ADDENDUM

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ABSTRACT

Maize, Zea mays, the second-most-widely-grown crop, yields 20 % of all consumed calories worldwide.¹ Despite its agronomic importance, research progress is limited by costly transformation. We recently described the Trojan horse method as a useful tool to study maize proteins in situ that circumvents timeand space-consuming whole plant transformation. The Trojan horse approach uses the protein-folding and secretory properties of the corn smut fungus Ustilago maydis to secrete maize proteins from fungal cells into the maize apoplast. Here, we discuss the timing and location of U. maydis during infection and the protein secretion site in relation to anther anatomy. This spatiotemporal analysis enables the study of apoplastic anther proteins in various premeiotic anther developmental stages, and could be adapted for larger screens.

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In the 2017–2018 harvest, 1,033 million tons of corn were produced worldwide, 2 making corn the most-grown cereal crop. People are highly dependent on this renewable food and energy source, motivating researchers to investigate new ways to increase yield.

Recent advances in maize genetic-editing using CRISPR-Cas9 technology³ make modern reverse genetics much easier than classical transposon insertion lines. Another milestone in stable maize transformation was achieved more recently with precise expression of maize BABYBOOM and WUSCHEL2 proteins enabling genotype-independent embryo transformation without time-consuming callus regeneration.^{$4,5$ $4,5$} Even so, the processes of vector construction and transformation are laborious, and consume greenhouse space. To overcome these drawbacks, we developed an alternative, the Trojan horse method.^{[6](#page-2-4),[7](#page-2-5)}

The biotrophic corn smut fungus U. maydis infects all aerial plant organs. During infection the hyphae never contact the plant cytoplasm; instead the plant plasma membrane invaginates resulting in a narrow apoplastic space between hyphae and plant cells. All interaction between the plant and fungus must traverse this tiny space, the biotrophic interac-tion zone.^{[8,](#page-2-6)[9](#page-2-7)} Upon infection *U. maydis* secretes diverse effector proteins to suppress the plant´s immune response. Sequences encoding signal peptides for classical secretion of these effectors have been defined.^{[10](#page-2-8)} Unconventional secretion of proteins has been reported and can be hijacked by researchers to secrete non-post-translationally modified proteins in *U. maydis* liquid culture.^{11-[13](#page-2-10)} The underlying mechanism is still poorly understood and has not yet been tested in a biotechnology setting in planta.^{[13](#page-2-10)-[15](#page-2-11)} Compared to its host, U. maydis has a small genome and is readily

transformed.¹⁰ After decades of work with this genetic model organism, an array of cloning vectors with a variety of epitope tags and promoters is available. U. maydis has also become a protein expression tool in biotechnology because of its folding and post-transcriptional modification abilities. $13,15,16$ $13,15,16$ $13,15,16$ Collectively, these features facilitate exploiting U. maydis to express and secrete maize proteins in situ. This approach permits single-cell resolution by tracking the Trojan horse hyphae during plant infection and allows comparison of maize cells receiving secreted protein to non-receiving cells in the same tissue.^{[7](#page-2-5)} Expression of foreign proteins by Trojan horse strains is driven by a strong in planta active promoter.^{[6](#page-2-4),[7,](#page-2-5)[17](#page-2-13)} This allows protein delivery to all infected tissue layers. To permit expression at specific time points or locations, alternative promoters may be used in the future. Even though U. maydis is known for its protein folding and secretion capabilities, secretion of foreign proteins of interest needs to be carefully monitored.^{[17](#page-2-13)} Also, the protein size limit for secretion is not yet known. The U. maydis effector Cmu1 (Chorismate mutase1) at nearly 300 amino acids long plus an mCherry-tag successfully complemented the cmu1 deletion mutant.^{[18](#page-2-14)}

To implement the Trojan horse method and achieve the highest possible spatiotemporal resolution, detailed growth timelines for both the host and the pathogen are required. For maize, detailed analyses of premeiotic anther develop-ment have been performed.^{19-[22](#page-3-0)} Premeiotic anthers double in length approximately once per day. During early stages, at least one new lobe tissue layer is formed each day over the course of four days of growth [\(Figure 1\)](#page-1-0). Correlations of tassel size and anther stages in different zones of the tassel (for

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maize inbred W23) showed that a tassel of 5–7 cm contains all stages of premeiotic anthers.^{[25](#page-3-1)} Consequently, the impact of infection by a Trojan horse strain can be assessed in all anther lobe cell types with just one tassel infection.

U. maydis infections were analyzed by various labs indicating reproducible patterns and timelines in all examined maize tissues.^{[8](#page-2-6)-[10](#page-2-8)[,24](#page-3-2)} This is true even though *U. maydis* deploys divergent effectors for infection of different host tissues and host tissue responses vary in turn.^{[26](#page-3-3)} An infectious hypha with an appressorium forms, and epidermal cell penetration starts within 12 h after contact with the plant. In the following 12 h infected epidermal cells are fully colonized ([Figure 1\)](#page-1-0). Two days post infection (dpi) two subepidermal layers are infected. After 3 dpi the entire organ is colonized.^{[7](#page-2-5)-[10](#page-2-8)[,24](#page-3-2)} In seedling leaves, U. maydis strain SG200-induced cell proliferation occurs 4-5 dpi.^{[27](#page-3-4)} In SG200-infected anthers, no significant formation of additional cells was observed 3 dpi. Subsequently, developmental studies using the Trojan horse approach were performed 3-4 dpi.^{[7](#page-2-5)}

To analyze the impact of the small secreted maize protein ZmMAC1 on Layer2-derived and archesporial cells present in 120–300 μ m anthers, tassels with 50–125 μ m anthers were infected. Three dpi, anthers were now 400–- [7](#page-2-5)00 μ m and were harvested for confocal imaging.⁷ By combining existing observations, a timeline for anther Trojan horse experiments can be estimated ([Figure 1](#page-1-0)). Presumably, infection of 50 μ m anthers should allow protein secretion to epidermal cells, primary parietal cells, secondary parietal cells, and the endothecium before observation of cellular responses is performed 3 dpi. At this time

Figure 1. Timeline for the Trojan horse approach in premeiotic maize anthers.

Schematic illustration of different developmental stages of premeiotic anther lobes. Maize anther lobe development is strongly linked to anther size, as exemplified
here for maize cultivar W[23](#page-3-6). New tissue layers form withi to days post infection (dpi) with *U. maydis* based on previous studies.^{[7](#page-2-5)-[10](#page-2-8)[,24](#page-3-2)}

point, epidermal cells should have received the secreted protein for two days, while primary parietal cells, secondary parietal cells, and the endothecium should have been treated for 1 day. To analyze the response of middle layer cells to a secreted protein of interest, anthers 350–450 µm long have to be infected.

In addition to developmental studies, the Trojan horse approach is a tool for analysis of proteins in plant-pathogen interactions.⁶ Here tumor formation is used as an easy measure of plant susceptibility or resistance to the fungus after receiving a secreted protein of interest. The study of the function of apoplastic maize peptide ZIP1 resulting from proteolytic cleavage of PROZIP1 is one example of this. During compatible maize-U. maydis interactions, release of ZIP1 is blocked. Secretion of ZIP1 by U. maydis reduces tumor formation, and hence increases resistance to *U. maydis.*^{[6](#page-2-4)} Thus, proving that not only proteins of interest but also individual protein domains can be functionally analyzed by this approach.

The Trojan horse approach can be used to analyze protein or protein-domain functions with high spatiotemporal resolution in situ without stable host transformation. Successful implementation relies on detailed knowledge of host developmental timelines and pathogen infection strategies. Easy transformation, understanding of secretion mechanisms and good protein folding capacity on the pathogen side are indispensable. Fortunately, many plant- and animal-pathosystems offer these features making the Trojan horse approach widely generalizable in the future for systems that lack easy host transformation technologies.

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