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Copolymerization of Ethylene and Vinyl Fluoride by Self-Assembled Multinuclear Palladium Catalysts

Qian Liu and Richard F. Jordan * 

Department of Chemistry, The University of Chicago, 5735 South Ellis Avenue, Chicago, IL 60637, USA; lqdenom1216@gmail.com

* Correspondence: rfjordan@uchicago.edu

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Abstract: The self-assembled multinuclear Pd^{II} complexes $\{(Li-OPO^{OMe2})PdMe(4\text{-}5\text{-nonyl-pyridine})_4Li_2Cl_2$ (**C**, $Li-OPO^{OMe2} = PPh(2-SO_3Li-4,5-(OMe)_2-Ph)(2-SO_3^-4,5-(OMe)_2-Me-Ph)$), $\{(Zn-OP-P-SO)PdMe(L)\}_4$ (**D**, $L = \text{pyridine or } 4\text{-}^t\text{Bu-pyridine}$, $[OP-P-SO]^{3-} = P(4\text{-}^t\text{Bu-Ph})(2-PO_3^{2--5-Me-Ph})(2-SO_3^-5-Me-Ph)$), and $\{(Zn-OP-P-SO)PdMe(\text{pyridine})\}_3$ (**E**) copolymerize ethylene and vinyl fluoride (VF) to linear copolymers. VF is incorporated at levels of 0.1–2.5 mol% primarily as in-chain $-CH_2CHFCH_2-$ units. The molecular weight distributions of the copolymers produced by **D** and **E** are generally narrower than for catalyst **C**, which suggests that the Zn-phosphonate cores of **D** and **E** are more stable than the Li-sulfonate-chloride core of **C** under copolymerization conditions. The ethylene/VF copolymerization activities of **C–E** are over 100 times lower and the copolymer molecular weights (MWs) are reduced compared to the results for ethylene homopolymerization by these catalysts.

Keywords: multinuclear catalyst; vinyl fluoride; copolymerization; fluorinated polyethylene

1. Introduction

The coordination–insertion copolymerization of ethylene with polar vinyl monomers by Pd^{II} catalysts has been extensively studied [1–8]. Vinyl halides are particularly challenging polar comonomers due to (i) their poor competition with ethylene for binding to Pd^{II} catalysts [9,10], (ii) the formation of inactive L_nPd-X complexes by β -X elimination of L_nPdCH_2CXR species (generated by 1,2 $CH_2=CHX$ insertion or 2,1- $CH_2=CHX$ insertion followed by chain walking) [11–20], and (iii) the low insertion reactivity of $L_nPdCHXCH_2R$ species formed by 2,1 $CH_2=CHX$ insertion [16,21]. We previously reported that $(PO)PdMe(L)$ catalysts (**A**, Figure 1) that contain phosphine-arenesulfonate ligands (PO^-) copolymerize ethylene and vinyl fluoride (VF) to linear copolymers with up to 0.55 mol% VF incorporation [22–25]. The catalyst activities are significantly reduced and the polymer molecular weights (MWs) are also reduced in ethylene/VF copolymerization compared to ethylene homopolymerization under the same conditions. For example, $\{P(2\text{-Et-Ph})_2(2-SO_3-Ph)\}PdMe(py)$ exhibits an activity of $296 \text{ kg}\cdot\text{mol}^{-1}\cdot\text{h}^{-1}$ in ethylene homopolymerization and produces polyethylene (PE) with $M_n = 16,560 \text{ Da}$ (300 psi ethylene, toluene, 80 °C), while in ethylene/VF copolymerization the activity and polymer M_n are reduced to $5 \text{ kg}\cdot\text{mol}^{-1}\cdot\text{h}^{-1}$ and 6800 Da, respectively (300 psi total pressure, VF/ethylene = 4/1, toluene, 80 °C) [22]. The ethylene/VF copolymers produced by $(PO)PdMe(L)$ catalysts contain internal $-CH_2CHFCH_2-$ units formed by 1,2 and/or 2,1 VF insertion into growing $(PO)PdR$ species. The copolymers also contain $-CH_2CHFCH_3$, $-CH_2CHF_2$, and $-CH_2CH_2F$ chain ends. The $-CH_2CHFCH_3$ units are formed by 2,1 VF insertion into $(PO)PdH$ species followed by chain growth. It was proposed that the $-CH_2CHF_2$ and $-CH_2CH_2F$ groups are generated by VF or ethylene insertion of $(PO)PdF$ species (formed by β -F elimination of $(PO)Pd(CH_2CHFR)$ species), followed by chain growth. Strong support for this proposal was provided by the demonstration that $(PO^{BpOMe})PdF(2,6\text{-lutidine})$

$[(\text{PO}^{\text{Bp,OMe}})]^- = [\text{P}(2',6'-(\text{OMe})_2\text{-biphenyl})(2\text{-OMe-Ph})(2\text{-SO}_3\text{-5-Me-Ph})]^-$ reacts with VF to yield the 1,2-insertion product $(\text{PO}^{\text{Bp,OMe}})\text{Pd}(\text{CH}_2\text{CHF}_2)(2,6\text{-lutidine})$ and with ethylene to yield PE with $-\text{CH}_2\text{CH}_2\text{F}$ end groups [23]. It was also demonstrated that $(\text{PO}^{\text{Bp,OMe}})\text{Pd}(\text{CH}_2\text{CHF}_2)(2,6\text{-lutidine})$ reacts with ethylene to produce PE with $-\text{CH}_2\text{CHF}_2$ chain ends. The ability of $(\text{PO})\text{PdF}$ species to undergo olefin insertions is key to the ability of $(\text{PO})\text{PdRL}$ species to catalyze ethylene/VF copolymerization. The copolymerization of ethylene with other $\text{CH}_2=\text{CHX}$ monomers by $(\text{PO})\text{PdRL}$ catalysts has been extensively studied [26–34].

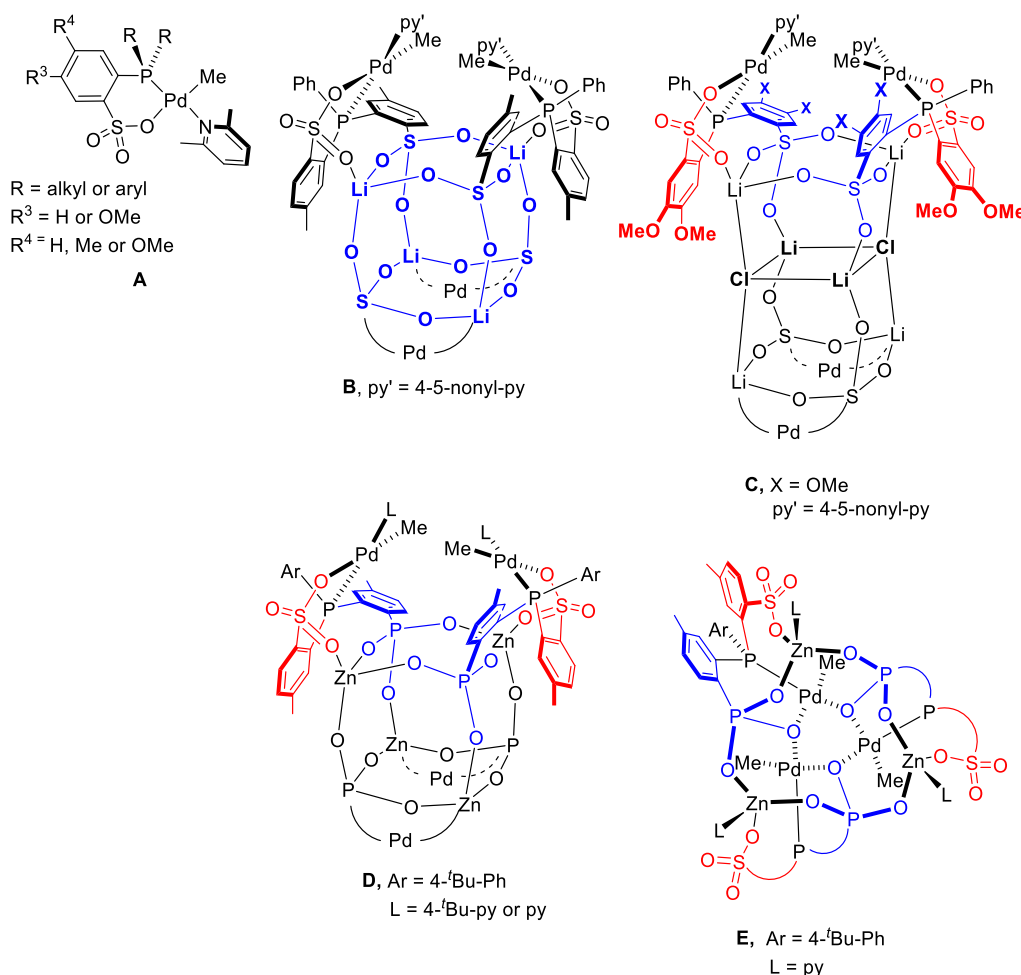


Figure 1. $(\text{PO})\text{PdR}$ Complexes. $\text{Py}' = 4\text{-}(5\text{-nonyl})\text{pyridine}$. The lower $(\text{Li-OPO})\text{PdMe}(\text{py}')$ or $(\text{OP-P-SO})\text{PdMeL}$ units in the schematic structure of **B**, **C**, and **D** are denoted by “Pd”.

The related $(\text{Li-OPO})\text{PdMe}(\text{py}')$ complex ($\text{py}' = 4\text{-5-nonyl-pyridine}$) based on the phosphine-bis(arenesulfonate) ligand $\text{PPh}(2\text{-SO}_3\text{Li-5-Me-Ph})(2\text{-SO}_3^-5\text{-Me-Ph})$ ($\text{Li}[\text{OPO}]^-$) self-assembles into the tetrameric $\{(\text{Li-OPO})\text{PdMe}(\text{py}')\}_4$ species **B**, which is held together by a $\text{Li}_4\text{S}_4\text{O}_{12}$ double 4-ring (D4R) cage (Figure 1) [35–37]. “Pd₄ cage” catalyst **B** produces linear PE with a high MW and linear ethylene/VF copolymers that contain up to 3.6 mol% VF (Table 1, entries 8,9). The ethylene/VF copolymers produced by **B** contain internal $-\text{CH}_2\text{CHFCH}_2-$ units as well as $-\text{CH}_2\text{CHFCH}_3$ and $-\text{CH}=\text{CHF}$ end groups, but no NMR-detectable $-\text{CH}_2\text{CF}_2\text{H}$ or $-\text{CH}_2\text{CH}_2\text{F}$ chain ends. The $-\text{CH}=\text{CHF}$ chain ends are most likely formed by 2,1 VF insertion into growing $(\text{Li-OPO})\text{Pd-R}$ species, followed by $\beta\text{-H}$ elimination. However, **B** undergoes partial disassembly to the monomeric $(\text{Li-OPO})\text{PdMe}(\text{py}')$ “Pd₁” species under polymerization conditions, which strongly influences the MWs and molecular weight distributions (MWDs) of the polyethylene and ethylene/VF copolymers it produces. For example, **B** produces a high-MW ethylene/VF copolymer with a broad bimodal MWD ($M_w = 494$ kDa, PDI = 310; Table 1,

entry 9) in hexanes suspension, due to competing copolymerization by intact **B** (which generates the high-MW fraction) and monomeric (Li-OPO)PdR species (which generates the low-MW fraction). In contrast, **B** produces a low-MW copolymer with a narrow MWD ($M_w = 4200$, PDI = 2.4; entry 8) in toluene solution, indicative of nearly complete dissociation to monomeric species.

Table 1. Ethylene/vinyl-fluoride copolymerization by catalysts **C**, **D**, and **E** [38].

Entry	Cat.	Solvent	$P_{C_2H_4}$ (psi)	P_{VF} (psi)	Activity ^d (kg·mol ⁻¹ ·h ⁻¹)	M_w ^e (10 ³)	PDI ^e	VF Incorp ^f (mol%)	T_m ^g (°C)
1 ^a	C	toluene	220	80	1.4	20.3	8.0	0.10	128.8
2 ^a	C	toluene	130	120	0.44	1.92	1.4	0.25	ND ^h
3 ^a	C	hexanes	220	80	12.0	498	18.2	0.87	134.4
4 ^a	C	hexanes	130	120	3.4	419	26.2	2.5	132.8
5 ^{a,b}	D	toluene + PhCl	130	120	2.0	42.2	4.1	0.96	131.4
6 ^{a,b}	E	toluene + PhCl	220	80	4.9	50.3	2.4	0.43	134.7
7 ^{a,b}	E	toluene + PhCl	130	120	1.0	23.0	2.3	1.1	131.6
8 ^{a,c}	B	toluene	130	120	1.9	4.2	2.4	2.4	127.8
9 ^{a,c}	B	hexanes	130	120	1.4	494	310	3.6	127.8

^a Conditions: [Pd] = 10 μmol, temperature = 80 °C, time = 2 h, 50 mL solvent. ^b 1 equiv B(C₆F₅)₃ per L, 49/1 toluene/chlorobenzene solvent. ^c cited from reference 36. ^d Activity is reported per mol Pd. ^e Determined by Gel Permeation Chromatography (GPC) [38]. ^f VF incorporation in copolymer determined by ¹H NMR. ^g DSC. ^h Not determined due to the limited quantity of copolymer.

We recently reported several new multinuclear Pd “cage” catalysts (**C-E**) [39,40]. The sterically-expanded phosphine-bis(arenesulfonate) ligand PPh(2-SO₃Li-4,5-(OMe)₂-Ph)(2-SO₃⁻-4,5-(OMe)₂-Me-Ph) ([Li-OPO^{OMe2}]⁻) directs the self-assembly of the tetranuclear complex (Li-OPO^{OMe2})PdMe(py')₄Li₂Cl₂ (**C**, Figure 1), in which four (Li-OPO^{OMe2})PdMe(py') units are arranged around the periphery of a Li₄S₄O₁₂•Li₂Cl₂ cage [39]. Compound **C** is much less susceptible to cage dissociation than **B**. For example, **C** undergoes only 6.5% dissociation to monomeric species in CDCl₂CDCl₂ solution at 80 °C ([Pd₄]_{initial} = 4.7 mM), whereas **B** undergoes 38% dissociation under these conditions. In toluene, **C** is only partially dissociated into monomeric species and therefore produces PE with broad MWD as expected for a multi-site catalyst. In contrast, in hexanes suspension at 80 °C, **C** is resistant to disassembly and exhibits nearly ideal single-site behavior in ethylene homopolymerization and produces high-MW PE ($M_w = 1473$ kDa, PDI = 2.3, Figure 2). The solvent effects on the MWDs of the PEs produced by **B** and **C** may appear to be opposite but simply reflect the relative stabilities of **B** and **C** toward disassembly to monomeric species.

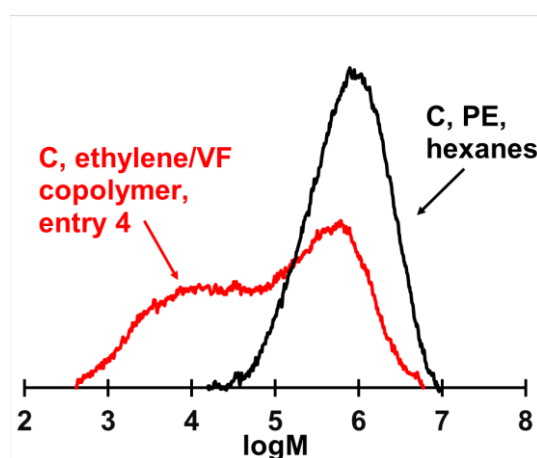


Figure 2. Molecular weight distributions of polyethylene (PE) [39] and ethylene/VF copolymer (Table 1, entry 4) generated by **C** determined by high temperature GPC. Polymerization conditions: hexanes solvent, 80 °C.

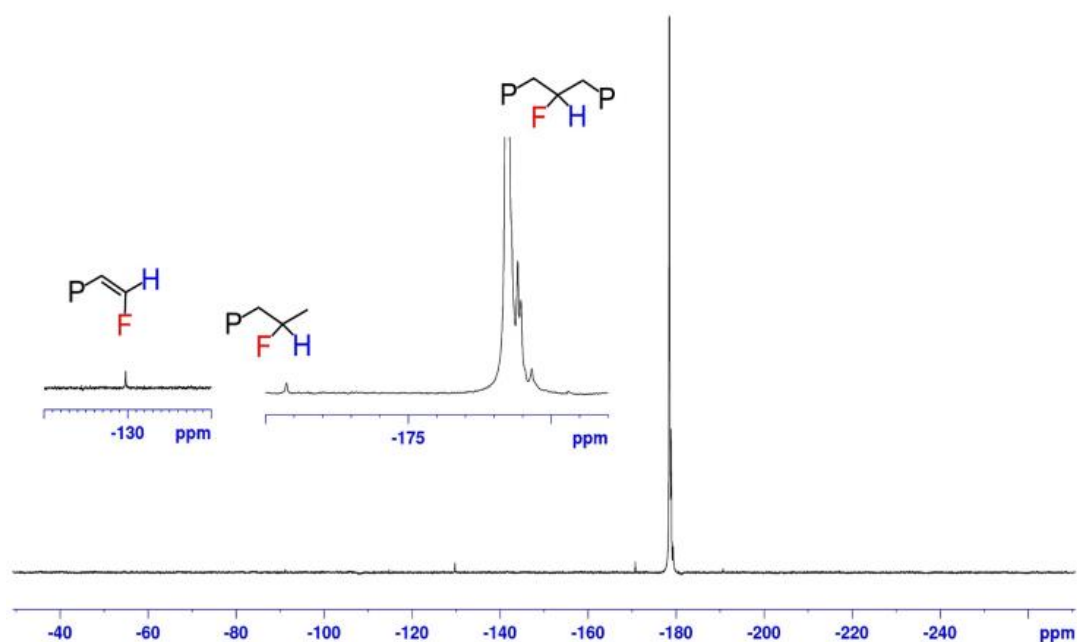


Figure 3. $^{19}\text{F}\{^1\text{H}\}$ NMR spectrum of ethylene/VF copolymer (*o*-dichlorobenzene- d_4 , 120 °C) produced by **C** (Table 1, entry 4).

The ethylene/VF copolymers produced by **C** in hexanes exhibit broad MWDs (Table 1, entries 3,4; Figure 2). Given the nearly ideal single-site behavior observed in ethylene homopolymerization by **C** in hexanes, it is unlikely that the broadening of the MWD of the ethylene/VF copolymers formed in this solvent is due to thermal disassembly of the cage structure (Figure 2). One possibility is that nucleophilic Pd-F species generated by 1,2-VF insertion and β -F elimination react with the Li^+ ions in the central cage, leading to the formation of new active Pd species [23,25,48–52]. Consistent with this proposal, $-\text{CH}_2\text{CH}_2\text{F}$ and $-\text{CH}_2\text{CHF}_2$ end groups, which are formed by ethylene and VF insertion of Pd-F species in copolymerization by mononuclear catalysts **A**, are not observed in the copolymer produced by **C** (Figure 3) [23]. This process provides a potential catalyst deactivation pathway.

Ethylene/VF copolymerization by **D** ($\text{L} = 4\text{-}^t\text{Bu-py}$) was investigated in a mixed toluene/chlorobenzene (49/1) solvent at a VF/ethylene feed ratio of 0.92 (Table 1, entry 5). **D** decomposes in the presence of free $4\text{-}^t\text{Bu-py}$ (and other Lewis bases). Therefore, 1 equiv $\text{B}(\text{C}_6\text{F}_5)_3$ per $4\text{-}^t\text{Bu-py}$ was added to a solution of **D** in chlorobenzene prior to dilution with toluene to sequester the $4\text{-}^t\text{Bu-py}$ that will be displaced by monomer, in order to minimize catalyst deactivation. Under these conditions, **D** produces a linear copolymer with 0.96 mol% VF incorporation. The MWD of the copolymer formed by **D** is unimodal but somewhat broadened ($\text{PDI} = 4.1$), which can be attributed to the presence of several diastereomeric forms of the catalyst. The copolymer contains in-chain $-\text{CH}_2\text{CFHCH}_2-$ and a small amount of VF-derived $-\text{CH}_2\text{CFHCH}_3$ chain ends. Catalyst **E** behaves similarly to **D**, incorporating 1.1 mol% VF in the form of both $-\text{CH}_2\text{CFHCH}_2-$ (major) and $-\text{CH}_2\text{CFHCH}_3$ (minor) units (Table 1, entry 7). The MWDs of the copolymers formed by **E** are narrow ($\text{PDI} \leq 2.4$), indicative of nearly ideal single-site catalysis. These results suggest that the multinuclear structures of **D** and **E** are substantially retained during ethylene/VF copolymerization.

The ethylene/VF copolymerization activities of **C–E** are over 100 times lower than the ethylene homopolymerization activities, and the copolymer MWs are reduced compared to the results for ethylene homopolymerization, as observed previously for **A** and **B** and mononuclear (PO)PdRL catalysts.

3. Conclusions

“Pd cage” catalysts C-E copolymerize ethylene with VF to linear copolymers with 0.1 to 2.5 mol% VF incorporation depending on the catalyst and reaction conditions. VF is incorporated predominantly as in-chain $-\text{CH}_2\text{CHFCH}_2-$ units, with minor $-\text{CH}_2\text{CFHCH}_3$ and, for C, *cis*- $\text{CH}=\text{CHF}$ end groups. C produces an ethylene/VF copolymer with a broad MWD in hexanes, which contrasts results for ethylene homopolymerization where nearly ideal single-site behavior is observed. These results suggest that the presence of VF promotes structural disruption of the catalyst, possibly through reaction of Pd-F species with the Li^+ ions in the cage. The ethylene/VF copolymerization behavior of C is very similar to that of B, except the extent of VF incorporation by C is somewhat lower than by B. D and E produce ethylene/VF copolymers with narrow MWDs, suggesting that the cage assemblies remain substantially intact during copolymerization. The activities of C–E in ethylene/VF copolymerization are >100 times lower than in ethylene homopolymerization. This inhibition effect is a major roadblock to the development of more practical ethylene/VF copolymerization catalysts.

Supplementary Materials: The following are available online at <http://www.mdpi.com/2073-4360/12/7/1609/s1>.

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