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Towards a structural understanding of plant-microbiota interactions using cryo-EM techniques

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**Towards a Structural Understanding
of Plant–Microbiota Interactions
using cryo-EM Techniques**

Janine Liedtke

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Towards a Structural Understanding of Plant–Microbiota Interactions using cryo-EM
Techniques

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Towards a Structural Understanding of Plant–Microbiota Interactions using cryo-EM Techniques

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Utrecht University

... für Papa

*Denn die einen sind im Dunkeln
Und die anderen sind im Licht.
Und man siehet die im Lichte
Die im Dunkeln sieht man nicht.*

Brecht

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