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1	Original research paper :
2	Title : Interrelated responses of tomato and the leafminer Tuta absoluta to nitrogen
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- 2 limitation, cross responses, interactions, plant defence, polyphenol oxidase, phenolics,
- 3 tomatine, growth

4 Abbreviations

- 5 CHAx: isomer x of caffeoyl hexaric acid
- 6 CP: caffeoylputrescine
- 7 FLA: flavonoids
- 8 FQA: feruloyl quinic acid
- 9 Hx: Harvest x
- 10 HCAD: hydroxycinnamic acid derivatives
- 11 HN: high nitrogen
- 12 KR: kaemperol rutinoside
- 13 LN: low nitrogen
- 14 N: nitrogen
- 15 NFT: nutrient film technique
- 16 pCoQA: p-coumaroyl quinic acid
- 17 PHE: phenolamide
- 18 PPO: polyphenol oxidase
- 19 PVPP: polyvinylpolypyrolidone
- 20 QAR: quercetin apiosyl-rutinoside
- 21 R: rutin
- 22 x-CGA: isomers of chlorogenic acid

1 ABSTRACT

Plant-insect interactions are strongly modified by environmental factors. This study evaluates,
for the first time, the means by which nitrogen fertilisation affects the interaction between
tomato plants (*Solanum lycopersicum* L. cv. Santa clara) and the leafminer *Tuta absoluta*(Meyrick) (Lepidoptera: Gelechiidae).

Greenhouse grown tomato plants were fed hydroponically on a complete nutrient solution
containing either a low nitrogen concentration (LN) limiting plant growth or a high nitrogen
concentration (HN) sustaining maximum growth. Insect-free plants were compared with
plants infested by *T. absoluta*. Seven and 14 days after an artificial oviposition leading to
efficacious hatching and larvae development, we measured vegetative tissue composition in
primary insect resources (total carbon, nitrogen, protein) together with defencive compounds
(phenolics, glycoalkaloids, polyphenol oxidase activity) in HN vs. LN plants.

It was only in the HN treatment that T. absoluta infestation slightly impaired leaf growth and 13 induced polyphenol oxidase (PPO) activity in the foliage. The concentration of phenolic 14 15 compounds and proteins, together with the total N distribution within the plant, were not affected by T. absoluta infestation. LN nutrition impaired the T. absoluta-induced PPO 16 17 activity. It decreased protein and total nitrogen plant organ concentrations and enhanced the 18 accumulation of constitutive phenolics and tomatine. Moreover LN nutrition impaired T. absoluta development by notably decreasing pupal weight and increasing the development 19 time from egg to adult. Nitrogen nutrition may thus be a means of altering the life cycle of T. 20 absoluta. 21

22 These results confirm for tomato, the existence of several cross-responses of plant

23 composition and *T. absoluta* development to nitrogen nutrition.

1 INTRODUCTION

2 Plants respond to herbivorous insect feeding by means of a set of resistance mechanisms. 3 These mechanisms involve localised and systemic synthesis/emission of secondary 4 metabolites (Zangerl et al. 2002), induction of defencive enzymes (Stout et al. 1994) and 5 tolerance mechanisms such as resource and metabolite remobilisation within plant organs 6 (Tiffin 2000). It is now established that plant responses are highly specific to the insect 7 feeding guild, this specificity being driven by the complex interaction of, at least, three major phytohormone signalling pathways *i.e.* jasmonic acid, ethylene and salicylic acid (Erb et al. 8 2012, Pieterse et al. 2009). Indeed, plants submitted to either phloem-feeding or chewing 9 10 insects have been shown to exhibit differences in defencive enzyme induction (Felton et al. 11 1994), phenolic compound accumulation (Olson & Roseland 1991), regulation of primary 12 metabolism (Schmidt et al. 2009) and transcriptomic responses (Kempema et al. 2007). By 13 contrast, plant responses to leaf miners have received less interest (Stout et al., 1994, Cardoso et al. 2014, Zhang et al. 2012). Leaf mining results from the ability of insect larvae to feed 14 and develop within plant tissues, mostly leaves and stems. From an ecological viewpoint, this 15 feeding strategy confers protection against natural enemies and allows larvae to avoid the 16 defence barriers (trichomes, spines...) on the leaf surface (Connor & Taverner 1997). 17 18 Furthermore, it is acknowledged that leaf miners provide a valuable model to study plant responses to insect damage due to the intimate interactions created by larvae developing 19 within plant tissues (Han et al. 2014, Inbar et al. 2001). 20

Plant-herbivorous insect interactions are highly dependent on environmental factors. Of these,
plant nutrition and particularly nitrogen (N) fertilisation, has been widely studied (Bentz *et al.*1995, Cates *et al.* 1987, Chen *et al.* 2010, Fischer & Fiedler 2000, Mattson 1980, Han et al.,
2014). N is an important macronutrient for both plants and herbivores. Plants require a great
deal of N to attain maximum growth and N concentration in the insect body tissues is more

1 concentrated than in their foodstuff (Schoonhoven et al. 2005). N deficiency (or limitation) at 2 the root level reduces tomato plant growth and N tissue concentration (Adamowicz & Le Bot 3 2008) whereas the concentrations of constitutive secondary compounds such as phenolic acids and flavonoids are increased (Fritz et al. 2006, Larbat et al. 2012a, Larbat et al. 2012b, Larbat 4 et al. 2014, Le Bot et al. 2009) and also glycoalkaloids (Royer et al. 2013) in Solanaceae. 5 From the viewpoint of the plant, N fertilisation affects the inducible defence in a complex 6 7 way that depends on the pathway considered. Indeed, N limitation reduces the induction of trypsin inhibitor and the accumulation of nicotine in infected tobacco, whereas it has no effect 8 9 on the induction of volatile terpenes (Lou & Baldwin 2004). From the insect viewpoint, plant 10 N fertilisation influences the development of various herbivores, especially lepidopterans, 11 through either (i) the plant nutritional value linked to tissue N concentration (Cates et al. 1987, Estiarte et al. 1994, Grundel et al. 1998, Han et al. 2014, Hunter & McNeil 1997, Inbar 12 et al. 2001, Schoonhoven et al. 2005), or (ii) the content of constitutive or induced chemical 13 compounds and mechanical plant defence (Gutbrodt et al. 2011, Koricheva 2002). The 14 relative importance of both effects on herbivore performance, however, is difficult to assess 15 and is likely pathosystem-specific. 16

17 The objective of this study was to evaluate the impact of N fertilisation on the ability of tomato to resist to the leafminer Tuta absoluta (Meyrick) (Lepidoptera: Gelechiidae). This 18 19 pest originating, from South America (Guillemaud et al. 2015), is now well established in Europe, Africa and the middle East since its first appearance in Spain in 2006 (Desneux et al. 20 21 2011, Desneux et al. 2010). Tuta absoluta larvae feed exclusively on Solanaceae, tomato plants being the major host. Leaf miners bring about serious leaf injuries, leading notably to 22 23 hydraulic damage and reduction in C acquisition (Aldea et al. 2005, Tang et al. 2006), which eventually result in plant death. In tomato, T. absoluta is considered as a serious pest causing 24 25 large production losses. Because it is an emergent pest in Europe, data in the literature remain

1 insufficient to fully characterise the response patterns of tomato to T. absoluta, although these 2 are necessary to develop pest control strategies, especially connected to management practice. 3 A recent study from our group identified that water and N availabilities can modulate the tomato susceptibility to T. absoluta infestation and the T. absoluta development (Han et al., 4 2014). In this study, the physiological clues underlying these effects were not assessed. The 5 present study was thus designed to assess the impact of N availability on the tomato plant 6 7 response to T. absoluta. Our experimental strategy was to grow tomato plants hydroponically in a greenhouse under two regimes of nitrogen fertilisation, one limiting plant growth (low 8 9 nitrogen: LN) and the other adequate for maximum growth (high nitrogen: HN). We measured tissue composition in primary insect resources (total carbon, nitrogen, protein) 10 11 together with defencive compounds (phenolic compounds, glycoalkaloids, polyphenol oxidase activity) in the vegetative parts of HN vs. LN tomato plants subjected to T. absoluta 12 infestation or maintained insect-free. The consequences of N fertilisation on leaf miner life 13 traits were determined and analysed in relation to the composition of plant tissues. 14

15 MATERIALS AND METHODS

17

Plant growth, Tuta absoluta infestation and harvests 16

This experiment was carried out under glasshouse conditions in Avignon (43°56'58" N, 18 4°48'32" E). Tomato seeds (Solanum lycopersicum L. var. Santa Clara) were sown in an NFT (nutrient film technique) system set up in a growth room as described in Larbat et al. (2012a). 19 Twelve-day-old plantlets were then transferred to the glasshouse and grown from May 30th to 20 21 June 21th 2012 under the following conditions: heating when air temperature $\leq 18^{\circ}$ C, ridge opening when $\geq 25^{\circ}$ C, mist spraying when humidity $\leq 55^{\circ}$. The glasshouse was whitewashed 22 to ease temperature control. Plants were grown hydroponically in six fully randomised blocks, 23 each providing a complete nutrient solution at two regimes of N concentration, respectively 24

representing high N (HN, 1.5 mM NO₃⁻) and low N (LN). In the latter, [NO₃⁻] was modified
once per day (from 10 µM to 30 µM) in order to maintain the daily NO₃⁻ uptake of LN plants
around 1/3rd of the value measured in the HN plants, using the Totomatix system (Adamowicz *et al.* 2012). During the experiment, [NO₃⁻] and pH were corrected hourly in the nutrient
solutions. In both solutions the sum (NO₃⁻ + SO₄²⁻) = 12 eq m⁻³, inferring constant
concentrations of other ionic species in all treatments (Le Bot *et al.* 2009).

7 Insect preparation and infestation

To obtain the T. absoluta eggs the method of Chailleux et al. (2013) was used. Ten couples of 8 9 adult insects were maintained for 24 hours in a double-cup system containing a fresh tomato 10 leaf and honey provided as foodstuff, under the following conditions: air temperature 25°C, 65% relative humidity, 12/12h photoperiod. The adults laid eggs on this leaf. On the 11 12 following day, an artificial oviposition was carried out whereby the eggs were gently transferred with a wet brush, to the terminal leaflet of the third leaf (counting from the base) 13 of nineteen-day-old plantlets growing in the greenhouse. A load of two eggs was placed on 14 each leaflet to maximise the chances of T. absoluta development. The infested leaves were 15 bagged *i.e.* enclosed using a nylon mesh (0.2 mm, 30 × 24 cm). The third leaf of the non-16 infested plants (controls) was similarly bagged to take into account any possible effect of the 17 mesh on leaf growth and metabolism. 18

19 Harvests and sample preparation

Three harvests were taken. The first (H1) was made on June 6th 2012 prior to *T. absoluta*oviposition, to characterise plant morphology and biochemical composition before infestation.

22 The second (H2) and the third (H3) were taken 7 and 14 days respectively after *T. absoluta*

- 23 oviposition. At each harvest, leaves, stems and roots were separated. Leaves were sub-
- sampled within the bulk foliage to separate the infested leaf (*i.e.* 3rd leaf from the base) and its

opposite leaf (*i.e.* 4th leaf), which served to test for systemic plant responses. Roots were
rinsed in deionised water and spin-dried (2 min at 2800 g). Plant parts were weighed, frozen
in liquid N₂ and stored at -80 °C until freeze-drying. Dried samples were weighed, ground to a
fine powder and stored under dry air in a desiccator at room temperature. In addition, at
harvests H2 and H3, the leaflets containing the larvae were weighed and a digital picture was
taken in order to calculate, by image processing, the number and the surface of mines dug by
the larvae.

8 Tuta absoluta survival and development

9 For each infested leaf harvested at H3, larvae survival was recorded. Each infested leaf was
10 thus kept in a double-cup system containing HN or LN nutrient solution as in the initial
11 treatment, until the larvae of *T. absoluta* pupated and the adult emerged. The pupae were
12 counted and weighed individually. The development time from egg to pupa or to adult was
13 recorded for all individuals.

14 Standards and chemicals

Chlorogenic acid (5-CGA), rutin, kaempferol rutinoside, ferulic, p-coumaric and caffeic acids
were purchased from Sigma (Steinheim, Germany). Solanine and tomatine were obtained
from Extrasynthese (Genay, France). Caffeoylputrescine was kindly provided by Dr. WerckReichhart (IBMP, Strasbourg, France).

19 Analyses of plant tissues

- 20 Total C and N concentrations were determined using an elemental auto-analyser (Flash EA
- 21 1112 series, Thermo Fisher Scientific, Courtaboeuf, France), on 3 mg of dry powder,
- 22 according to the Dumas method.

1 Phenolics and tomatine were extracted from 20 mg dry powder of the infested and opposite 2 leaves, stems and roots as described in Royer et al. (2013). For tomatine quantification, the extract was diluted fiftyfold in 70% MeOH containing 2 µM solanine as internal standard. 3 The compounds, from undiluted and diluted extracts respectively, were separated on a U-4 HPLC system (Prominence, Shimadzu, Japan) consisting in a binary solvent delivery pump 5 connected to a diode array detector. Two microliters of extract were separated on a C18 6 7 Zorbax Eclipse Plus (150 mm \times 2.1 mm, 1.8 µm) column (Agilent, USA) by using a gradient elution from 1 to 50% MeOH 0.1% formic acid (FA) in 7.1 min, then 99% MeOH 0.1% FA in 8 0.8 min with a flow rate of 430 µl min⁻¹. The column was rinsed during 2 min with 99% 9 10 MeOH 0.1% FA and re-equilibrated to the initial conditions for 2 min prior to the next run. 11 Phenolic quantification was based on the area under peak determined at 320 nm and expressed relative to calibration curves with ferulic acid (for FQA), chlorogenic acid (for 5-12 CGA, 1-CGA, 4-CGA), coumaric acid (for pCoQA), caffeic acid (for CHA1-6), 13 caffeoylputrescine (for CP). Regarding flavonoids, quantification was determined at 350 nm 14 and expressed relative to calibration curves with rutin (for R and QAR) and kaempferol 15 rutinoside (for KR). Tomatine was detected by mass analysis carried out in ESI positive ion 16 mode (ESI+) by following the major ion at m/z 529. The internal standard, solanine, was 17 18 followed at m/z 868. Tomatine was quantified relative to a tomatine calibration curve (0.2-10 μ M). Mass spectrometric conditions were previously described in Royer *et al.* (2013). 19

20 Polyphenol oxidase and total protein assays

Polyphenol oxidase (PPO) and total protein assays were made on H3 leaf extracts. All the
leaves of each individual plant were pooled, frozen in liquid nitrogen and crushed in a mortar.
Then, 50 mg of fresh weight (FW) were macerated, in a 2 ml tube, with 500 µl cold extraction
buffer (sodium phosphate 0.1 M pH 7 with 3% polyvinylpolypyrolidone (PVPP) and 1%
Triton X-100). This extract was mixed with a vortex for 1 min, then centrifuged at 10000 g

for 10 min. The supernatant was used for both the polyphenol oxidase assay and the total
 protein assay.

PPO assay was carried out by mixing 20 µl of leaf extract with 200 µl of pre-warmed reaction
buffer (sodium phosphate 0.1 M pH 7 containing 3 mM caffeic acid). PPO activity was
determined by monitoring the appearance of quinone products from caffeic acid at 470 nm at
25°C. PPO global activity was then expressed as the rate of absorbance change per mg of FW
and PPO specific activity as the rate of absorbance per mg of total protein.
Protein quantification followed the Bradford procedure using bovine serum albumin as

9 standard. Concentration was expressed as mg protein per g of leaf FW.

10 Data processing

Whole leaf and damaged areas, perimeters and mine numbers were determined by image 11 processing with Adobe Photoshop CS4 extended (Adobe systems Software, Ireland Ltd.). 12 Computations were performed using the R software (R project for statistical computing, 13 available at http://www.r-project.org) and statistical significance was set at p < 0.05. Analyses 14 of variance were performed using the aov procedure, with nutrition, and infestation as fixed 15 factors and blocks as random. Box-plots, Normal Q-Q plots and correlation between variance 16 17 and mean, assessed the data distribution and homoscedasticity. Square root transformation was necessary for homoscedasticity of third leaf weight data. Tuta absoluta survival on 18 19 tomato plants subjected to the nitrogen treatments was analysed using a log-linear model. Proportions of individuals alive at each developmental stage were compared by pairwise 20 21 Fisher's exact tests (with the Dunn–Sidak adjustment method). The effects of the nitrogen 22 treatment on development time from egg to pupa stage and from egg to adult stage were tested, as well as on pupal weight using a generalised linear model with a log-link function. 23

24 RESULTS

1 Impact of N and T. absoluta on plant growth

In the LN treatment, the insect-free plants showed no visual symptom of N deficiency 2 throughout the entire experiment. However, they were markedly reduced in shoot FW as 3 compared with the HN plants (Table 1). At the first harvest, shoot FW was significantly 4 decreased (p = 0.04) by 21%, the difference between the two N regimes in favour of the HN 5 treatment increasing at harvests H2 (42%, $p = 2 \ 10^{-8}$) and H3 (66%, $p = 8 \ 10^{-18}$). For both N 6 treatments, infestation by T. absoluta did not significantly alter the shoot FW at H2 and H3. 7 8 However, the infestation specifically decreased the FW of the infested leaves at H2 and H3 in 9 the HN treatment (p < 0.003).

10 Tissue N concentration and C/N ratio

11 At H3, tissue N concentration and C/N ratios were significantly altered by N nutrition (Fig 1.

12 A-B). Indeed, N limitation significantly reduced N concentrations (p < 0.001) and thus,

13 increased C/N ratios (p < 0.001) in all tissues. The effects were organ dependent (p < 0.001),

14 being more pronounced on stems. Insect feeding did not affect the N concentration or C/N

15 ratio of any tissue (p = 0.13 and 0.20 respectively).

16 Soluble phenolics

17 The phenolic composition was highly dependent on plant organs (Sup. data 1). Fourteen

18 phenolic compounds *i.e.* six isomers of caffeoyl hexaric acid (CHA1-6), three isomers of

19 chlorogenic acid (5-CGA, 3-CGA and 1-CGA), feruloyl quinic acid (FQA), coumaroyl quinic

20 acid (pCoQA), rutin (R), quercetin apiosyl-rutinoside (QAR), kaempferol rutinoside (KR) and

- 21 caffoylputrescine (CP) were investigated (12 in infested and opposite leaves, 10 in stems and
- 22 3 in roots). Analyses were carried out for the three harvest periods (H1 to H3, all data are

detailed in Sup. data 2), but for the benefit of the reader, the 14 phenolic compounds were

24 pooled into 3 groups *i.e.* hydroxycinnamic acid derivatives (HCAD, comprising CHA1-6, 1-

CGA, 3-CGA, 5-CGA, pCoQA and FQA), flavonoids (FLA, comprising R, QAR and KR)
 and phenolamides (PHE, comprising CP).

HCAD, FLA and PHE were distributed differently within the plant. HCAD and PHE were 3 detected in all plant organs (Sup. data 1-2) with the highest HCAD concentration in leaves 4 5 (infested and opposite, p < 0.001) and the highest PHE concentration in stems (p < 0.001). FLA were detected only in shoots (Sup. data 1). The concentration of all phenolic groups 6 7 varied between harvests in all organs, but not in the same way (Sup. data 2). HCAD 8 concentrations were highest at H1 (p < 0.001) but did not differ significantly between H2 and 9 H3. PHE concentrations were highest at H1 and H2 then dropped markedly at H3 to reach undetectable levels in opposite leaves. By contrast, FLA concentrations increased 10 significantly at H3 (p < 0.001). Nitrogen limitation did not affect HCAD and FLA 11 12 concentration at H1. However, LN clearly increased HCAD and FLA concentrations in all 13 organs at H2 and H3 (p < 0.001) (Fig. 2). LN reduced PHE concentrations in stems and roots at H3 only. Insect feeding did not affect HCAD, FLA and PHE concentrations at any harvest 14 15 nor in any organ.

16 *Tomatine*

The concentration of tomatine, the major tomato glycoalkaloid involved in plant defence, was 17 18 determined in all plant organs at H2 and H3 (Fig. 3). Tomatine responded significantly to N nutrition only in the stems, where its concentration increased under LN (H2 p < 0.001, H3 p <19 0.05). Insect feeding significantly affected tomatine concentration in stems and roots at H2 20 with opposite effects. Indeed, tomatine concentration decreased in the roots of infested plants 21 22 (p < 0.01) while it increased in stems (p < 0.01). These effects disappeared at H3. Insect feeding also brought about a small, non-significant (p = 0.06831) decrease in tomatine 23 concentration of infested leaves at H3. 24

1 Inducible responses of proteinaceous defence and total protein at H3

The effects of N limitation and insect infestation on polyphenol oxidase (PPO) activity and 2 total protein content were assessed in infested and control leaves at H3 (Fig. 4 A, B, C). The 3 global PPO activity expressed on a leaf FW basis (Fig. 4-A) increased under T. absoluta 4 5 infestation. The effect was significant under HN (p < 0.01), but not under LN (p = 0.32). The global PPO activity did not respond to N nutrition. By contrast, total protein concentration 6 (Fig. 4-B) responded significantly to N nutrition (infested and control leaves, p < 0.001) but 7 8 not to T. absoluta feeding (p = 0.29). The specific PPO activity expressed on a total protein basis (Fig. 4-C) increased significantly in response to insect feeding in the HN treatment only 9 (p < 0.05). In addition, the specific PPO activity was higher under LN (p < 0.001) than under 10 HN. 11

12 Tuta absoluta traits

The survival of *T. absoluta* (Fig. 5) decreased significantly in response to low N nutrition
(χ² = 4.8, df = 1, p = 0.028) and varied with the insect's developmental stage (χ² = 11.9,
df = 3, p = 0.008). The interaction between both factors was not significant (χ² = 2.5, df = 3,
p = 0.47). Under LN, the survival rate was mainly reduced during the larval stage while under
HN, the survival rate did not differ significantly between egg and larva or egg and pupa

stages. It is only between egg and adult stages that it was possible to observe a significant
decrease in the survival rate of *T. absoluta*.

20 Tuta absoluta development

21 Overall, the LN treatment significantly depressed pupal weight at H3 (Fig. 6-A; $\chi^2 = 6.4$,

df = 1, p = 0.011). Low N nutrition significantly increased the duration of insect development

from egg to pupa (χ² = 9.9, df = 1, p = 0.002, Fig. 6-B), as well as from egg to adult (χ² = 4.6,
df = 1, p = 0.032, Fig. 6-C).

3 DISCUSSION

4 Tuta absoluta infestation slightly impaired plant growth and modified tissue composition in 5 a N-dependent manner

6 Tuta absoluta infestation generated moderate plant responses. It decreased FW accumulation 7 in the infested leaves, it enhanced PPO activity in HN leaves and brought about discrete and transient modification of stem and root tomatine concentrations. Besides, as expected at this 8 9 short-term time scale, T. absoluta infestation did not alter the C/N ratio, or the concentrations of N, proteins and phenolics in the vegetative organs. The small egg load during oviposition 10 11 (only two per plant) may not be the main reason for this moderate plant response. Indeed, in a recent study, Mouttet et al. (2013) reported that the pre-infestation of tomato with three T. 12 absoluta larvae was enough to expand plant susceptibility to oïdium infection at a systemic 13 14 level, implying that plant physiology can respond to a small load of *T. absoluta* individuals. 15 Moreover, our recent unpublished data also confirm induction of response at a very low larval density. The design of the experiment and harvest procedure may also have hindered an 16 17 existing stronger local response at the leaflet scale, however the study form Mouttet et al. (2013) demonstrated the existence of a systemic tomato response to T. absoluta. 18

The low defensive response of tomato to *T. absoluta* could thus be explained by the insect
feeding mode. The plant response to *T. absoluta* has not been described previously neither on
tomato nor on other Solanaceous plants. But leaf mining effects on tomato have been
previously assessed in several studies on the serpentine leaf miner *Liriomyza sp* (Stout et al.,
1994, Inbar et al., 1999, Kawazu et al., 2012). Indeed, leaf mining by *Liriomyza trifolii* led,
also, to a smaller induction of defensive proteins in tomato plants (peroxidase, lysozyme) than

1 did other insects with other feeding guilds including leaf-chewing and sap-sucking (Stout et 2 al., 1994, Inbar et al., 1999). This stealthy effect may result from the signalling pathways 3 activated in response to the leaf miner infestation. A recent study highlighted that the jasmonic acid (JA) pathway, which activates the expression of a large range of defensive 4 proteins in tomato, was induced only moderately and transiently before the leaf miners enter 5 6 the tissue, but not once the larvaes were inside. On the contrary, once inside, the larvaes activated the salicylic acid (SA) pathway which acts antagonistically to the JA pathway 7 (Kawazu et al., 2012). The authors postulated that this SA activation pathway may be a 8 strategy of the leaf miner to decrease the JA-induced tomato defence. Since the low response 9 10 of tomato to L. trifolii and T. absoluta compared, a more in depth study should be conducted 11 to identify possible similarities in the plant response to these two leaf miners.

12 Tomato plants responded to *T. absoluta* infestation by activating some resistance mechanisms. 13 We observed in particular an increase in PPO activity in the leaves of the high N treatment. Such an increase in PPO activity is a well-known response to chewing insects, and pathogens 14 15 (Mayer 2006, Stout *et al.* 1998) but, to our knowledge, this paper is the first to report such a 16 response for the tomato-T. absoluta pathosystem. The enzyme PPO catalyses the oxidation of phenolic compounds into quinones, which can bind to amino acids. This accumulation of by-17 products alters the plant nutritional quality and may also be toxic to the larvae (Constabel & 18 19 Barbehenn 2008). Additionally, the soluble phenolic compound concentrations in leaves and other vegetative organs were not affected by T. absoluta infestation. A similar pattern 20 21 (induction of PPO with no impact on soluble phenolics) was previously shown in tomato infested by the chewing insect Helicoverpa zea (Stout et al. 1998), indicating common tomato 22 response traits to different insect feeding guilds. 23

Although tomatine and more generally glycoalkaloids were previously described as toxic

compounds for many insect larvae (for review, see Friedman 2002), the effect of herbivory on

these compounds is scarcely documented and seems to depend on the insect feeding guild
(Fragoyannis *et al.* 2001, Hlywka *et al.* 1994). Our data show that *T. absoluta* infestation had
no significant effect on the foliar tomatine concentration but it induced limited and opposite
responses on stem (increase) and root (decrease) concentration at the second harvest, 7 days
after *T. absoluta* oviposition. This observation suggests the transport of tomatine from roots to
stems. However, a specific experiment is needed to confirm this hypothesis.

Leaf FW accumulation and PPO activity responded significantly to T. absoluta infestation 7 8 only under the HN treatment. Our results for PPO activity differ from other studies dealing 9 with leaf response to damage, which show that inductions of PPO and proteinase inhibitor, another protein-based plant defence, were not affected by N availability in tomato (Stout et al. 10 1998, Tan et al. 2012). Our data, however, are in agreement with those of Lou & Baldwin 11 (2004), who showed that low nitrogen fertilisation reduced the magnitude of damage-induced 12 13 signalling pathways together with the accumulation of nicotine and trypsin inhibitor in Nicotiana attenuata infested with the chewing insect Manduca sexta. 14

15

16 Low N nutrition altered T. absoluta development, plant growth and tissue composition

17 LN nutrition significantly impaired the development of *T. absoluta*, by decreasing the survival rate from egg to larvae, pupa and adult and by increasing the development time from egg to 18 19 adult. LN also reduced the pupa weight. These results accord fully with the observations of Han et al. (2014) for the same pathosystem, and raise the question of how LN nutrition is able 20 21 to impair *T. absoluta* development. Our analyses of plant tissue composition (and particularly 22 the leaves) clearly indicate two possible explanations. Firstly, LN nutrition lowered the plant nutritional value for T. absoluta by decreasing total N and protein concentrations and 23 increasing the C/N ratio. Since N concentration is higher in the herbivore than in plant tissues, 24

1 and because N-based compounds (notably proteins) are essential for larval growth, LN 2 conditions necessarily lower the grazing efficiency of the insect for biomass production, thus 3 impairing its development. The second explanation is that LN increased the level of tomato plant constitutive defence, by increasing the concentration of glycoalkaloids and phenolic 4 compounds but not phenolamides. All these compounds are known to be toxic or repellant to 5 a large array of organisms including insects. Thus, we hypothesise that reduction in plant 6 nutritional quality and increase in constitutive defence both contribute to the observed effect 7 8 of LN on T. absoluta development. Our finding that PPO activity is not induced under LN 9 reinforces this view, and might also indicate low plant responsiveness to T. absoluta under N 10 limitation. To confirm these assumptions, however, it will be necessary to determine the 11 impact of LN on other tomato inducible responses. These include the induction of the methyljasmonate pathway, the activity of other defencive enzymes and the emission of volatile 12 organic compounds, which have recently been shown to increase in T. absoluta infested 13 tomato plants (Strapasson et al. 2014). 14

15 From a practical viewpoint, lower N fertilisation input induced a lower survival and a sub-16 optimal development status in T. absoluta, which may offer a possible means of pest 17 management strategy via manipulation of fertilisation regimes in managed cropping systems (*i.e.* glasshouse tomato production). However, the efficiency of such a strategy should be 18 19 tested at population scales, by integrating the impact of *T. absoluta* infestation and the impact of N itself on tomato yield, using different levels of N supply, since N supply is a detrimental 20 elemental governing tomato yield (Warner et al., 2004). Furthermore, the induction of higher 21 22 chemical defence by T. absoluta may influence the fitness of other pest insects (i.e. aphids and 23 whiteflies), or the infection of plant pathogens, often coexisting in tomato crops (Mouttet et 24 al. 2013). This might be even more complex when organisms from the higher trophic level 25 are involved (i.e. predators) (Bompard et al. 2013). Overall, the net effect of N fertilisation on

tomato plant health (or yield) depends on the interactions of various factors: the occurrence of
insect pests (single or multiple species), plant pathogens and natural enemies introduced into
the system.

4 The present study provided some clues concerning cross-responses of tomato plants and the leaf miner T. absoluta to N nutrition. T. absoluta infestation led to a slight plant response, 5 restricted to induction of PPO activity and reduction of fresh weight accumulation in HN 6 plants. LN nutrition impeded PPO induction in infested plants but also impaired the 7 8 development of T. absoluta. This effect of LN nutrition on T. absoluta development may be explained by the tissue composition of the LN-fed plants, which were depleted in primary 9 10 resources (total N and protein) and enriched in constitutive soluble defence molecules (phenolic compounds and tomatine). Further investigation is necessary to assess the relative 11 contribution of the primary resource depletion and the constitutive defence accumulation on 12 13 *T. absoluta* development.

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11

1 FIGURE LEGENDS

2 Figure 1: Effects of N supply and T. absoluta infestation on C/N ratio (A) and N

3 concentrations (B) in tomato vegetative organs at the third harvest (H3). Inf L: Infested leaf,

4 Opp L: Leaf opposite to the infestation site, S: Stem, R: Root. Error bars are standard error of
5 means (± SEM, n=6).

6 Figure 2: Effects of N supply and *T. absoluta* infestation on HCAD (A), FLA (B) and PHE

7 (C) concentrations in tomato vegetative organs at the third harvest (H3). Inf L: Infested leaf,

8 Opp L: Leaf opposite of the infestation site, S: Stem, R: Root. Error bars are standard error of
9 means (± SEM, n=6)

10 Figure 3: Effects of N supply and *T. absoluta* infestation on tomatine concentrations at the

second (H2) and third (H3) harvests in infested leaf (A), leaf opposite to the infestation site

12 (B), stem (C) and root (D). Error bars are standard error of means (\pm SEM, n=6).

Figure 4: Effects of N supply and *T. absoluta* infestation on global PPO activity (A), total
protein content (B) and specific PPO activity (C) in leaves at the third harvest (H3). Error bars
are standard error of means (± SEM, n=6).

Figure 5: Survival rate of *T. absoluta* individual eggs reaching larva, pupa or adult stage
feeding on HN or LN tomato plants (HN: high nitrogen; LN: low nitrogen; n=24) For each
nitrogen treatment, bars followed by the same letter are not significantly different (pairwise
Fisher's exact tests with Dunn–Sidak adjustment method).

Figure 6: (A) Mean pupal weight (mg, \pm SEM, n = 9-16); Mean development time (B) from

egg to pupa (days, \pm SEM, n = 9-16) and (C) from egg to adult (days, \pm SEM, n = 7-15) of *T*.

22 absoluta individuals feeding on HN or LN tomato plants (HN: high nitrogen; LN: low

23 nitrogen). * p < 0.05, ** p < 0.01 GLM analysis).

1	Supplemental data 1: U-HPLC chromatograms of soluble phenolics from vegetative organs of
2	tomato (cv. Santa Clara) grown under HN (high nitrogen) nutrition. The profiles were
3	recorded at 300 nm. A: leaf; B: stem; C: root. IS: Internal Standard.
4	Supplemental data 2: Concentration of phenolic compounds and tomatine in different tomato
5	organs in Control (= insect-free) or T. absoluta infested (= Tuta) plants harvested immediately
6	prior to infestation (H1), 7 days (H2) or 14 days (H3) after the infestation. Plants were grown
7	hydroponically in the greenhouse and supplied with a complete nutrient solution with either
8	high nitrogen (HN) or low nitrogen (LN) concentration. Inf L: Infested leaves, Opp L:
9	opposite leaves. All concentrations are given in $\mu g g^{-1} DW$ except for tomatine (in mg g ⁻¹
10	DW). Molecules abbreviations as follows: CHA1-6 = six isomers of caffeoyl hexaric acid; 5-
11	CGA, 4-CGA and 1-CGA = three isomers of chlorogenic acid; FQA = feruloyl quinic acid;
12	pCoQA = coumaroyl quinic acid; R = rutin; QAR = quercetin apiosyl-rutinoside;
13	KR = kaemperol rutinoside; CP = caffoylputrescine; HCAD = hydroxycinnamic acid
14	derivatives (CHA1-6 + 1,4,5 CGA + FQA + pCoQA); FLA = flavonoids (R + KR + QAR).
15	Values are means of 6 replicates and are given \pm SE. nd: not defined.



Figure 1: Effects of N availability and *T. absoluta* infestation on CN ratio (A) and N concentrations (B) in tomato vegetative organs at the third harvest (H3). Inf L: Infested leaf, Opp L: Leaf opposite to the infestation site, S: Stem, R: Root. Error bars are standard error of means (± SEM, n=6).



Figure 2: Effects of N availability and *T. absoluta* infestation on HCAD (A), FLA (B) and PHE (C) concentrations in tomato vegetative organs at the third harvest (H3). Inf L: Infected leaf, Opp L: Leaf opposite to the infestation site, S: Stem, R: Root. Error bars are standard error of means (± SEM, n=6).



Figure 3: Effects of N availability and *T. absoluta* infestation on tomatine concentrations at the second (H2) and third (H3) harvests in infested leaf (A), leaf opposite to the infestation site (B), stem (C) and root (D). Error bars are standard error of means (± SEM, n=6).



Figure 4: Effects of N availability and *T. absoluta* infestation on global PPO activity (A), total protein content (B) and specific PPO activity (C) in leaves at the third harvest (H3). Error bars are standard error of means (± SEM, n=6).



Figure 5: Survival rate of *T. absoluta* individual eggs reaching larva, pupa or adult stage feeding on the tomato plants treated with HN or LN input (HN: high nitrogen; LN: low nitrogen; n=24) For each nitrogen treatment, bars followed by the same letter are not significantly different (pairwise Fisher's exact tests with Dunn–Sidak adjustment method).



Figure 6: (A) Pupal weight (mean \pm SEM, n = 9-16); Development time from (B) egg to pupa (mean \pm SEM, n = 9-16) and(**C**) from egg to adult (mean \pm SEM, n = 7-15) of *T. absoluta* feeding on high nitrogen (HN) vs. low nitrogen (LN) tomato plants. (* *p*< 0.05, ** *p* < 0.01 GLM analysis).

		Insect	-free	Infest	ed		
	Harvests	HN	LN	HN	LN		
Shoot	1	4.1 (0.4)	3.2 (0.3)	-	-		
	2	38.4 (2.2)	22.3 (1.6)	34.3 (1.9)	23.6 (2.1)		
	3	227.0 (13.1)	77.3 (3.9)	183.6 (12.3)	81.0 (5.5)		
3rd Leaf	2	5.3 (0.4)	3.4 (0.4)	4.6 (0.4)	3.1 (0.2)		
	3	9.8 (0.6)	4.4 (0.3)	6.6 (0.7)	4.4 (0.3)		

Table 1 : Impact of N availability and *T. absoluta* feeding on the plant fresh weight (g/plant). n=12, standard errors are between brackets



Supplemental data 1: U-HPLC chromatograms of soluble phenolics from vegetative organs of tomato (cv. Santa Clara) grown under HN (high nitrogen) nutrition. The profiles were recorded at 300 nm. A: leaf; B: stem; C: root. IS: Internal Standard.

Supplemental data 2: Concentration of phenolic compounds and tomatine in different tomato organs in Control (= insect-free) or T. absoluta-infested (= Tuta) plants harvested immediately prior to infestation (H1), 7 days (H2) or 14 days (H3) after the infestation. Plants were grown hydroponically in the greenhouse and supplied with a complete nutrient solution with either high nitrogen (LN) or low nitrogen (LN) concentration. Inf L: Infested leaves, Opp L: opposite leaves. All concentrations are given in µg g-1 DW except for tomatine (in mg g-1 DW). Molecules abbreviations as follows: CHA1-6 = six isomers of caffeoyl hexaric acid; 5-CGA, 4-CGA and 1-CGA = three isomers of chlorogenic acid; FQA = feruloyl quinic acid; pCoQA = coumaroyl quinic acid; R = rutin; QAR = quercetin apiosyl-rutinoside; KR = kaemperol rutinoside; CP = caffoylputrescine; HCAD = hydroxycinnamic acid derivatives (CHA1-6 + 1,4,5 CGA + FQA + pCoQA); FLA = flavonoids (R + KR + QAR). Values are means of 6 replicates and are given ± SE. nd: not defined.

Inf L																		
Harvests	Nutr	Treat	CHA1	CHA2	CHA3	CHA4	CHA5	CHA6	5CGA	1CGA	AR	R	KR	FQA	СР	Tomatin (mg/g DW ⁻ ¹)	HCAD	FLA
H1	HN	Control	514 ± 91	37 ± 7	739 ± 136	5070 ± 588	137 ± 25	420 ± 58	3000 ± 228	133 ± 34	109 ± 7	636 ± 20	241 ± 10	36 ± 2	80 ± 13	nd	10088 ± 1114	986 ± 25
	LN	Control	788 ± 65	54 ± 4	1206 ± 110	6422 ± 667	108 ± 25	575 ± 69	2949 ± 288	135 ± 29	112 ± 5	694 ± 70	250 ± 9	34 ± 2	67 ± 10	nd	12273 ± 1194	1057 ± 69
	HN	Control	293 ± 20	18 ± 2	541 ± 44	2875 ± 175	29 ± 2	340 ± 17	1601 ± 109	70 ± 4	59 ± 16	651 ± 42	99 ± 5	28 ± 2	107 ± 14	7.5 ± 1.0	5796 ± 324	810 ± 52
⊔າ	HN	Tuta	281 ± 32	20 ± 2	518 ± 62	2814 ± 253	27 ± 2	340 ± 26	1589 ± 131	75 ± 7	63 ± 6	695 ± 55	104 ± 12	30 ± 2	113 ± 11	7.2 ± 0.6	5691 ± 490	862 ± 72
п∠	LN	Control	478 ± 19	27 ± 4	951 ± 51	4171 ± 153	46 ± 2	574 ± 39	1848 ± 197	81 ± 9	88 ± 5	1024 ± 73	165 ± 10	32 ± 1	118 ±8	7.8 ± 1.2	8208 ± 242	1277 ± 85
	LN	Tuta	422 ± 31	27 ± 2	836 ± 73	3714 ± 234	44 ± 5	506 ± 22	1638 ± 144	73 ± 7	79 ± 9	914 ± 80	152 ± 12	29 ± 2	113 ± 16	9.2 ± 0.8	7289 ± 404	1145 ± 99
	HN	Control	385 ± 44	16 ± 4	805 ± 95	2574 ± 300	42 ± 9	535 ± 36	967 ± 158	54 ± 16	326 ± 26	1606 ± 150	185 ± 13	43 ± 4	45 ± 11	9.4 ± 0.7	5421 ± 462	2117 ± 167
цэ	HN	Tuta	405 ± 31	18 ± 3	860 ± 70	2563 ± 155	41 ± 6	634 ± 64	1004 ± 142	52 ± 13	431 ± 129	1617 ± 227	179 ± 21	40 ± 7	36 ± 7	6.5 ± 1.0	5619 ± 309	2226 ± 245
ПЗ	LN	Control	660 ± 89	25 ± 3	1534 ± 225	3947 ± 392	61 ± 9	981 ± 127	1710 ± 228	48 ± 2	409 ± 49	2195 ± 7287	263 ± 37	67 ± 10	28 ± 6	7.7 ± 0.4	9723 ± 709	3117 ± 299
	LN	Tuta	539 ± 38	22 ± 3	1280 ± 85	3618 ± 248	52 ± 6	855 ± 50	1868 ± 115	65 ± 16	487 ± 22	2607 ±111	304 ± 19	86 ± 9	26 ± 5	6.3 ± 1.0	8388 ± 420	3398 ± 149
Opp L	-	-		-						_				-				
Harvests	Nutr	Treat	CHA1	CHA2	СНАЗ	CHA4	CHA5	CHA6	5CGA	1CGA	AR	R	KR	FQA	СР	Tomatin (mg/g DW ⁻	HCAD	FLA
	HN	Control	329 ± 52	22 ± 6	637 ± 146	2889 ± 567	38 ± 9	360 ± 43	1141 ± 253	56 ± 13	50 ± 8	510 ± 98	83 ± 16	24 ± 4	8 ± 3	5.7 ± 0.8	5497 ± 1100	644 ± 120
	HN	Tuta	363 ± 40	24 ± 5	708 ± 80	3126 ± 312	43 ± 5	358 ± 22	1197 ± 185	63 ± 8	59 ± 9	681 ± 93	99 ± 14	25 ± 3	16 ± 5	7.7 ± 0.9	5908 ± 634	839 ± 115
HZ	LN	Control	651 ± 20	29 ± 7	1316 ± 40	5372 ± 268	62 ± 2	550 ± 18	1539 ± 79	66 ± 5	98 ± 8	968 ± 64	157 ± 13	31 ± 2	14 ± 4	7.7 ± 0.9	9617 ± 307	1223 ± 81
	LN	Tuta	508 ± 53	33 ± 4	1050 ± 110	4209 ± 498	52 ± 6	464 ± 48	1234 ± 222	54 ± 11	78 ± 12	767 ± 132	136 ± 21	24 ± 3	11 ± 4	9.2 ± 0.6	7629 ± 858	980 ± 164
	HN	Control	185 ± 11	10 ± 1	440 ± 27	1322 ± 66	23 ± 2	542 ± 16	236 ± 417	25 ± 4	95 ± 7	532 ± 39	58 ± 4	13 ± 2	0	6.7 ± 0.9	2797 ± 125	685 ± 47
	HN	Tuta	170 ± 22	8±2	407 ± 49	1223 ± 145	21 ± 3	488 ± 57	224 ± 30	18 ± 2	78 ± 13	574 ± 113	57 ± 10	11 ± 2	0	8.8 ± 1.6	2570 ± 300	709 ± 135
ПЗ	LN	Control	560 ± 38	27 ± 1	1398 ± 77	3237 ± 237	61 ± 4	1355 ± 54	659 ± 86	56 ± 9	214 ± 27	1100 ± 184	147 ± 24	23 ± 3	0	8.8 ± 0.8	7377 ± 541	1461 ± 233
	LN	Tuta	534 ± 38	25 ± 2	1357 ± 104	3075 ± 195	58 ± 6	1361 ± 91	525 ± 37	38 ± 2	185 ± 21	905 ± 146	120 ± 21	18 ± 2	0	9.7 ± 1.2	6991 ± 423	1210 ± 187
Stems																	I	
Harvests	Nutr	Treat	CHA1	СНАЗ	CHA4	4CGA	5CGA	1CGA	AR	R	FQA	p CoQA	СР	Tomatin (mg/g DW ⁻¹)	HCAD	FLA		
	HN	Control	228 ± 31	B01 ± 44	4 1614 ± 89	34 ± 6	2392 ± 165	55 ± 4	nd	594 ± 43	89 ± 5	21 ± 2	188 ± 23	nd	4733 ± 298	594 ± 43		
HI	LN	Control	291 ± 24	875 ± 34	1438 ± 131	39 ± 9	2173 ± 162	52 ± 2	nd	485 ± 58	94 ± 5	19 ± 2	1689 ± 16	nd	4482 ± 303	485 ± 59		
	HN	Control	39 ± 6	67±9	403 ± 60	49 ± 8	847 ± 139	56 ± 11	42 ± 4	351 ± 28	45 ± 5	11 ± 1	197 ± 33	4.6 ± 0.3	1734 ± 96	407 ± 31		
	HN	Tuta	46 ± 4	81±8	436 ± 34	66 ± 8	967 ± 53	66 ± 7	47 ± 8	367 ± 24	51 ± 3	12 ± 1	201 ± 12	5.2 ± 0.4	1725 ± 98	413 ± 32		
H2	LN	Control	98 ± 10	167 ± 18	616 ± 36	70 ± 5	1112 ± 66	57 ± 5	74 ± 7	527 ± 27	64 ± 5	13 ± 1	153 ± 12	6.9 ± 0.2	2198 ± 119	600 ± 33		
	LN	Tuta	69±11	120 ± 19	501 ± 85	46 ± 9	908 ± 163	52 ± 11	59 ± 7	484 ± 43	53 ± 7	11 ± 1	145 ± 23	8.5 ± 0.6	2052 ± 68	549 ± 54		
	HN	Control	31 ± 6	50 ± 10	175 ± 36	104 ± 21	513 ± 157	13 ± 3	216 ± 15	695 ± 62	27 ± 6	29 ± 5	32 ± 13	4.4 ± 0.6	943 ± 225	911 ± 73		
	HN	Tuta	33 ± 4	56±6	237 ± 30	132 ± 8	769 ± 154	17 ± 2	232 ± 30	689 ± 47	47 ± 7	33 ± 5	59 ± 19	5.0 ± 0.4	1324 ± 211	921 ± 72		
H3	LN	Control	79 ± 9	146 ± 17	423 ± 37	185 ± 17	1311 ± 110	36 ± 2	484 ± 36	.270 ± 13	47 ± 11	59 ± 8	38 ± 9	7.5 ± 1.1	2287 ± 176	1754 ± 165		
	LN	Tuta	81 ± 6	144 ± 10	429 ± 48	231 ± 16	1393 ± 166	38 ± 2	566 ± 24	1332 ± 49	61 ± 12	58 ± 5	23 ± 3	7.2 ± 0.7	2435 ± 239	1898 ± 69		

ROOLS									
Harvests	Nutr	Treat	5CGA	1CGA	СР	Tomatin (mg/g DW ⁻¹)	HCAD		
LI1	HN	Control	670 ± 109	5.4 ± 0.6	72 ± 17	nd	675 ± 110		
пі	LN	Control	557 ± 73	5.5 ± 1.1	51 ± 12	nd	562 ± 74		
	HN	Control	265 ± 41	29 ± 6.5	27 ± 7	8.9 ± 0.5	294 ± 44		
⊔ 2	HN	Tuta	272 ± 49	23 ± 5.3	34 ± 5	5.9 ± 0.8	295 ± 51		
Π∠	LN	Control	363 ± 46	35 ± 4.9	22 ± 7	7.9 ± 0.5	399 ± 46		
	LN	Tuta	449 ± 74	37 ± 6.1	19 ± 3	6.9 ± 0.6	486 ± 69		
	HN	Control	154 ± 39	9.9 ± 1.7	22 ± 5	13 ± 0.8	163 ± 39		
цэ	HN	Tuta	162 ± 20	5.6 ± 0.7	19 ± 2	13 ± 0.8	167 ± 20		
ПЭ	LN	Control	439 ± 71	9.0 ± 1.0	2.3 ± 0.4	13 ± 1.2	448 ± 72		
	LN	Tuta	652 ± 82	12 ± 0.8	3.2 ± 0.7	14 ± 2.0	664 ± 84		

Roots