
Leah Winhard, PI (WMRD, ARL)
Melissa Garner (ARL Summer student)

ARL's contribution focused on the evaluation of the mechanochemical nitration technique developed by NJIT as viable nitration technique for a range of substrates of interest

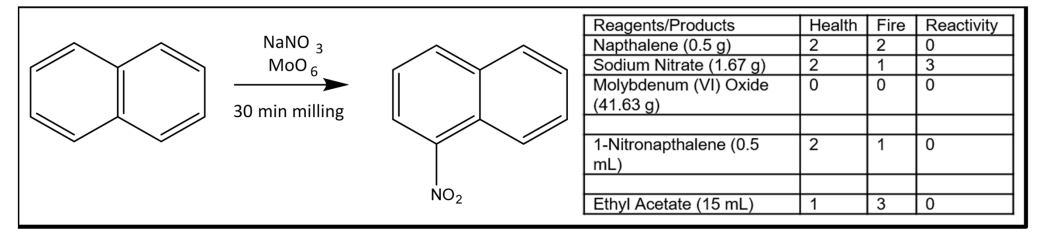
- SOP approved
- Process applied to both commercially available substrates and non-energetic precursors designed and synthesized at ARL



Reaction using toluene as a substrate



Reaction using naphthalene as a substrate



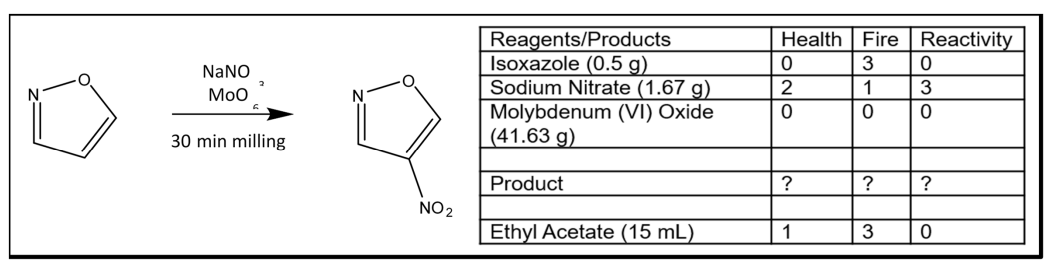
Additional substrates

Milling was also tested with the following list of commercially available substrates:

- ☐ Isoxazole
- ☐ 1,2-Benzisoxazole
- □ Maleimide
- □ Pyrazole
- □ Pyrrole
- All reaction mixtures showed some production of nitrated products via IR.
- NMR analysis revealed low yields (<20%) of nitrated products along with unreacted substrate and decomposition products
- For pyrazole substrate, a highly hygroscopic product was formed, and decomposed too rapidly for accurate analysis



Reaction using isoxazole as a substrate



ARL-designed precursors

- 5,5'-Dihydroxymethyl-3,3'-bis-isoxazole
- 2,2'-([3,3'-biisoxazole]-5,5'-diyl)bis(Ethan-1-ol)
 - Both readily form their respective nitrate ester energetic compounds in high yields (>90%) through traditional wet nitration chemistry.
 - The nitrated compounds are relatively insensitive to impact and friction, concern with milling nitrate esters prompted using very small amounts of precursor (100 mg or less) in the milling process.
- Analysis showed some degree of nitration, yield was low
 - NJIT comment: milling conditions were not optimized, e.g., no premilling catalyst with nitrate was used

