

REGIOSELECTIVITY INVESTIGATION FOR THE PYROLYSIS OF XANTHATES: A COMPUTATIONAL STUDY

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Received 17 February 2013

Accepted 14 July 2013

Published 23 October 2013

MP2/6-31+G(d,p)//MP2/6-31G(d) method was employed to investigate the pyrolyses of O-*sec*-butyl S-methyl xanthate (Chugaev reaction) and S-*sec*-butyl O-methyl xanthate, which gave regioselective products of E-butene, Z-butene and 1-butene. Both procedures were found to have 13 possible pathways, of which nine pathways would generate the alkene products. For O-*sec*-butyl S-methyl xanthate, the computational results indicated that the most favorable three pathways corresponded to a two-step mechanism, with the rate-determining step to be a thion sulfur atom involved six-membered ring transition states. The calculated products distribution was consistent with the experimental observations. However, for S-*sec*-butyl O-methyl xanthate, thiol-participated four-membered ring transition states were found to be more energetically favored than the six-membered ring transition state to produce 1-butene, which can be attributed to a larger sulfur atomic size than an oxygen atom. As the calculation result,

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only trace amount of 1-butene could be obtained with a major product being E-butene and Z-butene as a minority.

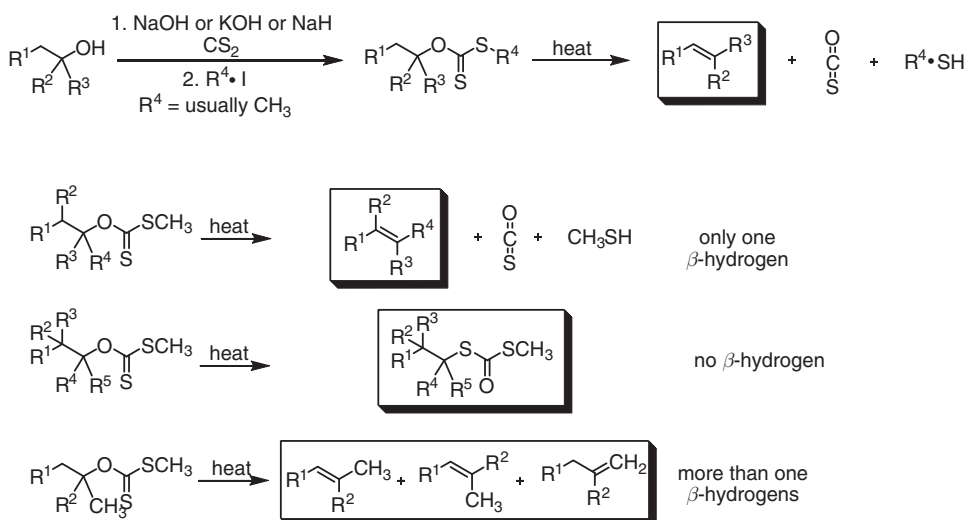
Keywords: MP2/6-31+G(d,p)//MP2/6-31G(d); O-*sec*-butyl S-methyl xanthate; S-*sec*-butyl O-methyl xanthate; regioselectivity, pyrolysis.

1. Introduction

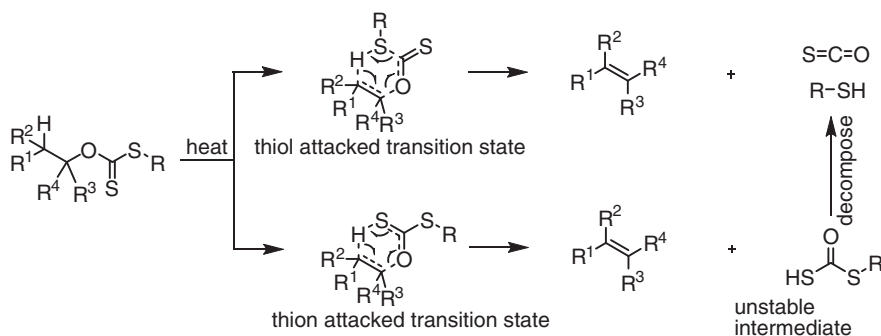
As a valuable synthetic route to synthesize alkenes without rearrangement of the carbon skeleton, pyrolysis of xanthate precursors was first reported by L. Chugeav in 1899 in their studies on the optical properties of xanthates.^{1,2} This method had attracted considerable interests in polymer chemistry for the synthesis of poly-paraphenylene-vinylene (PPV) samples, which had applied as promising characteristics for fabrication of light-emitting devices.³⁻⁶

Xanthates can be easily prepared by the reaction of the corresponding alcohols with a base and carbon disulfide, followed by trapping with an alkyl iodide (usually methyl iodide).⁷ At least one β -hydrogen of the xanthate is required to ensure that the pyrolysis will lead to the alkenes, together with the elimination of by-products (carbonyl sulfide and a thiol). However, if more than one β -hydrogen is present, regioselectivity will need to be considered, which will limit the application of the Chugeav reaction. Xanthates with no β -hydrogen will undergo thione-to-thiol rearrangements, giving S,S-dialkyl dithiocarbonates (Scheme 1).

The mechanism of the Chugeav reaction was proposed to involve two possible pathways (Scheme 2). The first pathway reacts in a one-step mechanism, yielding an alkene, a carbonyl sulfide and a thiol. In the second mechanism, a thione group, rather than a thiol group, will attack the β -hydrogen, giving an alkene and an



Scheme 1. Chugeav reaction with different β -hydrogens.



Scheme 2. Two proposed mechanisms for the pyrolysis of xanthates.

unstable S-alkyl dithiocarbonate intermediate, which will decompose to carbonyl sulfide and a thiol. The first pathway was supported by W. Hückel in 1940,⁸ while D. H. Barton⁹ and D. J. Cram¹⁰ preferred the second mechanism. Experimental studies were also applied to investigate the Chugeav mechanisms. Alexander and Mudrak proved the *cis*-elimination mechanism.^{11–13} Bader and Bourns¹⁴ made isotope effect studies for the pyrolysis of *trans*-2-methyl-1-indanyl xanthate and found solid evidence for the second pathway and further kinetic and thermodynamic studies concluded that the Chugeav reactions were homogenous and unimolecular.

As for the theoretical studies, Erickson and Kahn¹⁵ first calculated the activation energies of HSC(S)OEt at MP2/6-31G(d)//HF/6-31G(d) level in 1994. Deleuze group¹⁶ investigated the E_i reaction of EtSC(S)OMe and its substituent effects (such as α -CN, β -CN, *trans*-stilbene and *cis*-stilbene) at different theoretical levels (HF, B3LYP, MPW1K, MP2, MP3, MP4, CCSD, CCSD(T)) and different basis sets (6-31G*, 6-31G**, 6-311G**, 6-311++G**, cc-pVDZ). In comparison with all these methods and basis sets, DFT calculations gave with longer or shorter bond lengths and underestimated the activation energies, whereas HF method tended to significantly overestimate the activation energy. However, MP2 barriers matched nicely with benchmark CCSD(T) activation energies. Harano¹⁷ utilized B3LYP/6-31G(d) theoretical level to study the transition states of MeSC(S)OEt. Recently, Vélez group¹⁸ considered five possible mechanisms for the thermal decomposition of a series of xanthates at the level of MP2/6-31G(d). Similar theoretical studies were also applied to the other computational systems.^{19–23} However, all the previous studies focused on either the various computational methods or different transition states, none of them had considered the regioselectivity of the Chugeav reaction. As for the experimental results, Goeders group²⁴ pyrolyzed different xanthates by dropping into external heated glass columns. The products were collected by liquid nitrogen and analyzed by gas phase chromatography. The distributions of regioselective products were determined titrimetrically or gravimetrically. Of all the xanthates, O-*sec*-butyl S-methyl xanthate gave products of Z-butene, E-butene and 1-butene with a distribution ratio to be 40:19:14, after normalization (the original ratio is

40:19:41), which was also selected in this paper, together with S-*sec*-butyl O-methyl xanthate, to investigate the regioselectivity of the Chugeav reaction at the computational level of MP2/6-31+G(d,p)//MP2/6-31G(d).

2. Calculation Methods

All calculations were performed with the help of Gaussian 03 package.²⁵ As mentioned by Deleuze group¹⁶ that the MP2 level²⁶ gave nice activation barriers for the pyrolysis of xanthates as the CCSD(T) method, which was also applied by Vélez¹⁸ with the basis set of 6-31G(d),²⁷ the same computational level was also selected for the structural optimization. For the transition state calculations, force constants were also calculated with MP2/6-31G(d) method and noeigentest was applied to facilitate the transition state optimization. A higher basis set of 6-31+G(d,p) was employed to calculate the single-point energy based on the optimized structures in order to obtain more accurate results. The frequency calculations were evaluated at 623.15 K, as the experimental temperature.²⁴ A scaling factor²⁸ of 0.9670 for the zero-point vibrational energies was used for correction. All zero-gradient structures were characterized by a vibrational analysis with no imaginary frequency. All of the transition-state structures had only one imaginary frequency, which was interpreted as a negative vibrational mode, and the intrinsic reaction coordinate (IRC)²⁹ was followed to make sure that each transition state connects the expected reactant and product.

Classic transition-state theory (TST)^{30,31} was selected to calculate the relative rate constant, $k(T)$, for the rate-determining steps, which was also proportional to the product distributions. The equation is shown below, which assumes that the transmission coefficient is equal to unity:

$$k(T) = \frac{k_B T}{h} e^{\frac{-\Delta G^\ddagger(T)}{RT}},$$

where k_B , h and R are the Boltzmann constant, Planck's constant and the universal gas constant, respectively, and $\Delta G^\ddagger(T)$ is the standard free energy of activation at the absolute temperature T .

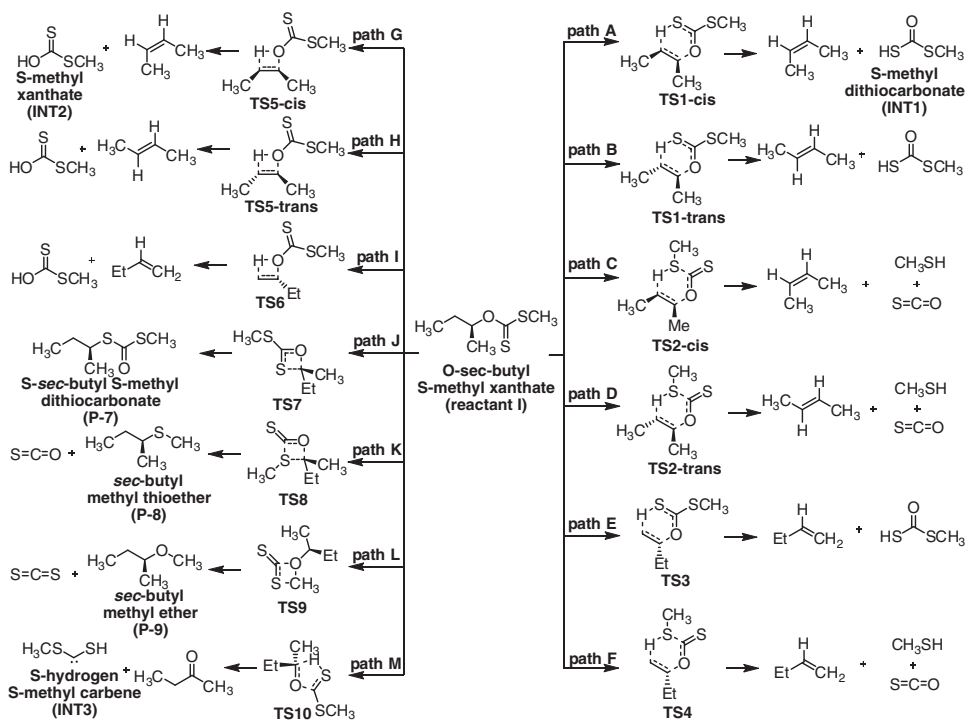
3. Results and Discussion

Five possible mechanisms had been calculated by Vélez group for the pyrolysis of O-alkyl (Et, *i*-Pr and *t*-Bu) S-methyl xanthates and S-alkyl (Et, *i*-Pr and *t*-Bu) O-methyl xanthates.¹⁸ Of all the five mechanisms, only two mechanisms with six-membered ring transition states resulted in the desired alkene products, while ignoring that four-membered ring transition states could also give alkene products. Besides, five-membered ring transition states were also found to give ketones or thioketones. Herein, we presented a detailed mechanism study for the thermal decomposition of O-*sec*-butyl S-methyl xanthate and S-*sec*-butyl O-methyl xanthate.

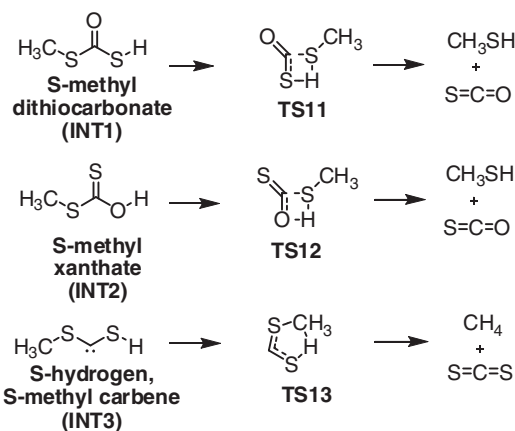
3.1. Pyrolysis of *O*-*sec*-butyl *S*-methyl xanthate

The MP2/6-31+G(d,p)//MP2/6-31G(d) calculated results for the pyrolysis of *O*-*sec*-butyl *S*-methyl xanthate in the gas phase were shown in Schemes 3 and 4.

As can be seen from the schemes, 13 possible pathways were considered for the mechanism study, of which pathways A to I ended up with the Chugeav reaction products. In pathways A, B and E, a thion sulfur atom attacked a β -hydrogen *via* a six-membered ring transition state, giving the desired *Z*-butene, *E*-butene and 1-butene, together with an unstable intermediate, *S*-methyl dithiocarbonate (INT1), which rapidly decomposed into carbonyl sulfide and methanethiol. For pathways C, D and F, the same products would be obtained, but a thiol sulfur atom was participated with a one-step process. Besides a sulfur atom, an oxygen atom could also act as a nucleophile, as shown in pathways G, H and I, to extract a β -hydrogen atom through a four-membered ring transition state. Herein, *S*-methyl xanthate (INT2) was generated as an intermediate, which also pyrolyzed into methanethiol and carbonyl sulfide. Pathway J resulted in a thion-to-thiol rearrangement product, *S*-*sec*-butyl *S*-methyl dithiocarbonate (P-7). Pathways K and L gave off carbonyl sulfide and carbon disulfide, leaving *sec*-butyl methyl thioether (P-8) and *sec*-butyl methyl ether (P-9), respectively. Pathway M was revealed with a five-membered transition state



Scheme 3. Thirteen possible pathways for the thermolysis of *O*-*sec*-butyl *S*-methyl xanthate.



Scheme 4. Thermal decomposition of INT1, INT2 and INT3.

to give butanone and S-hydrogen S-methyl carbene (INT3), which was unstable and easily decomposed into methane and carbon disulfide.

The corrected Gibbs energies for the reactants, transition states and products involved in the pyrolysis of *O-sec*-butyl S-methyl xanthate were collected in Table 1 at MP2/6-31+G(d,p)//MP2/6-31G(d) level, while the detailed electronic energies, zero-point vibrational energies, thermal corrections to enthalpies, entropies and the calculation methods were displayed in Table S1 in the supporting information. The Gibbs free energy profiles for all pathways were depicted in Figs. 1 and 2.

According to Figs. 1 and 2, the lowest barriers for the pyrolysis of *O-sec*-butyl S-methyl xanthate corresponded to pathways B, A and E, all of which were exothermic pathways with late transition states of TS1-trans, TS1-cis and TS3. The activation energies for TS1-trans, TS1-cis and TS3 were 38.30, 39.65 and 40.54 kcal·mol⁻¹, respectively, with the differences of only 1.35 kcal·mol⁻¹ (TS1-cis VS TS1-trans) and 0.89 kcal·mol⁻¹ for TS3 versus TS1-cis. According to above

Table 1. Corrected Gibbs energies, in Hartrees, for the reactants, transition states and products involved in the pyrolysis of *O-sec*-butyl S-methyl xanthate at MP2/6-31+G(d,p)//MP2/6-31G(d).

Species	Corrected G	Species	Corrected G	Species	Corrected G
reactant I	-1105.3364184	TS8	-1105.2376825	P-9	-272.0430996
TS1-cis	-1105.2732393	TS9	-1105.2123258	INT1	-948.6994613
TS1-trans	-1105.2753856	TS10	-1105.2222227	INT2	-948.6877888
TS2-cis	-1105.2548255	TS11	-948.6426505	INT3	-873.5555020
TS2-trans	-1105.2552343	TS12	-948.6279945	butanone	-231.7332780
TS3	-1105.2712175	TS13	-873.5104950	CH ₃ OH	-115.3972433
TS4	-1105.2529187	Z-butene	-156.6593760	CH ₃ SH	-438.0027460
TS5-cis	-1105.2520597	E-butene	-156.6604383	CH ₄	-40.3634193
TS5-trans	-1105.2537616	1-butene	-156.6560539	S=C=S	-833.3169485
TS6	-1105.2503782	P-7	-1105.3559477	O=C=S	-510.7380868
TS7	-1105.2591576	P-8	-594.6560415		

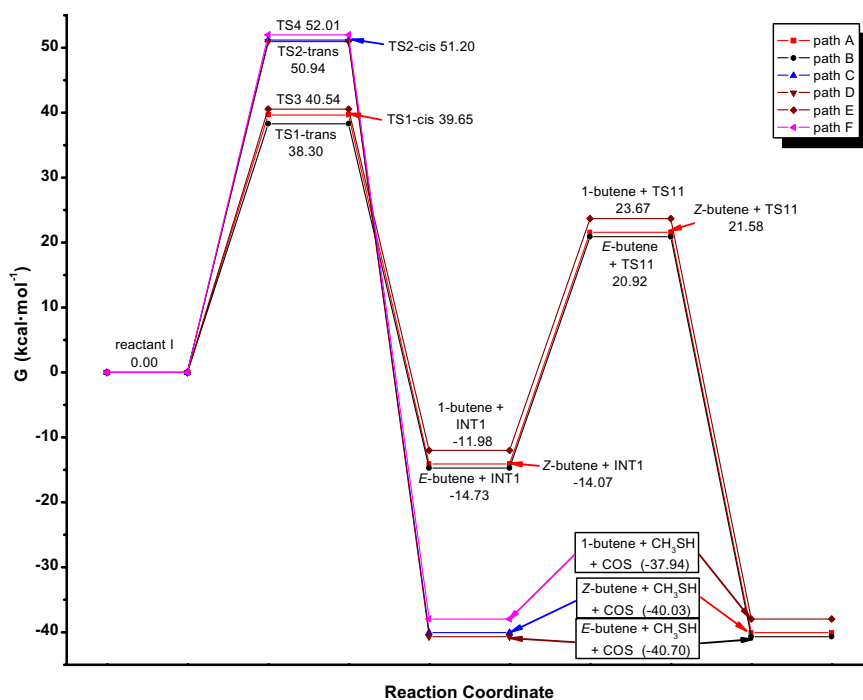


Fig. 1. Gibbs energy profile of pathways A to F for the pyrolysis of *O-sec-butyl S-methyl xanthate*.

mentioned classic TST equation, the relative rate constants were calculated to be 40:13:7 for TS1-trans, TS1-cis and TS3. This theoretical result agreed with the experimental observation, of which three alkenes were measured to be 40:19:14 for *E*-butene, *Z*-butene and 1-butene after normalization.²⁴ All three transition states produced the corresponding alkene together with a same intermediate, *S*-methyl dithiocarbonate (INT1). INT1 would decompose into methanthiol and carbonyl sulfide *via* a four-membered ring transition state with the energy barrier to be 35.65 kcal·mol⁻¹. The reaction energies for the whole procedures were found to be -40.70, -40.03 and -37.94 kcal·mol⁻¹, respectively, indicating that they were all exothermic.

For TS2-cis, TS2-trans and TS4 (Fig. 1), the corresponding activation energies raised to 51.20, 50.94 and 52.01 kcal·mol⁻¹, which was consistent with the experimental observation for the isotope effect of ³²S/³⁴S⁸ and clearly indicated that a thion sulfur atom was highly preferred over a thiol sulfur atom to attack a β -hydrogen. For the four-membered ring transition states (Fig. 2), TS5-cis, TS5-trans and TS6 had activation energies of 52.94, 51.87 and 53.99 kcal·mol⁻¹, resulting in *Z*-butene, *E*-butene, 1-butene and INT2. INT2 would convert to methanthiol and carbonyl sulfide *via* TS12 with a 37.52 kcal·mol⁻¹ energy barrier. TS7, TS8 and TS9 were similar transition states as E. Vélez group considered.¹⁸ TS7 was 48.48 kcal·mol⁻¹ higher than reactant I, producing P-7. TS8 (61.96 kcal·mol⁻¹) and

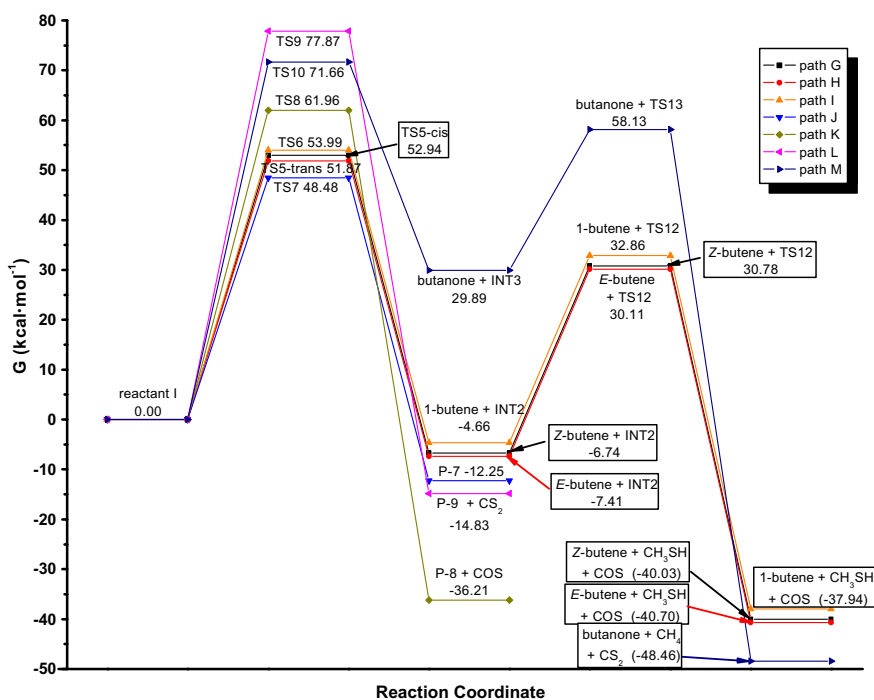


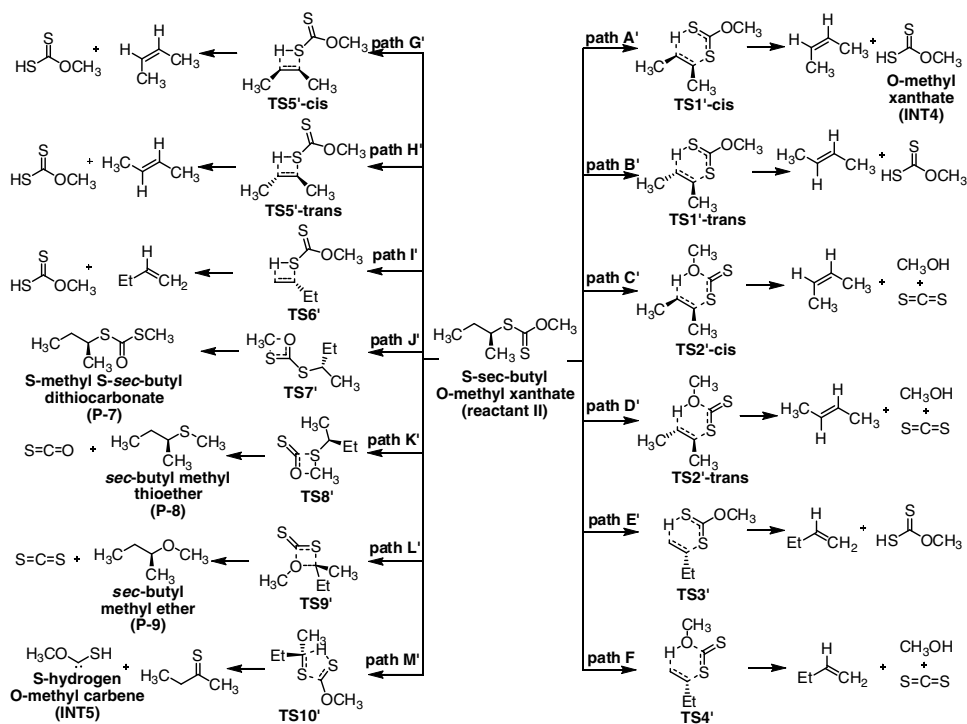
Fig. 2. Gibbs energy profile of pathways G to M for the pyrolysis of *O-sec-butyl S-methyl xanthate*.

TS9 (77.87 kcal·mol⁻¹) generated P-8 and P-9. The five-membered ring transition state (TS10) had activation energy as high as 71.66 kcal·mol⁻¹, giving butanone and INT3. INT3 decomposed into methane and carbon disulfide *via* TS13 (28.24 kcal·mol⁻¹) with an exothermicity of 37.94 kcal·mol⁻¹.

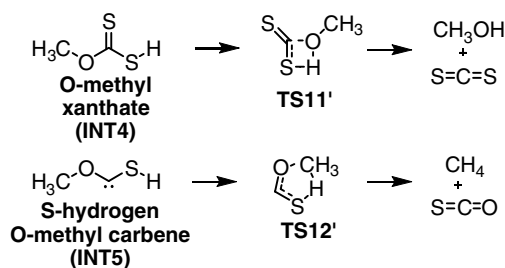
3.2. Pyrolysis of *S-sec-butyl O-methyl xanthate*

Similar to the thermal decomposition of *O-sec-butyl S-methyl xanthate*, the mechanisms for the pyrolysis of *S-sec-butyl O-methyl xanthate* were displayed in Schemes 5 and 6. Table 2 collected the corrected Gibbs energies for the reactants, transition states and products involved in the pyrolysis of *S-sec-butyl O-methyl xanthate* at MP2/6-31+G(d,p)//MP2/6-31G(d) level, while the detailed electronic energies, zero-point vibrational energies, thermal corrections to enthalpies, entropies and the calculation methods were listed in Table S2 in the supporting information. Meanwhile, Figs. 3 and 4 displayed the free energy profiles for all pathways.

About 13 possible pathways were also found for the thermal decomposition of *S-sec-butyl O-methyl xanthate*. For pathways A' to I', Z-butene (for pathways A', C' and G'), E-butene (for pathways B', D' and H') and 1-butene (for pathways C' F' and I') were obtained, together with same by-products, methanol and carbon disulfide. Pathway J' underwent a thion-ether to thiol-carbonyl rearrangement, giving



Scheme 5. 13 possible pathways for the pyrolysis of S-sec-butyl O-methyl xanthate.



Scheme 6. Thermal decomposition of INT4 and INT5.

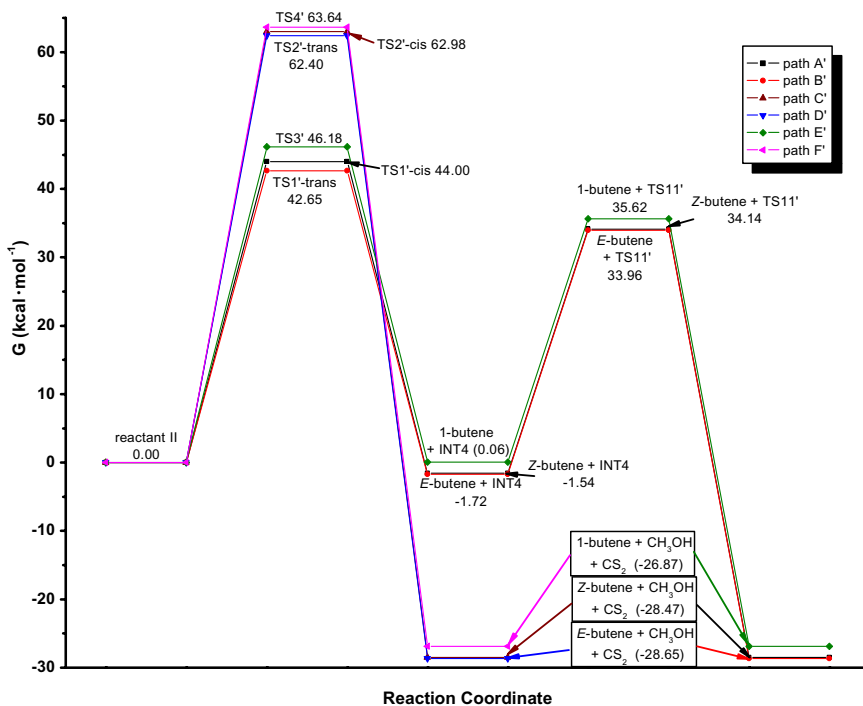
S-methyl S-sec-butyl dithiocarbonate (P-7). Pathways K' and L' resulted in sec-butyl methyl thioether and sec-butyl methyl ether with the emission of carbonyl sulfide and carbon disulfide. Pathway M' passed through a five-membered ring transition state, giving butane-2-thione and S-hydrogen O-methyl carbene (INT5), which decomposed into methane and carbonyl sulfide.

As depicted in Figs. 3 and 4, the lowest six transition states were TS1'-trans (42.65 kcal·mol⁻¹), TS1'-cis (44.00 kcal·mol⁻¹), TS5'-cis (46.04 kcal·mol⁻¹), TS5'-trans (46.08 kcal·mol⁻¹), TS3' (46.18 kcal·mol⁻¹) and TS6' (47.08 kcal·mol⁻¹). It

Table 2. Corrected Gibbs energies, in Hartrees, for the reactants, transition states and products involved in the pyrolysis of *S-sec-butyl O-methyl xanthate* at MP2/6-31+G(d,p)//MP2/6-31G(d).

Species	Corrected G	Species	Corrected G	Species	Corrected G
reactant II	-1105.3281974	TS7'	-1105.2447600	P-8	-594.6560415
TS1'-cis	-1105.2580784	TS8'	-1105.2180214	P-9	-272.0430996
TS1'-trans	-1105.2602262	TS9'	-1105.2190422	INT4	-948.6712764
TS2'-cis	-1105.2278313	TS10'	-1105.2163551	INT5	-550.9532339
TS2'-trans	-1105.2287505	TS11'	-948.6144155	CH ₃ OH	-115.3972433
TS3'	-1105.2545976	TS12'	-550.9079445	CH ₄	-40.3634193
TS4'	-1105.2267838	Z-butene	-156.6593760	S=C=S	-833.3169485
TS5'-cis	-1105.2548272	E-butene	-156.6596631	O=C=S	-510.7380868
TS5'-trans	-1105.2547608	1-butene	-156.6568291	butane-1-thione	-554.3133414
TS6'	-1105.2531660	P-7	-1105.3559477		

should be mentioned that TS5'-cis, TS5'-trans and TS6' had much lower energy barriers as compared with TS2'-cis (62.98 kcal·mol⁻¹), TS2'-trans (62.40 kcal·mol⁻¹) and TS4' (63.64 kcal·mol⁻¹), which was different from TS5-cis, TS5-trans and TS6 in the pyrolysis of *O-sec-butyl S-methyl xanthate*. This could be attributed to the atomic sizes of different nucleophiles. Sulfur had a much larger atomic size than oxygen, which could dramatically release the ring strain of the transition states. Both TS1'-trans and TS5'-trans produced E-butene, together with O-methyl

Fig. 3. Gibbs energy profile of pathways A' to F' for the pyrolysis of *S-sec-butyl O-methyl xanthate*.

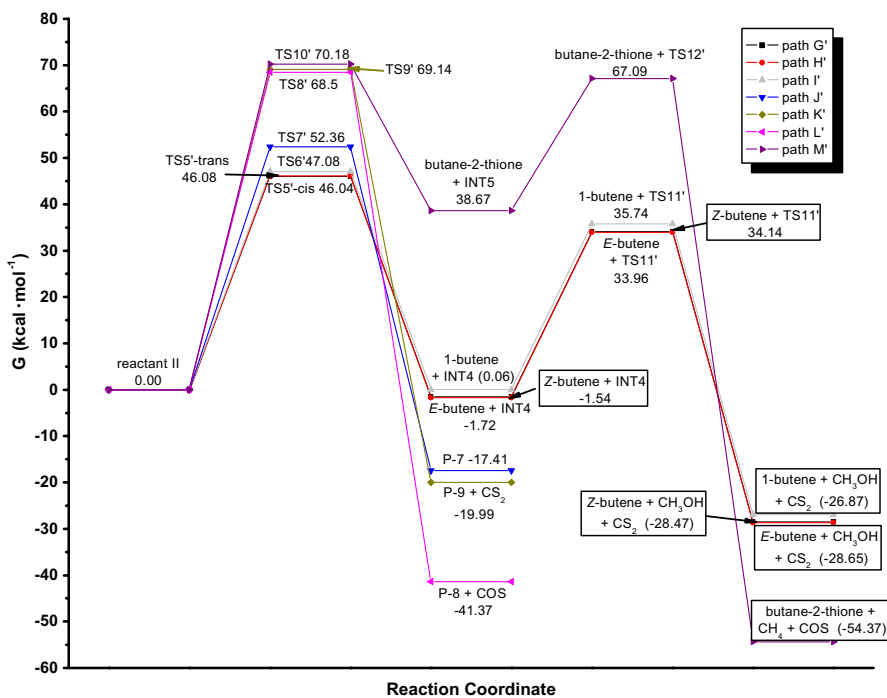


Fig. 4. Gibbs energy profile of pathways G' to M' for the pyrolysis of S-sec-butyl O-methyl xanthate.

xanthate (INT4), which converted to methanol and carbon disulfide through TS11' (35.68 kcal·mol⁻¹). The final products had exothermicity of -28.65 kcal·mol⁻¹ as compared with reactant II. Z-butene could be obtained from TS1'-cis and TS5'-cis, of which TS1'-cis was 1.35 kcal·mol⁻¹ higher than TS1'-trans and TS5'-cis was slightly lower (0.04 kcal·mol⁻¹) than TS5'-trans. The final products were also exothermic of -28.47 kcal·mol⁻¹. TS3' and TS6' gave 1-butene as the product with activation energies of 46.18 kcal·mol⁻¹ and 47.08 kcal·mol⁻¹, which were 3.53 kcal·mol⁻¹ and 4.43 kcal·mol⁻¹ higher than TS1'-trans, respectively. The final products were 26.87 kcal·mol⁻¹ lower than reactant II. As discussed above, E-butene would be collected as the major product with a minor product to be Z-butene, while trace amount of 1-butene could be obtained.

For the rest four mechanisms, TS7' had an energy barrier of 52.36 kcal·mol⁻¹, but the rearranged product could give off 69.77 kcal·mol⁻¹, resulting in an exothermic procedure. The following two transition states (TS8' and TS9') had even higher activation energies of 68.50 kcal·mol⁻¹ and 69.14 kcal·mol⁻¹, respectively, both of which resulted in the same products as TS8 and TS9. For TS10', it had the highest activation energy of 70.18 kcal·mol⁻¹, resulting in butane-2-thione and INT5, which decomposed into methane and carbonyl sulfide *via* TS12' (28.42 kcal·mol⁻¹) with exothermicity of 93.04 kcal·mol⁻¹, making the total pathway to be 54.37 kcal·mol⁻¹ lower than reactant II.

4. Conclusion

The pyrolyses of O-*sec*-butyl S-methyl xanthate and S-*sec*-butyl O-methyl xanthate were calculated at the MP2/6-31+G(d,p)//MP2/6-31G(d) level. Thirteen possible pathways were considered for both procedures. In the case of O-*sec*-butyl S-methyl xanthate, the lowest energy pathways, B, A and E, corresponded to the formation of E-butene, Z-butene and 1-butene *via* a two-step mechanism with the first step being a rate-determining step. This theoretical result was consistent with experimental product distribution. As for S-*sec*-butyl O-methyl xanthate, pathways G', H' and I' were found to be favored over pathway E', which could be attributed to a larger sulfur atomic size than an oxygen atom. Besides, TS3' and TS6' were 3.53 kcal·mol⁻¹ and 4.43 kcal·mol⁻¹ higher than TS1'-*trans*, which would result in trace amount of 1-butene with E-butene to be a major product and Z-butene as a minor product.

Supplementary Materials

Tables S1 and S2 illustrated the calculated electronic energies, zero-point vibrational energies, thermal corrections to enthalpies, entropies, and the corrected free energies for all species at the level of MP2/6-31+G(d,p)//MP2/6-31G(d). The optimized Cartesian coordinations for all species were collected in Tables S3 and S4. The IRC for all transition states were depicted in Figs. S1-S31.

Acknowledgment

This work is financially supported by Natural Science Foundation of Shandong Province of China (ZR2009BL004), National Science Foundation of China (21272236, 21207136) and Natural Science Fund of Education Department of Anhui province (1208085QB32).

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