Carbonization Mechanisms of PET



The Chemical Understanding

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INTRODUCTION



The understanding of charring mechanisms in polymer thermal decomposition is of paramount importance because it may provide a more suitable, environmentally friendly approach to fire retardance of polymer materials. Indeed, charring occurs in competition and at the expenses of formation of volatile combustible products from the polymer exposed to fire conditions. For what concerns the PET:

•No exhaustive literature can be found regarding the PET carbonization mechanism.

• Disagreements exist about the primary chain-scission mechanism due to thermal degradation.

Particular attention was addressed to the ANALYSIS OF CHARRED SOLID RESIDUE structural changes by IR

characterization at different temperatures of thermal degradation. Moreover, the hyphenated technique TG-IR

has allowed the definition of the temperature ranges within which the volatile species evolved during the heating

process adopted for the experiment. Based on the results obtained from TGA analysis on neat PET, pyrolysis

•Only Holland and Hay have provided a study of the structural evolution of PET charred solid residues during thermal or thermoxidative degradation.



Identify which of the mechanisms reported in the literature are relevant to PET thermal decomposition, thus providing an integrated mechanism for both volatilisation and charring of PET in the combustion process by a detailed molecular characterization of the volatiles and of the residues produced by thermal degradation. The results of this work can be useful for designing fire retardants for PET.

LITERATURE BACKGROUND

is still not clear whether heterolytic or homolytic scission of aliphatic fragments is the first step of degradation.



They provided an explanation for the







TGA of neat PET Nitrogen

Single stage of thermal degradation. T_{max} = 450°C

Air

Two stages of thermal decomposition. @ 590°C: complete oxidation to volatile products of carbonization residues.

 N_2

CO2 CO2 Gas phase Water → Water Acid Acid

Characterization approach

experiments were performed at 375,400 and 425°C.

THERMAL DEGRADATION \mathbb{N}_2

DIFFERENT TECHNIQUES HAVE PERMITTED TO DETECT THE SPECIES **PRODUCED IN THE THREE PHASES**

The release of non condensable IR was monitored bv gases TG-IR during spectroscopy experiment.

CONCLUSION: Hypothesis of mechanism

Temperature [°C]

----- PET N₂ ----- PET Air

20°C/min

To hypothesize a mechanism of PET residue thermal degradation the greatest difficulty lies in the comparison of literature data that was provided by different thermal degradation approaches. The authors have focalized their attention on the final solid residue structure depicting paths of hetero or

homolytic degradation.

In fact it is known that for the gas phase pyrolysis more common is an Ei mechanism and radical reactions dominate the process. However, for polymers where the pyrolysis takes place in condensed phase, E2 and E1 mechanisms are not excluded. A type of mechanism which involves a cyclic transition state, which may be four-, five- or six-membered could be accompanied by radical reactions at the same

Acid-catalyzed Hydrolysis ROLE

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conjugated aromatic systems. It is reasonable to ponder also on the conjugation of aromatic structures with carbonyl groups of ketones, acids, esters and anhydrides. The condensation of aromatic rings is accompanied by the crosslinking of linear chain segments which does not break the conjugation between the aromatic domains.

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