



Frustrated Lewis Pairs

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Ultra-High-Molecular-Weight Polymers Produced by the Immortal Phosphine-Based Catalyst System

Yun Bai, Jianghua He, and Yuetao Zhang*

Abstract: A strong organophosphorus superbase, N-(diphenylphosphanyl)-1,3-diisopropyl-4,5-dimethyl-1,3-dihydro-2H-imidazol-2-imine (IAP3) was combined with a sterically encumbered but modestly acidic Lewis acid (LA), (4-Me-2,6- 1 Bu₂-C₆H₂O)Al 1 Bu₂ ((BHT)Al 1 Bu₂), to synergistically promote the frustrated Lewis pair (FLP)-catalyzed living polymerization of methyl methacrylate (MMA), achieving ultrahigh molecular weight (UHMW) poly(methyl methacrylate) (PMMA) with M_n up to 1927 kgmol⁻¹ and narrow molecular weight distribution (MWD) at room temperature (RT). This FLP catalyst system exhibits exceptionally long lifetime polymerization performance even in the absence of free MMA, which could reinitiate the desired living polymerization after the resulting system was held at RT for 24 h.

here have been continuing efforts in the development of living polymerization systems to produce polymers with controlled molecular weight (MW) and narrow molecular weight distribution (MWD) since the properties of polymeric materials are closely related to their MW and MWD.[1-4] However, it remains as a challenging task to synthesize UHMW polymers, ^[5] especially PMMA with $M_n > 10^6$ g mol⁻¹, only a handful of examples have been achieved under harsh reaction conditions so far, such as PMMA with M_n up to 3.6 × 10⁶ g mol⁻¹ prepared through free radical polymerization under high pressure of 500 MPa and about 60°C.[6,7] On the other hand, the synthesis of UHMW PMMA by anionic polymerization was restricted by difficulty in obtaining the ultra-purified MMA and solvents and the requirement of low temperature. [8,9] There is no report on the production of UHMW PMMA with $M_{\rm n} > 10^6 \, {\rm g \, mol^{-1}}$ and narrow MWD under mild condition up to date. Therefore, it is a great impetus to research efforts to develop new catalyst system to overcome these limitations.

As a newly emerging polymerization technique, Lewis pair polymerization (LPP) has demonstrated extraordinary activity towards various monomers and exhibits very promising prospect in polymer synthesis in recent years, [10] since the well-established application of FLP in small-molecule chemistry. [11] Despite the development of a wide range of

effective LP systems, the application of LPP was still hampered by both low initiation efficiencies and chain-termination side reactions. Until recently, by choosing appropriate combination of Lewis base (LB) and LA with matching acidity and steric hindrance, the first example of authentic FLP catalyst system has been developed to successfully remove these restriction barriers and achieve living (co)polymerization of MMA and benzyl methacrylate, producing PMMA with M_n up to 351 kg mol⁻¹.

As a strong nucleophile superior than many organobases such as alkylphosphines and N-heterocyclic carbenes, [15] organophosphorus(III) superbase, imidazolin-2-ylidenamino substituted phosphines (IAPs) are capable of capturing CO₂ and SO₂ molecules to form stable phosphine-based $adducts^{[16-18]}$ and activating extremely inert greenhouse gas sulfur hexafluoride. [19] Although strong donor ability and easily adapted steric hindrance of IAPs should enpower them to be a very promising LB candidate for an FLP system, there has been no report on the employment of IAPs for LPP yet. Herein five sterically hindered IAPs with the same imidazoline framework but varied substituents on the N atom (Ndiisopropylphosphanyl, **IAP1**; *N*-di-tert-butylphosphanyl, **IAP2**; *N*-diphenylphosphanyl, **IAP3**; *N*-dimesitylphosphanyl, *N*-(bis(6,6,6,6,6-pentafluoro-6λ8-hexa-1,3,4-triyn-1yl))phosphanyl, IAP5, Scheme 1) were synthesized and employed to combine with a series of organoaluminum LAs to achieve an immortal FLP catalyst system for living polymerization of MMA at RT, producing PMMA with medium to UHMW (up to 1927 kg mol⁻¹), narrow MWD (*Đ* as low as 1.06) and high to quantitative initiation efficiencies. To the best of our knowledge, this is for the first time UHMW PMMA with $M_n > 10^6 \,\mathrm{g\,mol^{-1}}$ and narrow MWD was achieved in polymer synthesis under mild conditions so far.

 $\textbf{Lewis acids (LA):} \qquad \text{Al}(C_6F_5)_3, \qquad (\text{BHT})_2\text{AlMe}, \quad (\text{BHT})_2\text{Al/Bu}, \quad (\text{BHT})\text{Al/Bu}_2$

Scheme 1. Structures of the Lewis acids, Lewis bases, monomers, and the resulting (co)polymers investigated in this study.

Supporting information (including experimental details, NMR spectra, and further tabular data) and the ORCID identification number(s) for the author(s) of this article can be found under: https://doi.org/10.1002/anie.201811946.

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When combined with a strongly acidic Al(C₆F₅)₃, IAPbased LPs exhibited extraordinarily effective catalytic activity for polymerization in an 800:1:2 $MMA/IAP/Al(C_6F_5)_3$ ratio, achieving quantitative monomer conversion by IAP1 to IAP4-based LPs in 30 s (Table 1, runs 1-4) but only 41% MMA conversion by IAP5-based LP in 24 h (Table 1, run 5) probably due to the lower nucleophilicity of IAP5 resulting from the presence of electronwithdrawing pentafluorophenyl substituent. Since IAP3 exhibited the highest initiation efficiency I*% of 92 among the investigated IAPs, it was employed for the following polymerization. MALDI-TOF MS spectrum indicated the exclusive presence of a cyclic chain end derived from backbiting cyclization side reactions in the low-MW PMMA produced by $IAP3/Al(C_6F_5)_3$ LP (Supporting Information, Figur-

es S36 and S37), which revealed that polymerization catalyzed by $IAP3/Al(C_6F_5)_3$ LP is not a living process.

Previously, we have demonstrated that both the acidity and steric hindrance of LA play significantly important role in the reactivity of polymerization. Therefore, a series of organoaluminum LAs with descending acidity on a relative scale[20] were examined for MMA polymerization in an 800:1:2 MMA/IAP3/LA ratio $(Al(C_6F_5)_3 (100\%) >$ $(BHT)_2AlMe (86\%) \approx (BHT)_2Al^iBu (85\%) > (BHT)Al^iBu_2$ (74%); Supporting Information, Table S1). It turned out that LP composed of bulky (BHT)2AlMe with lower acidity relative to Al(C₆F₅)₃ fully converted MMA to PMMA with $M_{\rm n} = 88.3 \text{ kg mol}^{-1}$, $D = 1.19 \text{ and } I^*\% = 91 \text{ in } 10 \text{ min (Table 1,}$ run 6). However, MALDI-TOF MS spectrum indicated that the final product still contained a cyclic chain end (Supporting Information, Figures S38 and S39). Using (BHT)₂AlⁱBu with similar acidity but larger steric hindrance to that of (BHT)₂AlMe, we could successfully suppress backbiting cyclization (Supporting Information, Figures S40 and S41) at the expense of polymerization activity (Table 1, run 7). Excitingly, switching to a (BHT)Al'Bu₂ MMA adduct with lower acidity but reduced steric hindrance led to the significantly enhanced polymerization activity, producing PMMA with $M_n = 92.8 \text{ kg mol}^{-1}$, D = 1.10, thus furnishing I*% = 86 (Table 1, run 9). Moreover, chain-end analyses revealed that the produced polymer is a linear, living chain, capped with IAP3/H chain ends, and showed no evidence for the formation of cyclic backbiting chain end (Figure 1). These results further highlight the importance of considering both electronic and steric factors when matching LA with LB for generating a suitable LP that can promote living polymerization. The living feature of the polymerization could be

Table 1: Results of polymerization by IAP-based LPs. [a]

Run	LB	LA	М	M/LB/LA	t	Conv. ^[b]	$M_n^{[c]}$	MWD ^[c]	/ ☆[d]
						[%]	$[kg mol^-]$		[%]
1	IAP1	$Al(C_6F_5)_3$	MMA	800:1:2	30 s	100	100	1.17	80
2	IAP2	$AI(C_6F_5)_3$	MMA	800:1:2	30 s	100	154	1.21	52
3	IAP3	$AI(C_6F_5)_3$	MMA	800:1:2	30 s	100	86.9	1.11	92
4	IAP4	$AI(C_6F_5)_3$	MMA	800:1:2	30 s	100	99.2	1.14	81
5	IAP5	$AI(C_6F_5)_3$	MMA	800:1:2	24 h	41	392	1.35	8
6	IAP3	(BHT)₂AlMe	MMA	800:1:2	10 min	100	88.3	1.19	91
7	IAP3	(BHT)₂Al ⁱ Bu	MMA	800:1:2	340 min	100	72.0	1.13	112
8	IAP3	(BHT)Al ⁱ Bu ₂ ·MMA	MMA	400:1:2	40 s	100	54.0	1.06	75
9	IAP3	(BHT)Al ⁱ Bu₂·MMA	MMA	800:1:2	2 min	100	92.8	1.10	86
10	IAP3	(BHT)Al ⁱ Bu₂·MMA	MMA	1600:1:2	10 min	100	153	1.06	105
11	IAP3	(BHT)Al ⁱ Bu ₂ ·MMA	MMA	3200:1:2	30 min	100	297	1.07	108
12	IAP3	(BHT)Al ⁱ Bu₂·MMA	MMA	6400:1:2	135 min	100	656	1.09	98
13	IAP3	(BHT)Al ⁱ Bu₂·MMA	MMA	10000:1:2	11 h	100	1029	1.08	97
14	IAP3	(BHT)Al ⁱ Bu ₂ ·MMA	MMA	15 000:1:2	21 h	100	1568	1.08	96
15	IAP3	(BHT)Al ⁱ Bu₂·MMA	MMA	20 000:1:2	48 h	89	1927	1.10	92
16	IAP3	(BHT)Al'Bu ₂ ·EMA	EMA	400:1:2	2 min	100	49.7	1.16	90
17 ^[e]	IAP3	(BHT)₂Al ⁱ Bu	MMBL	800:1:2	340 min	100	85.6	1.12	105
18	IAP3	(BHT)Al ⁱ Bu ₂	DMAA	800:1:2	5 min	100	833	1.64	
							44.4	1.39	
19	IAP3	(BHT)Al ⁱ Bu ₂	MS	400:1:2	24 h	16	n.d	n.d	n.d

[a] Conditions: carried out at RT in toluene using procedure A; for a 800MMA/1LB/2LA ratio, $[M] = [MMA]_0 = 0.936 \,\text{m}$ and $[LA]_0 = 2[LB]_0 = 2.34 \,\text{mm}$. n.d. = not determined. [b] Monomer conversions measured by ¹H NMR. [c] M_n and D determined by GPC relative to PMMA standards in DMF. [d] Initiator efficiency $(I^*)\% = M_n(\text{calcd})/M_n(\text{exptl}) \times 100$, where $M_n(\text{calcd}) = [\text{MW}(\text{MMA})]([\text{MMA}]_0/[\text{I}]_0)$ (conversion) + MW of chain-end groups. [e] Performed in CH₂Cl₂.

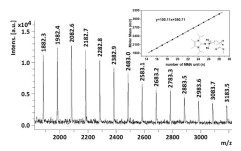


Figure 1. MALDI-TOF MS spectrum of the low-MW PMMA sample produced by IAP3/(BHT)AliBu, LP in toluene at RT. Inset: Plot of m/z values vs. the number of MMA repeat units (n).

confirmed by the linearly ($R^2 = 0.9996$; Supporting Information, Figure S35) increased M_n values of PMMA with an increase in [MMA]₀/[IAP3]₀/[(BHT)AlⁱBu₂·MMA]₀ ratio from 400:1:2 to 20000:1:2 (Table 1, runs 8–15), while \mathcal{D} values remained narrow (from 1.06 to 1.10; Figure 2a). Worth noting that UHMW PMMA with M_n up to 1927 kg mol⁻¹ were achieved by FLP for the first time (Table 1, run 13-15). The livingness and robustness also enable successful chain-extension experiments performed with the sequential addition of three batches (1000/1000/ 1000 equiv) of MMA (Figure 2b; Supporting Information, Table S5) and the synthesis of well-defined diblock copolymer PMMA-b-PEMA and triblock copolymer PMMA-b-PEMAb-PMMA (Supporting Information, Table S6, runs 2 and 3; Figure 2c) by using ethyl methacrylate (EMA) as co-monomer. Next, we investigated the effectiveness of IAP3-based LPs for polymerization of other monomers and found that it also exhibited well control on polymerization of both EMA

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Figure 2. GPC traces of a) PMMA produced by IAP3/(BHT)Al † Bu₂ LP at various [MMA]₀/[IAP3]₀/[(BHT)Al † Bu₂]₀ ratio at RT. Conditions: [MMA]₀/[IAP3]₀/[(BHT)Al † Bu₂]₀ = 400:1:2, 800:1:2, 1600:1:2, 3200:1:2, 6400:1:2, 10000:1:2, 15 000:1:2, 20 000:1:2, [MMA]₀ = 0.936 M and b) PMMA samples obtained from chain extension experiments in toluene at RT, [MMA]₀ = 0.47 M and c) homopolymer, diblock, and triblock copolymer produced from the sequential block copolymerization of MMA and EMA by IAP3/(BHT)Al † Bu₂ in toluene at RT: polymerizing MMA first, [MMA]₀ = 0.936 M.

(Table 1, run 16) and renewable monomer, γ -methyl- α -methylene- γ -butyrolactone (MMBL) (Table 1, run 17). Polymerization of N,N-dimethylacrylamide (DMAA) produced polymers with a bimodal MWD, indicating the coexistence of different active species (Table 1, run 18) whereas only 16% monomer conversion was obtained for polymerization of methyl sorbate (MS) in 24 h (Table 1, run 19).

To gain further insight into the above-described polymerization behavior, we systematically investigated polymerization by IAP3/(BHT)Al'Bu₂ LP system. It is generally accepted that a catalyst system should meet the following requirements to synthesize UHMW polymers: First, extraordinarily effective polymerization activity. Second, exclusive active species to initiate polymerization. Third, long lifetime polymerization performance and no interference with other side reactions such as chain-termination. As an organophosphorus superbase, the combination of IAP3 with (BHT)-AliBu₂ could achieve high to quantitative monomer conversion for polymerization with [MMA]₀/[IAP3]₀/ [(BHT)Al'Bu₂]₀ ratio ranging from 400:1:2 to 20000:1:2. Different from NHO alone could initiate polymerization of MMA,[14] IAP3 itself is completely ineffective whereas (BHT)Al'Bu2 only exhibits negligible effect for polymerization in up to 24 h (Supporting Information, Table S2). We performed a series of stoichiometric reactions at RT and observed the formation of classic Lewis adduct for the reaction of IAP3 and Al(C₆F₅)₃ (Supporting Information, Figures S16-S18) and the exclusive generation of non-interacting, true FLP for the reaction of **IAP3** with (BHT)₂AlMe, (BHT)₂AlⁱBu or (BHT)AlⁱBu₂, respectively (Supporting Information, Figures S19–S24). As revealed by ¹H and ³¹P NMR spectroscopy, the reaction of **IAP3** and (BHT)Al'Bu₂ MMA immediately and exclusively yielded a clean zwitterionic species IAP3- $CH_2C(Me)$ =C(OMe)OAl-(BHT)iBu2, which possessed analogous structure with that of IAP2-CH₂C(Me)=C(OMe)OAl(C_6F_5)₃ (Figure 3) generated from the reaction of IAP2 and Al(C₆F₅)₃·MMA adduct (Supporting Information, Scheme S1, Figures S30–S33). Above results indicated that such zwitterionic species is the exclusive active species for polymerization and there is no

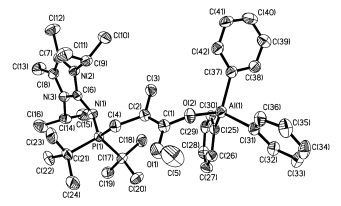


Figure 3. X-ray crystal structure of IAP2-CH₂C(Me)=C(OMe)OAl(C_6F_5)₃. Hydrogen and fluorine atoms are omitted for clarity and ellipsoids are set at 50% probability.

unwanted reaction resulting from other active species. Comparative studies performed with different activation procedures (see the Supporting Information for details of procedures A and B) afforded the exactly same PMMA as revealed by their completely overlapped GPC traces (Figure 4; Supporting Information, Table S3), which further confirmed the uniqueness of zwitterionic species. In addition to meet the requirement of no occurrence of side reaction (see above), IAP3/[(BHT)AlⁱBu₂ FLP system exhibited exceptionally long-life catalytic performance, which is significantly important for synthesis of UHMW polymer. The living polymerization system is immortal and could reinitiate the desired living polymerization after 400 equiv of MMA was fully consumed and the resulting system was held in the absence of free MMA at RT for 24 h by the sequential addition of another batch (400 equiv) of MMA, producing PMMA with $M_n = 96.1 \text{ kg mol}^{-1}$, D = 1.18 (Supporting Information, Table S4, run 2), which is similar with that $(M_n = 92.8 \text{ kg mol}^{-1}, D = 1.10, \text{ Table 1, run 8})$ obtained from polymerization in an 800:1:2 [MMA]₀/[IAP3]₀/ [(BHT)AlⁱBu₂·MMA]₀ ratio (Supporting Information, Figure S34). As discussed above, IAP3/(BHT)AliBu₂ FLP

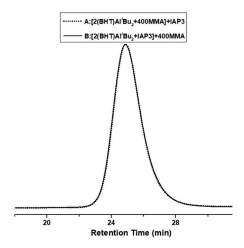


Figure 4. GPC traces of PMMA obtained from the polymerization performed using different activation procedures.

system meets all requirements for synthesis of UHMW polymers and thus accomplishing the goal.

In summary, the first example of UHMW PMMA with $M_n = 1927 \text{ kg mol}^{-1}$ and small D = 1.10 was produced from thenon-interacting, authentic IAP3/(BHT)AlⁱBu₂ FLP-promoted living polymerization of MMA at RT. Systematic investigation revealed that the combination of extraordinarily effective polymerization activity, exclusive zwitterionic active species and immortal polymerization process without observance of side reaction or chain termination enabled IAP3/ (BHT)AliBu, FLP system to accurately control over polymerization of MMA and achieve UHMW PMMA in the end. These findings not only provide significantly important foundation to the synthesis of UHMW polymer in the future, but also enable us to synthesize advanced polymers such as thermoplastic elastomer or self-assembling polymers by this highly effective and immortal catalyst system. Relevant research work is in progress.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: frustrated Lewis pairs \cdot living polymerization \cdot organoaluminum · organophosphorus superbases · ultrahigh-molecular-weight polymers

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