

Preparation of Trifluoroiodomethane *via* Vapour-Phase Catalytic Reaction Between Pentafluoroethane and Iodine

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(Received: 4 August 2012;

Accepted: 18 February 2013)

AJC-13031

A new route for preparing CF₃I has been developed *via* a reaction between C₂HF₅ and I₂. The influence of reaction temperature and active components of the catalysts on the amount of CF₃I was investigated. The result suggests that the selectivity of the CF₃I can be controlled by reaction conditions and active component of catalyst. The process for the formation of CF₃I and by-products is also discussed.

Key Words: Trifluoriodomethane, Vapour-phase catalytic, Pentafluoroethane, Difluorocarbene, Trifluoromethyl radical.

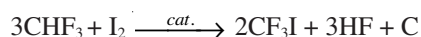
INTRODUCTION

Trifluoriodomethane (CF_3I) possesses a short atmospheric lifetime and low global warming potential^{1,2}, which makes it become the preferred alternative for halon and other halohydrocarbons in the application of fire extinguishing agent. Of note, CF_3I is expected for various other applications³⁻⁶, such as freezing medium, etching gas and raw materials for fluorochemicals, *etc.*

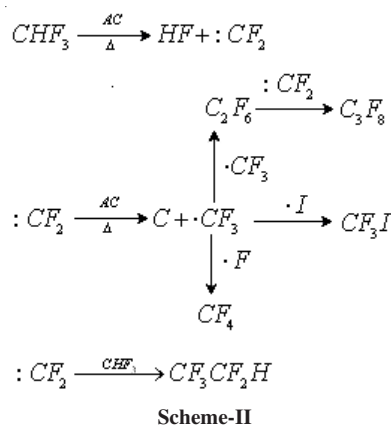
Nagasaki *et al.*^{7,8} have reported a continuous vapour-phase catalytic process for the manufacture of CF₃I *via* the reaction between trifluoromethane (CHF₃) and iodine (I₂), which makes the CF₃I industrialization possible (**Scheme-I**). Yang *et al.*^{9,10} have investigated the above mentioned synthetic mechanism of CF₃I and by products, and suggested that the reaction proceeds *via* difluorocarbenes (CF₂ carbenes) formed on the catalyst surface as intermediates, followed by CF₂ carbene disproportionation to trifluoromethyl (CF₃) radicals, furthermore CF₃ radical react with I₂ to form CF₃I (**Scheme-II**). But the raw material CHF₃ will be stopped little by little based on the montreal protocol, because the global warming potential of CHF₃ is as high as 12000 and the atmospheric lifetime is as long as 250 years. The raw material pentafluoroethane (C₂HF₅) possesses a low global warming potential and the atmospheric lifetime is only 33 years¹¹.

In our previous report¹², a consideration of the reaction mechanism for the formation of pentafluoroethyl iodide (C_2F_5I)

via the reaction between C_2HF_5 and I_2 suggested the reaction proceeds also *via* carbenes formed on the catalyst surface as intermediates. Based on this investigation, a new route for preparing CF_3I was developed *via* a vapour-phase catalytic reaction between C_2HF_5 and I_2 . We reported here the detailed study on the effect of catalytic activities especially the selectivity of the CF_3I to reaction conditions and catalysts, and the process for the formation of CF_3I and by-products is also discussed.



Scheme-I



EXPERIMENTAL

Preparation of catalyst: A commercial wooden AC (Tanxin Activated Carbon Co. Ltd., Shanghai; specific surface area 1322 m²/g) was used as the catalyst support in this study. After HF (2 M) washing, the AC had an ash content lower than 1.2 wt %, then AC was subjected to 2 M HCl for about 24 h at room temperature in order to decompose the carboxyl and lactone groups, increase the relative amounts of the phenolic and carbonyl groups¹³⁻¹⁵, and then the AC was washed with distilled water and dried in air at 103-105 °C overnight. After that, the sample was added to the aqueous metal salts solution, impregnated at room temperature for about 6 h and dried by a rotary evaporator at about 80 °C for 4 h. The pre-dried catalyst was then calcined at 600 °C for 2 h under nitrogen flow.

Synthesis of CF₃I: 15 mL catalysts were packed into a fixed bed reactor and were heated to 550 °C under nitrogen. Then nitrogen flow was stopped and C₂HF₅ was introduced to reactor at a rate of 60 mL/min. After 2 h, I₂ was provided by bubbling C₂HF₅ into molten I₂ which was placed in a stainless steel evaporator and heated up to 190 °C in an oil bath. The products were neutralized by Ca(OH)₂ solution, dried with CaCl₂ and analyzed by GC-MS. The results were listed as follows:

1. CF₄, *m/z*: 69, ⁺CF₃; 50, ⁺CF₂; 31, ⁺CF; 19, ⁻F.
2. CF₃CF₃, *m/z*: 119, ⁺CF₃CF₂; 100, ⁺CF₂CF₂; 69, ⁺CF₃; 50, ⁺CF₂; 31, ⁺CF.
3. CHF₃, *m/z*: 69, ⁺CF₃; 51, ⁺CHF₂; 31, ⁺CF.
4. CF₃CF₂CF₃, *m/z*: 169, ⁺CF₃CF₂CF₂; 119, ⁺CF₃CF₂; 100, ⁺CF₂CF₂; 69, ⁺CF₃; 50, ⁺CF₂; 31, ⁺CF.
5. CF₃I, *m/z*: 196, ⁺CF₃I; 177, ⁺CF₂I⁺; 127, ⁻I; 69, ⁺CF₃; 31, ⁺CF;
6. CF₃CF₂I, *m/z*: 246, ⁺CF₃CF₂I; 227, ⁺CF₂CF₂I; 177, ⁺CF₂I⁺; 158, ⁺CFI; 127, ⁻I; 119, ⁺CF₃CF₂; 100, ⁺CF₂CF₂; 69, ⁺CF₃; 50, ⁺CF₂; 31, ⁺CF.

Characterization of samples: Products were analyzed by means of GC-1002 produced by Beijing analytical instrument factory. The capillary column was a CP-PoraPLOTQ with 0.32 mm diameter and 30 m length from J & W Scientific Inc. The column was programmed as follows: the initial temperature was set at 60 °C for 6 min; then the temperature was increased at the rate of 40 °C/min and finally reached to 200 °C and held for 11 min. The instrumental parameters were set up as follows: injection port temperature, 200 °C; FID detector, 250 °C; the carrier gas was N₂ (0.1 MPa). The products were identified by comparison of their GC retention times and mass spectra with authentic samples. Quantitative analysis of the product standard ratios was obtained by comparison with mixtures prepared for calibration purposes. The relative response coefficients and retention time of each compound are given in Table-1.

TABLE-1
RELATIVE RESPONSE COEFFICIENTS
(*k_f/k_{std}*) DETERMINED BY GC-FID

Sample	Retention time/min	<i>k_f/k_{std}</i>
CHF ₃	8.3	3.6
CF ₃ CHF ₂	8.5	1.0
CF ₃ I	9.5	1.45
C ₂ F ₅ I	10.4	1.29

GC-MS (EI, 70 eV) spectra were measured using the Shimadzu GCMS-QP2010 system equipped with GC-2010. The column was CP-PoraPLOT Q with 0.32 mm i.d. and 30 m length from J & W Scientific Inc. The column was programmed as above-mentioned GC conditions. Injection port temperature, 200 °C; the carrier gas rate, 4.1 cm³ He/min.

RESULTS AND DISCUSSION

Synthesis of CF₃I on different temperature: Effect of the reaction temperature on product distribution in the gas-phase catalytic reaction for the synthesis of CF₃I between C₂HF₅ and I₂ in the presence of K₂CO₃/AC catalyst was examined. The results were shown in Fig. 1 and the product distribution was listed in Table-2.

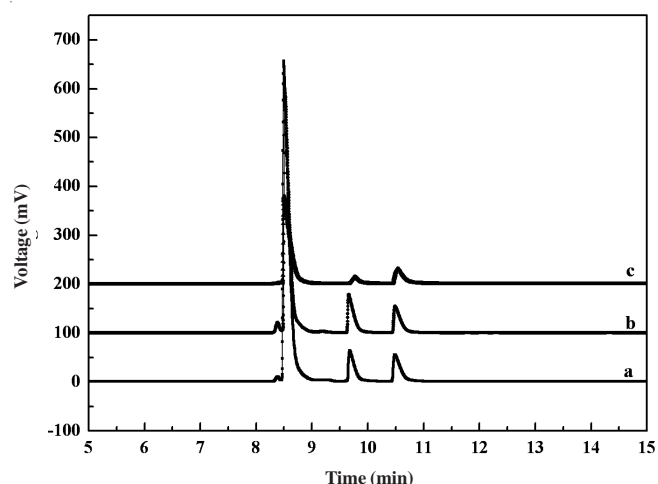


Fig. 1. GC spectrum of gas products; a-reaction at 500 °C, b- reaction at 550 °C, c- reaction at 600 °C

TABLE-2
EFFECT OF REACTION TEMPERATURE
ON PRODUCT DISTRIBUTION

Temperature (°C)	Ratio of relative volume of products (%)				
	C ₂ HF ₅	CHF ₃	CF ₃ I	C ₂ F ₅ I	Others
450	100	-	-	-	-
500	79.4	1.4	8.4	10.1	0.7
550	68.7	0.4	17.2	13.7	-
600	68.2	1.2	8.9	19.8	1.9

- - no products detected

The conversion of C₂HF₅ increased with the increasing reaction temperature and when the reaction temperature was 450 °C, reaction did not occur and only raw material C₂HF₅ was detected in gas product. When the reaction temperature was 550 °C, the conversion of C₂HF₅ reached 31.3 % and CF₃I was produced in 17.2 % relative volume of the gaseous products for 4 h, accompanying with CF₄, CHF₃, C₂F₆, C₃F₈ and C₂F₅I as by-products. But the ratio of CF₃I formed gradually decreased when the reaction temperature was 600 °C, the concentration of CF₃I reached only in 8.4 % (Table-2). It is suggested that the selectivity of the CF₃I is controlled by reaction conditions.

Synthesis of CF₃I on different catalysts: The tested catalysts were made from various alkali metal salts and alkaline earth metal salts with AC as a supporter. The catalytic activities of these catalysts were tested after 4 h reaction (Table-3).

TABLE-3
EFFECTS OF ALKALI METAL AND ALKALINE EARTH METAL
SALT CATALYSTS ON THE REACTION OF C₂HF₅ AND I₂

Component	C ₂ HF ₅ conversion (%)	CF ₃ I selectivity (%)	C ₂ F ₅ I selectivity (%)
NaNO ₃	11.2	0	0
KNO ₃	18.7	53.5	29.0
RbNO ₃	30.4	44.0	40.0
CsNO ₃	17.3	36.5	32.5
KF	22.0	49.5	48.7
RbF	18.5	34.0	51.3
CsF	<1	0	0
K ₂ CO ₃	31.3	54.9	43.8
Mg(NO ₃) ₂	5.9	13.6	12.4
Ba(NO ₃) ₂	9.8	20.1	11.9

Reaction temperature: 550 °C; C₂F₅H/I₂ (mol ratio):1.5; SV: 130 h⁻¹

As shown in Table-3, alkali metal salts shows a higher catalytic activity than the alkaline earth metal salts. This is because the electronegativity of alkali metal is less than the alkaline earth metal and the basicity of the alkali metal oxide decomposing from alkaline metal is better than alkali earth metal oxide decomposing from alkaline earth metal. Zhu *et al.*¹⁶ have reported that the decomposition of alkaline earth metal nitrates becomes more difficult with the atomic number increases. The highest activity of the alkali metal catalysts is RbNO₃ and the highest selectivity is potassium such as KF and K₂CO₃. While the anions, counterion to the metal cations, have also effect on the reaction rate, NO₃⁻ with high selectivity and F⁻ with high activity.

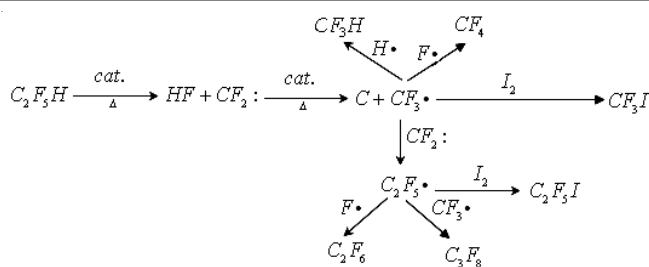
Based on exhaustive studies, it is found that the most suitable catalyst for this reaction is the synergistic combination of KF with high selectivity and RbNO₃ with high activity, especially the mass ratio of RbNO₃ and KF is 3:1 (Table-4). This ensures that the selectivity of the CF₃I is also controlled by active component of catalyst.

TABLE-4
EFFECT OF THE RATIO OF TWO ACTIVE COMPONENTS
ON CATALYTIC ACTIVITY

KF/RbNO ₃ (mass ratio)	C ₂ HF ₅ conversion (%)	CF ₃ I selectivity (%)	C ₂ F ₅ I selectivity (%)
1:1	34.6	50.2	32.5
1:2	46.0	42.4	41.2
2:1	40.0	43.8	39.3
1:3	46.5	53.1	28.3
3:1	38.3	51.2	32.8

Reaction temperature: 550 °C; C₂F₅H/I₂ (mol ratio):1.5; SV: 130 h⁻¹

The compounds containing CF₃ group such as CHF₃, CF₄, C₂F₆, C₃F₈, C₂F₅I and CF₃I were formed. We propose the following process for the formation of CF₃I and by products. As shown in **Scheme-III**, C₂HF₅ is decomposed to form the intermediates CF₂ carbene. Then CF₂ carbene is absorbed on the surface of the catalysts and disproportionation to CF₃ radicals, furthermore CF₃ radical react with I₂ to form CF₃I, whereas with H and F to form CHF₃ and CF₄. H radical¹⁷⁻¹⁹ maybe comes from carboxyls and phenols on the AC surface and F radical⁹ comes from the decomposing of CF₂ carbene on heat. Alternatively, CF₃ radical reacts with CF₂ carbene to form CF₃CF₂ radical, which generates C₂F₅I, C₂F₆ and C₃F₈.



Scheme-III: Suggested process for the formation of CF₃I and by-products

Conclusion

A new method for the preparation of CF₃I has been developed *via* a reaction between C₂HF₅ and I₂. The effect of the temperature on the synthesis of CF₃I was studied and the results indicated that the conversion of C₂HF₅ increased with the increasing reaction temperature, but the ratio of CF₃I formed gradually decreased. Also the conversion of C₂HF₅ and the selectivity of CF₃I reached 46.5 % and 53.1 % respectively when the reaction was conducted in the presence of Rb-K/AC catalyst with the mass ratio of 3:1 (RbNO₃:KF). The results suggest that the selectivity of the CF₃I can be controlled by reaction conditions and active component of catalyst. Furthermore, the catalyst promotes disproportionation of CF₂ carbene to generate CF₃ radical, which reacts with I₂ to form CF₃I.

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