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Destroy, create, transform and sublimate: laboratory dissociation studies on polycyclic aromatic hydrocarbons and analogues

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Destroy, Create Transform and Sublimate

Laboratory Dissociation Studies on Polycyclic Aromatic
Hydrocarbons and Analogues

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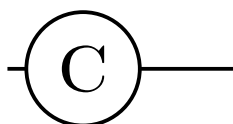
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"Ik heb 'm nooit gezien, maar toch moet tie echt bestaan."



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