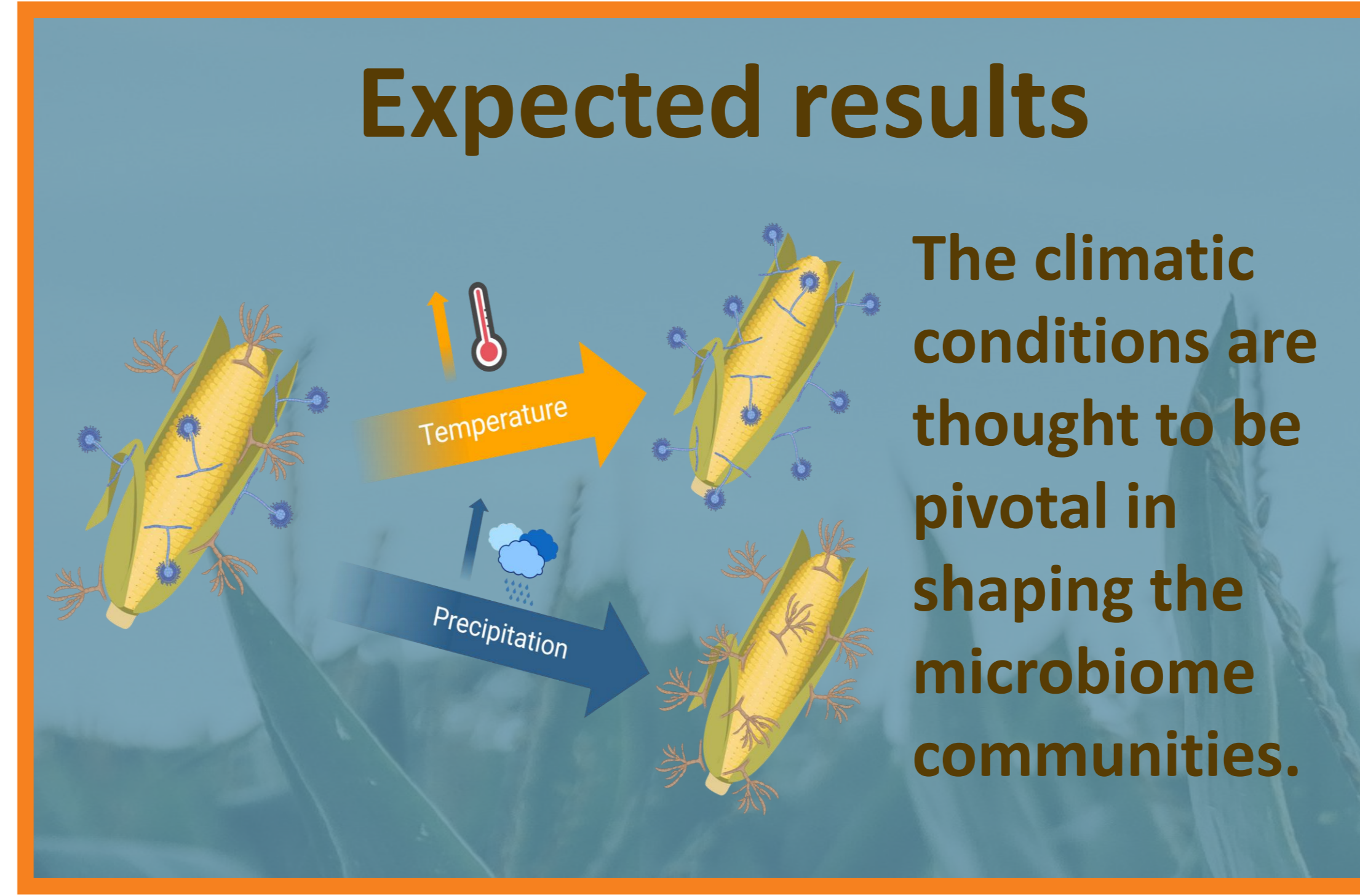
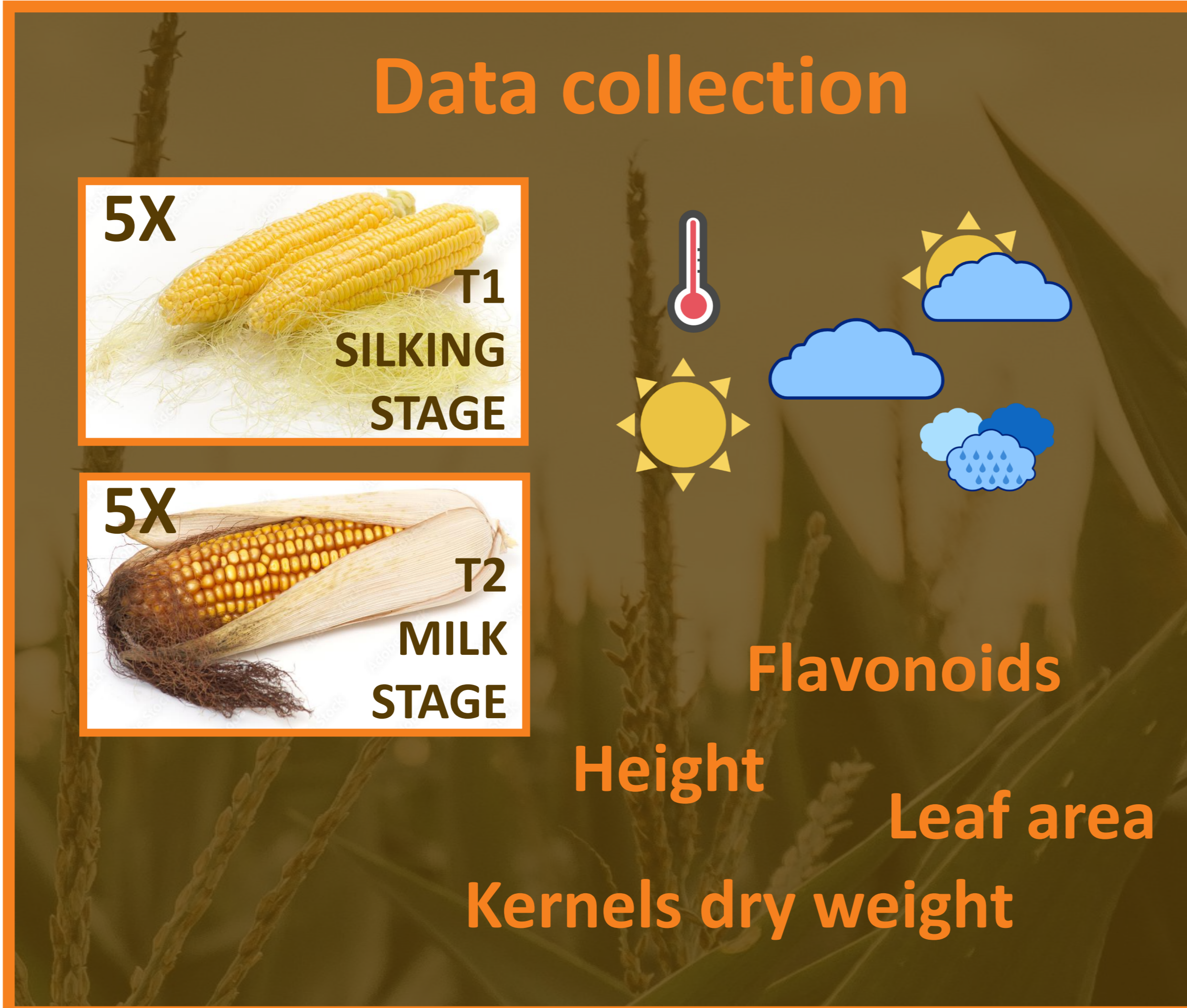
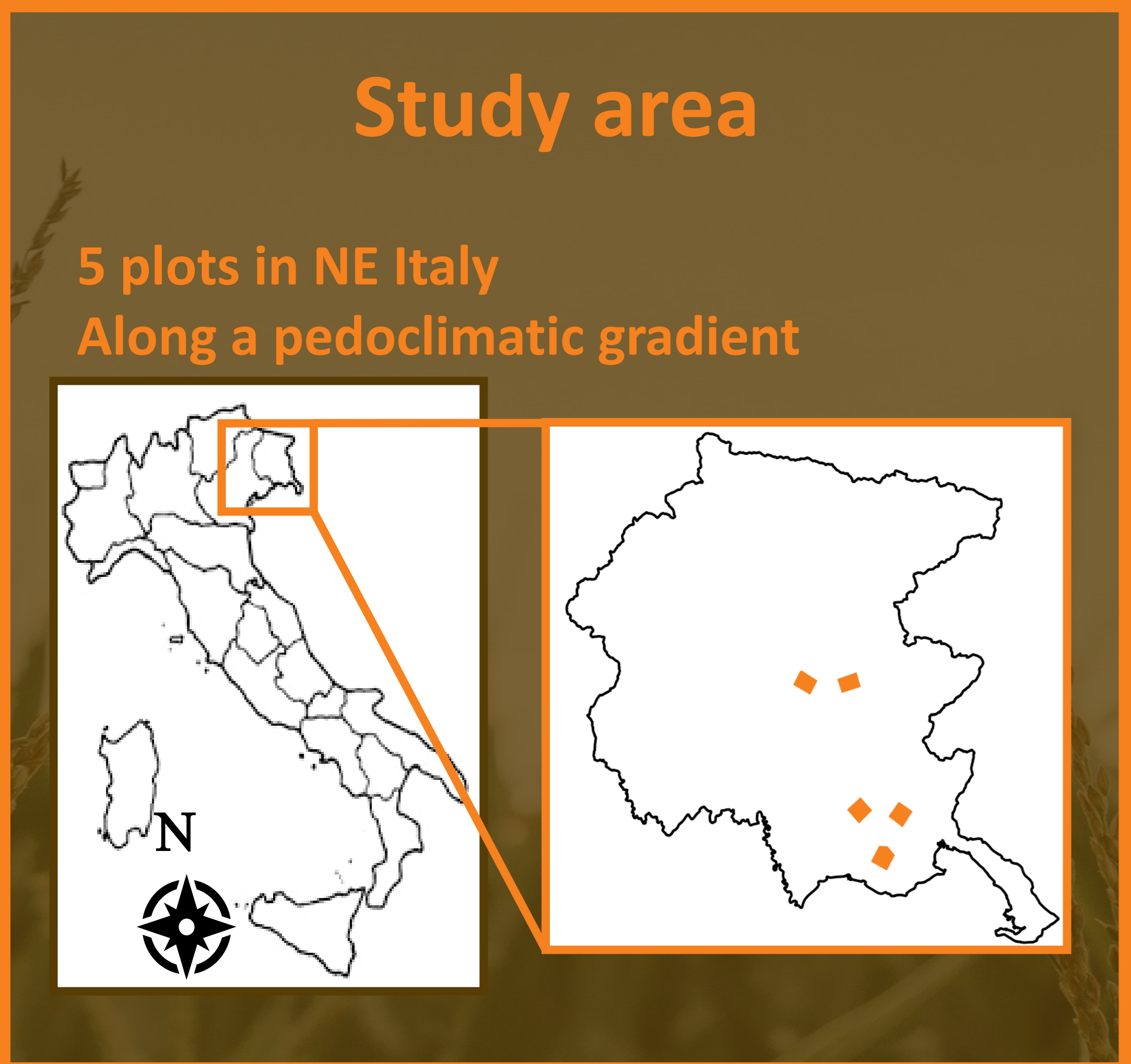


Interactions between plant response to environment and fungal microbiome in developing maize silks in relation to mycotoxin risk

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Conclusions

The communities are diverse in T1 and T2.

The species richness (R) increases from T1 to T2.

Temperature is statistically significant in shaping the composition of communities.

