Adaptations and responses of the common dandelion to low atmospheric pressure in high-altitude environments

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Abstract
1. Atmospheric pressure is an important, yet understudied factor that may shape plant ecology and evolution.
2. By growing plants under controlled conditions at different experimental stations in the Swiss alps, we evaluated the impact of ecologically realistic atmospheric pressures between 660 and 950 hPa on the growth and defence of different dandelion populations.
3. Low atmospheric pressure was associated with reduced root growth and defensive sesquiterpene lactone production. Defence suppression only occurred in populations originating from lower altitudes. Populations from higher altitudes constitutively produced less sesquiterpene lactones and did not suffer from suppression under low atmospheric pressure.
4. Synthesis. We conclude that atmospheric pressure modulates root growth and defence traits, and that evolutionary history shapes plant phenotypic responses to atmospheric pressure. Our findings have important implications for our understanding of altitudinal gradients and the future use of plants as a source of food and bioactive metabolites in extraterrestrial habitats.

KEYWORDS
adaptation, altitude gradient, atmospheric pressure, dandelion, defence, intraspecific variation, phenotypic plasticity, secondary metabolites

1 INTRODUCTION
Plants successfully colonize a wide range of terrestrial habitats. Their capacity to survive under variable and challenging environmental conditions is a prerequisite for their contribution to biodiversity and sustainable agriculture in the face of climate change, and their potential use as a source of food and medicine in future extraterrestrial habitats (Jump & Peñuelas, 2005; Wheeler, 2017). Atmospheric pressure plays a potentially important role in terrestrial and extraterrestrial environments, albeit for different reasons. On earth, climate change research often relies on altitudinal gradients to understand how plants and terrestrial ecosystems will be affected...
by future climates. Such studies typically assume that differences in climate at different altitudes account for changes in plant performance, and that effects of atmospheric pressure are negligible. In contrast, space plant biology has long been interested in understanding effects of atmospheric pressure on plants, as it would be more practical to maintain low pressure greenhouses in future moon or mars colonies (Corey et al., 1997; Iwabuchi & Kurata, 2003; Richards et al., 2006). Thus, understanding the impact of atmospheric pressure on plants and the capacity of plants to withstand and adapt to different atmospheric pressures is of substantial interest for the present and future of humanity. Yet, to date, our understanding of how atmospheric pressure influences plants is not well-developed.

How are plants affected by low atmospheric pressure? One consequence of low atmospheric pressure at high altitudes is the decrease of partial pressures of $O_2$ and $CO_2$, which are essential substrates for respiration and photosynthesis (Xu et al., 2015; Zabalza et al., 2009). Changes in $O_2$ and $CO_2$ partial pressures have been linked to changes in plant physiology and growth (He et al., 2007; Kammer et al., 2015; Paul et al., 2004; Zhou et al., 2017). Effects are complex, since the decrease of the partial pressure of the atmospheric gases is accompanied by an increase in diffusion rates, which may compensate for the low ambient concentration of the essential gases (Terashima et al., 1995). Further, as the diffusion coefficient for water vapour is increased, transpiration increases (Smith & Geller, 1979), which can impose water stress on plants growing under reduced atmospheric pressure (Iwabuchi & Kurata, 2003; Paul et al., 2004; Richards et al., 2006).

Can plants exhibit phenotypic plasticity to cope with low atmospheric pressure? Several studies suggest that plants respond dynamically to reduced atmospheric pressure (He et al., 2003; Iwabuchi & Kurata, 2003; Richards et al., 2006; Spanarkel & Drew, 2002). In the mountain plant Arabis alpina, for instance, stomatal density increases and stomata aperture narrows at low atmospheric pressure (Kammer et al., 2015). This induced response likely benefits the plant, because at low partial pressures of atmospheric gases, a higher stomatal density may ensure optimal supply of $CO_2$ for photosynthesis (Woodward & Bazzaz, 1988; Xu et al., 2016), while a narrow aperture of the stomata can restrict water loss to counteract increased transpiration rate (Buckley, 2005). At a molecular level, it is assumed that low atmospheric pressure represents an environmental stress to which plants must respond with changes in their metabolic pathways in order to survive successfully (Ferl et al., 2002; Paul & Ferl, 2006). Recent research has documented extensive changes in gene expression patterns in Arabidopsis thaliana when exposed to a low atmospheric pressure environment, including genes associated with hypoxia and water loss (Paul et al., 2004; Zhou et al., 2017).

Can plants adapt to low atmospheric pressure over evolutionary time? Vascular plants colonize habitats between 0 and 6,150 m above sea level corresponding to atmospheric pressures between 101 and 46 kPa (Angel et al., 2016). Since atmospheric pressure influences plant performance, and many plant species occur along altitudinal gradients, local adaptation to atmospheric pressure can be expected (Kammer et al., 2015; Ward et al., 2000; Ward & Strain, 1997). Using growth chambers simulating high- and low-altitude pressure conditions, Kammer et al., (2015) found that Arabis alpina adjusts stomatal density in response to low atmospheric pressure, while the low-altitude plant Arabidopsis thaliana does not. Evidence for genetic differences in the response of stomatal density to pressure conditions was also found within species. Woodward and Bazzaz (1988) showed that in the grass Nardus stricta, plants from higher altitudes developed greater declines in stomatal density at experimentally increased $CO_2$ partial pressure than plants from lower altitudes. Apart from stomatal development, very little is known about evolutionary adaptations of plants to low atmospheric pressure.

Plant secondary or specialized metabolites play important roles in plant responses and adaptations to diverse environments and stress factors (Hartmann, 2007; Moore et al., 2014), including herbivores and pathogens (Bednarek & Osbourn, 2009; Ehlich & Raven, 1964; Kessler & Baldwin, 2001; Moles et al., 2013), abiotic stress (Arbona et al., 2013; Nakabayashi & Saito, 2015; Ramakrishna & Ravishankar, 2011), mutualists (Pichersky & Gershenzon, 2002; Schlemann et al., 2008; Stevenson et al., 2017) and other plants (Baldwin et al., 2006; Semchenko et al., 2014). The production, transport and storage of specialized metabolites is assumed to be costly (Neilson et al., 2013), and plants therefore constantly fine-tune their chemical arsenal to the demands of their environment and internal condition through phenotypic plasticity (Metlen et al., 2009). Over evolutionary times, environmental conditions may act as selective forces on plant genotype selection and can shape genetically determined chemical profiles of plants (Agrawal et al., 2012; Cunningham et al., 1999; Kessler & Kalske, 2018; Züst et al., 2012). With increasing altitude, mountain habitats impose different environmental demands on plants, including harsher abiotic conditions and a lower intensity of biotic interactions (Buckley et al., 2019; Midolo & Wellstein, 2020; Rasmann et al., 2014). Several studies found evidence that the genetic variation of plant secondary metabolites is shaped by these environmental gradients (Bakhitari et al., 2019; Bernal et al., 2013; Bont et al., 2020; Buckley et al., 2019; Moreira et al., 2018). However, the role of decreasing atmospheric pressure at increasing altitudes in shaping the evolution and expression of plant secondary metabolites is poorly understood. Levine et al. (2008) found that the glucosinolate content in the roots of radish was decreased under hypobaric, normoxic conditions. In lettuce, the concentration of phenolics, anthocyanins and carotenoids in the leaves was increased under hypobaric and hypoxic conditions (He et al., 2013). However, all these experiments were conducted in low-pressure chambers under conditions far beyond the natural range of plants. How a plant’s evolutionary history shapes its response to ecologically realistic low-pressure environments is unknown.

Along environmental gradients, closely related asexual and sexual plant taxa often have different distribution patterns—a phenomenon called geographic parthenogenesis (Glesener & Tilman, 1978). It is generally predicted that apomictic plants with asexual reproduction tend to have larger distribution ranges including higher latitudes and higher altitudes, and thus lower atmospheric pressure, than their sexual relatives (Bierzychudek, 1985; Cosendai et al., 2013;
Hypotheses to explain these environmental distribution patterns include different colonization abilities of sexual and asexual organisms as well as different capacities to co-evolve with other organisms (e.g. reviewed in Hörandl, 2006 and in Tilquin & Kokko, 2016). Although many studies empirically support the predicted geographical parthenogenesis patterns, some taxa with asexual and sexual organisms show opposite or mixed trends in geographic distribution. For the common dandelion *Taraxacum officinale* agg. (Asteraceae), a species complex that largely consists of diploid sexuals and triploid asexuals (Verduijn et al., 2004), larger latitudinal distribution of triploids has been observed, with triploids colonizing more extreme environments in the north of Europe (Menken et al., 1995; Van Dijk et al., 2003; Verhoeven & Bierne, 2013). Surprisingly however, triploids are less frequent at higher altitudes than diploids, at least along certain transects (Bont et al., 2020; Calame & Felber, 2000). The reason for this pattern is currently unclear. One possibility is that high-altitude environments impose selection pressures that are different from high-latitude environments. Therefore, one hypothesis to explain the lower success of triploids at high altitude is that they are constrained by low resistance to low atmospheric pressure.

*Taraxacum officinale* possesses a reservoir of secondary metabolites stored in specialized cells, so called laticifers, throughout almost all organs. The latex of *T. officinale* is most abundant in the taproot. The latex is characterized by three major classes of secondary metabolites: hydroxyphenylacetate inositol esters with either two or three side chains (di-PIEs and tri-PIEs), the sesquiterpene lactone taraxinic acid β-d-glucopyranosyl ester (TA-G), and triterpene acetates (TritAc) (Huber et al., 2015). Latex is mainly connoted with defensive functions against herbivores and pathogens (Konno, 2011), and our previous work on the bioactivity and ecological role of the latex metabolites of *T. officinale* confirms this hypothesis (Bont et al., 2017; Huber, Bont, et al., 2016; Huber, Epping, et al., 2016). TA-G in particular reduces the attractiveness of *T. officinale* to white grubs and thereby increases plant performance (Huber, Epping, et al., 2016). Interestingly, in our recent study on the latex metabolites of 63 natural *T. officinale* populations across Switzerland we found a strong association of the latex metabolites TA-G and di-PIEs with the climatic history of the natural populations, which may be suggestively of a role of latex secondary metabolites in abiotic stress tolerance (Bont et al., 2020). Inside the laticifers, latex is maintained at positive pressure, which allows it to be expelled after tissue disruption. Latex likely responds to turgor pressure (Agrawal & Konno, 2009), which in turn is determined by the water balance of the plant. At low atmospheric pressure, evaporation is increased, which could therefore have a direct or indirect effect on latex quality or quantity.

Here, we took advantage of experimental stations in Switzerland between 526 and 3,450 m a.s.l. with standardized abiotic conditions to study the effects of atmospheric pressure on the growth and latex composition of *T. officinale*. We included offspring from nine natural populations from Switzerland from different altitudes, including populations containing both diploid and triploid cytotypes. This setup allowed us to test (a) whether low atmospheric pressure affects plant growth and latex composition through environmental plasticity; (b) whether there are signatures of local adaptation to atmospheric pressure, with populations from higher altitudes performing better at low atmospheric pressure and (c) whether low atmospheric pressure has a stronger impact on triploid than diploid plants, thus potentially restricting the expansion of triploids towards higher altitudes.

## 2 | MATERIALS AND METHODS

### 2.1 Study species and seed collection

The common dandelion is a latex-producing species complex with a world-wide, cosmopolitan distribution (Stewart-Wade et al., 2002). The wind-dispersed perennial herb can tolerate a broad range of environmental conditions and can be found from sea level to altitudes of up to 4,000 m a.s.l. (Molina-Montenegro et al., 2013; Sandoya et al., 2017). In Switzerland, the plant occurs at altitudes of up to 2,000 m a.s.l. (Calame & Felber, 2000). In this study, plants from nine natural populations of Switzerland were included. The nine populations are a subset of 63 populations that were characterized in our previous work (Bont et al., 2020). They cover an altitudinal range from 302 to 1,607 m a.s.l (Figure 1a,b; Table S1). Each population was located within a maximal distance of 1 km from a meteorological monitoring station of MeteoSwiss, the Swiss Federal Office for Meteorology and Climatology, which enabled us to obtain long-term data on climatic conditions. According to Bont et al. (2020), 10 variables were selected from the MeteoSwiss database, representing average air pressure, temperature, precipitation and light conditions of the populations for the years 1996–2015. For most variables, data were available for all nine populations (Figure S1c). Seeds from the natural populations were collected and F2 plants were generated for each population as described in Bont et al. (2020). The cytotype distribution of each population was determined by analysing the ploidy level of the plants with flow cytometry (Bont et al., 2020). The nine F2 populations consisted of populations with only diploid plants (five populations) and populations with both triploid and diploid plants (four populations; Table S1).

### 2.2 Experimental design

In order to investigate the effect of atmospheric pressure on the growth and chemical defence of *T. officinale*, we cultivated the F2 plants from the nine natural populations in four experimental stations at 526 m a.s.l. (Bern), 1,046 m a.s.l. (Adlemsried), 2,061 m a.s.l. (Kleine Scheidegg) and 3,450 m a.s.l. (Jungfraujoch) in Switzerland (Figure 1). Although *T. officinale* does not occur naturally in Switzerland at 3,450 m a.s.l. due to the cold climate, the plant is frequently reported at this altitude in the South American Andes (Sandoya et al., 2017) and is therefore not restricted in its occurrence...
by the associated low atmospheric pressure. In the experimental stations, the plants were grown inside under controlled light supply, temperature and relative humidity and ambient indoor air quality. This allowed us to specifically test the influence of the atmospheric pressure gradient, ranging from 950 to 660 hPa (Figure 1c).

For experiments, seeds from 6 to 8 mother plants (F1) per population were germinated on moist seedling substrate and transplanted into individual 1l pots filled with potting soil (5 parts field soil, 4 parts peat, 1 part sand) after 18 ± 2 days. All plants were grown under controlled conditions (25 ± 2°C, 60 ± 5% RH, 16:8 light:dark cycle) in Bern. 24 hr after transplantation, the plants were distributed among the four experimental stations, so that 6–8 plants from each of the nine populations were cultivated in each station. The atmospheric pressure at the experimental stations was measured at the beginning of the experiment and 2 weeks later. With 0.3%–0.8% difference between the two measurements, the variation was negligible, and we thus used atmospheric pressure as a constant environmental factor in the experiment. The temperature and relative humidity were measured hourly at all stations with RHT10-Data Logger (Extech Instruments, China). To standardize light supply, we grew the plants in rooms with reduced natural light and supplied them with LED lights (400 LED beads, LED type SMD 5730, 400 ± 20 µmol m⁻² s⁻¹) placed 1 meter above the plants. The plants were watered three times a week and fertilized once a week.

Figure 1 Geographical positions of the origins of the *Taraxacum officinale* populations (blue squares 1–9) and of the experimental stations (orange circles). (a) Spatial distribution of the sites across Switzerland. © Photographs: Bern, CC0 public domain; Adlemsried, Zoe Bont; Jungfraujoch, Julius Silver, CC BY-SA 4.0; Kleine Scheidegg, Grindel1, CC BY-SA 3.0. (b) Altitude and atmospheric pressure of the sites and representation of the linear relationship between these two parameters. (c) Visualization of plant growth conditions at experimental stations, including controlled abiotic parameters (grey bars) and varying atmospheric pressure (orange bars). Temperature and relative humidity: mean ± SD (measurement every hour)
2.3 **Assessment of plant traits and chemical analysis**

After 45 days of growth in the experimental stations, we transported the plants back to Bern to analyse growth and defence traits. To assess vegetative growth, we quantified the plant biomass by measuring the dry weight of roots and shoots separately and then calculated root:shoot ratios. To study defence traits, we measured the amount of taproot latex and quantified the concentration of the latex secondary metabolites TA-G, di-PIEs and tri-PIEs.

To quantify latex traits, plants were cut 0.5 cm below the tiller and the taproot latex that was released over 20 s was collected and weighted. Two microlitres of latex was immediately transferred into 200 μl methanol for chemical analysis. The roots were carefully washed with tap water and roots and shoots were placed in a drying oven at 50°C until constant dry mass was reached. The chemical analysis of the latex metabolites was carried out as described in Bont et al. (2017). Briefly, the samples were vortexed for 10 min, ultrasonicated for 10 min, centrifuged at 4°C and 14,000 rpm for 20 min and supernatants were used for further analysis. Relative concentrations of TA-G, di-PIEs and tri-PIEs were determined by injecting the latex extracts into an Acquity UPLC-PDA-MS (Waters) with electrospray ionization in positive mode, consisting of an ultra-performance liquid chromatograph (UPLC) coupled to a photodiode array detector (PDA) and a single quadrupole mass detector (QDa). For quantification, peak areas were integrated at 245 nm for TA-G and at 275 nm for di- and tri-PIEs, while concurrently recorded characteristic mass features were used to confirm compound identities (Bont et al., 2017). For absolute quantification of TA-G, we established an external standard curve with loganin (CAS: 18524-94-2, Sigma-Aldrich Chemie GmbH) and calculated the corresponding response factor to pure TA-G. (Bont et al., 2017). For di- and tri-PIEs, relative concentrations were calculated separately.

2.4 **Statistical analysis**

All statistical analyses were performed in R 4.0.2 (R Core Team, 2017). To represent the climatic conditions associated with the different altitudes of the populations we first conducted a principal component analysis (PCA), as some of the meteorological variables of the population origins were highly correlated (Figure S1c). This approach is widely used to analyse the impact of climate on the evolution of plant traits (Keller et al., 2009; Kooyers et al., 2015; Villaverde et al., 2017). We applied the function ′prcomp′ on scaled variables to reduce dimensionality of the data and selected the axis that explained most of the cumulative variance and represented the climatic conditions associated with the different altitudes at which the populations evolved (climPCA1).

For data exploration, we calculated a correlation matrix of all parameters included in the experiment. Then, we analysed all data with linear mixed-effects models (LMMs; Bates, 2020). The models were fit using the function ′lmer′ from the package ′lme4′ (Bates et al., 2015) with restricted maximum likelihood estimation (REML). Variables representing fixed effects were scaled and centred prior to computation to reduce nonessential multicollinearity (Iacobucci et al., 2016). If necessary, log-transformation was applied to the response variable to improve distribution of variance. Model assumptions were validated using ′plotresid′ from the package ′RVAdMEmoRe′ (Hervé, 2018). The significances of the fixed effects were estimated using the package ′lmerTest′ (Kuznetsova et al., 2017) by calculating type II analysis of variance tables with Kenward-Roger’s approximation to degrees of freedom (Halekoh & Hojsgaard, 2014).

To test the overall effect of the varying atmospheric pressure of the experimental stations on plant growth and defence, we performed LMMs separately for each plant trait (root dry weight, shoot dry weight, root:shoot ratio, taproot latex, TA-G, di-PIEs, tri-PIEs). We used the mean value per trait per population as response variable, tested the fixed effect of the atmospheric pressure of the experimental station (′PStaion′) and included ′(1|Population)′ as random effect and grouping factor to allow for varying intercepts between populations. We performed a similar analysis to assess the overall effect of the climatic conditions associated with the different altitudes of the population origins on plant growth and defence by testing the fixed effect of ′climPCA1′ and including ′(1|Station)′ as random effect and grouping factor to allow for varying intercepts between experimental stations. To visualize the significant effects, linear regression analyses were performed, and adjusted R²-values were calculated.

In order to further investigate interacting effects of the climatic conditions associated with different altitudes of the population origins (genetic effects) and the atmospheric pressure of the experimental stations (environmental effects) on the plants, we next conducted a full model analysis for each plant trait, including the abiotic environment of the stations as covariate. Since temperature and relative humidity were highly correlated (Pearson's r = −0.95, Figure S3), we used only temperature as covariate in the model. Trait values of individual plants were used as response variables, with ′(1|Population)′ as random effect to allow the calculation of separate intercepts for each population. Concurrently, the impact of ploidy was tested by adding the ploidy level (diploid or triploid) of each plant as fixed effect. The full model syntax was the following: plant trait −climPCA1 × PStaion + PStaion × temperature + PStaion × Ploidy + (1|Population). As expected, temperature influenced all experimental parameters (LMMs: Temp, p < 0.05, Table 1), but did not interact with atmospheric pressure (LMMs: PStaion × Temp, p > 0.05, Table 1). A confounding effect of residual variation in temperature on the detected effects of atmospheric pressure is, therefore, unlikely. To visualize the significant interactive effect of ′climPCA1 × PStaion′ on TA-G and on di-PIEs concentration, we then used the package ′effects′ (Fox et al., 2019) for model prediction with unscaled and uncentred fixed effects, excluding the non-significant interaction of ′PStaion × temperature′ from the model. Likewise, we predicted and visualized the nonsignificant effects of PStaion × Ploidy on all measured plant traits. All results
### TABLE 1 Effects of experimental atmospheric pressure (P\textsubscript{Sta\textregistered}), of climatic conditions associated with different altitudes of population origins (climPCA1), of the plant’s ploidy level (Ploidy) and of selected interactions on performance and latex profile of Taraxacum officinale are shown. Temperature of experimental station (Temp) is included as control variable. Results of full mixed-effects model analyses are displayed separately for each performance parameter (root dry weight, shoot dry weight, root:shoot ratio, amount of latex) and for each class of latex secondary metabolites. Significances of fixed effects were assessed by \( F \) tests. Estimated \( F \)-values (NumDF, DenDF) are shown. Levels of statistical significance are indicated with asterisks (**\( p < 0.001 \); *\( p < 0.01 \); *\( p < 0.05 \); (*) \( p < 0.1 \)).

| Performance | Root | Shoot | Root:shoot | Latex | TA-G | di-PIEs | tri-PIEs^a |
|-------------|------|-------|------------|-------|------|--------|-----------|
| climPCA1    | 0.83 (1.4) | 2.08 (1.4) | 0.00 (1.6) | 5.20 (1.4) | 11.00 (1.4) | 2.05 (1.6) | 0.35 (1.6) |
| \( P_{\text{station}} \) | 10.75 (1.209) | 1.29 (1.209) | 18.22 (1.209) | 0.01 (1.209) | 14.12 (1.211) | 0.01 (1.211) | 5.60 (1.211) |
| Temp | 6.26 (1.210) | 4.16 (1.210) | 1.62 (1.209) | 11.41 (1.210) | 4.17 (1.211) | 8.03 (1.211) | 8.21 (1.211) |
| Ploidy | 0.06 (1.108) | 0.92 (1.107) | 0.38 (1.143) | 0.06 (1.114) | 0.23 (1.101) | 1.36 (1.95) | 0.38 (1.209) |
| climPCA1 × \( P_{\text{station}} \) | 0.72 (1.209) | 2.47 (1.209) | 0.40 (1.209) | 0.08 (1.209) | 7.26 (1.211) | 6.49 (1.211) | 2.06 (1.211) |
| \( P_{\text{station}} \) × Temp | 0.49 (1.210) | 0.13 (1.210) | 1.33 (1.210) | 3.32 (1.210) | 0.27 (1.212) | 0.00 (1.212) | 0.09 (1.212) |
| \( P_{\text{station}} \) × Ploidy | 0.67 (1.209) | 1.75 (1.209) | 1.08 (1.209) | 0.50 (1.209) | 0.91 (1.211) | 0.48 (1.211) | 3.30 (1.211) |

Abbreviations: di-PIEs, di-4-hydroxyphenylacetate inositol esters; TA-G, taraxinic acid β-D-glucopyranosyl ester; tri-PIEs, tri-4-hydroxyphenylacetate inositol esters.

^a\( \log \)-transformed.

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were visualized using ‘ggplot2’ (Wickham, 2016) and ‘effects’ (Fox et al., 2019).

### 3 | RESULTS

#### 3.1 | Validation of experimental setup

We first tested whether our efforts to standardize environmental parameters across experimental sites were successful. We found that indoor temperature and relative humidity varied slightly between sites (Figure 1c), but neither variable was correlated to atmospheric pressure (Figure S3). As plants were also grown using identical potting soil, watering, lighting and comparable indoor environments with respect to air pollution, we estimate that major abiotic parameters were successfully standardized to allow for an assessment of the role of atmospheric pressure on T. officinale (see discussion for a critical assessment of this aspect).

By growing T. officinale plants from seeds of diploid and triploid populations originating from different altitudes, we tested (a) the effect of atmospheric pressure on growth and defence; (b) the impact of population origin on plant responses to atmospheric pressure (as an indicator for local adaptation to atmospheric pressure) and (c) the importance of a plant’s cytotype (diploid vs. triploid) on its capacity to grow under different atmospheric pressure, as a test of the hypothesis that low atmospheric pressure restricts the establishment of triploids at high altitudes. To this end, we first describe environmental (Figure 2) and heritable contributions (Figure 3) to variation in defence and growth independently. Using a full model, we then address adaptive phenotypic plasticity (Table 1; Figure 4) and differences between cytotypes (Table 1; Figure 5).

#### 3.2 | Atmospheric pressure modulates root growth and defences

Across all populations and irrespective of population origin and cytotype, T. officinale plants showed distinct phenotypes as a function of atmospheric pressure at the different experimental sites (Figure 2). Both root growth and root:shoot ratio were overall lower at higher altitudes with lower atmospheric pressure (LMEM, \( p = 0.022 \), Figure 2 resp. LMEM, \( p < 0.001 \), Figure 2).

By contrast, no significant effects on shoot growth and on the amount of exuded taproot latex were observed (LMEMs, \( p > 0.05 \), Figure 2). Thus, growth under low atmospheric pressure is specifically associated with lower root growth and lower accumulation of a defensive sesquiterpene lactone in T. officinale.

#### 3.3 | Climatic history associated with altitude shapes heritable variation in root defences

To characterize the climatic conditions under which the T. officinale populations evolved, we conducted a principal component analysis...
(PCA) and then scaled variables to reduce the dimensionality of the data. The first two axes (climPCA1 and climPCA2) together explained 90% of the cumulative variance in climate parameters (Figure S1a). 54.3% of the variance was explained by the first axis (climPCA1) which mainly represents variation in temperature-related parameters and atmospheric pressure (Figure S1a,c). ClimPCA1, but not climPCA2 was highly correlated with the altitude of the population origin (Pearson’s $r = 0.941$, $p < 0.001$, Figure S1b). We therefore used climPCA1 to represent the climatic conditions associated with the different altitudes of the populations.

Across experimental stations, we found no significant association between climPCA1 and T. officinale growth (LMEMS, $p > 0.05$, Figure 3). However, climPCA1 was correlated to the amount of taproot latex and the concentration of TA-G, with populations originating from lower altitudes producing more latex and more TA-G than populations collected at high altitudes (LMEMS, $p < 0.001$, Figure 3). No clear effects were observed for di- and tri-PIEs (LMEMS, $p > 0.05$, Figure 3). Thus, climatic conditions that are correlated with altitude are associated with heritable differences in the production and chemical composition of root latex, with plant originating from higher altitudes producing less latex and defensive sesquiterpene lactones.

### 3.4 Root defences show patterns of adaptation to high atmospheric pressure

In a next step, we constructed a full model to detect interactions between the climatic history of the populations (climPCA1) and the atmospheric pressure of the experimental sites at which the plants were grown ($P_{\text{Station}}$). The full model confirmed that the concentration of TA-G in the latex is influenced by climPCA1 (LMEM: $\text{climPCA1}, p = 0.017$, Table 1, Figure 2), and that atmospheric pressure at the experimental sites is associated with changes in root growth (LMEM: $P_{\text{Station}}, p = 0.001$, Table 1, Figure 3), root:shoot ratio (LMEM: $P_{\text{Station}}, p < 0.001$, Table 1, Figure 2) and TA-G concentration in root latex (LMEM: $P_{\text{Station}}, p < 0.001$, Table 1, Figure 2). In addition, we detected a negative effect of atmospheric pressure on the concentration of tri-PIEs (LMEM: $P_{\text{Station}}, p = 0.019$, Table 1, Figure 2). We detected no significant interactions between the climatic history of the populations (climPCA1) and atmospheric pressure at the experimental sites for root or shoot growth (LMEM: climPCA1 $\times P_{\text{Station}}, p = 0.05$, Table 1). However, a significant interaction was observed for TA-G concentration (LMEM: climPCA1 $\times P_{\text{Station}}, p = 0.008$, Table 1), indicating natural selection for phenotypic variability of TA-G production under different atmospheric pressure. Closer inspection of the data...
revealed that TA-G concentrations in populations originating from low-altitude environments were more strongly influenced by atmospheric pressure than in populations from high-altitude environments (Figure 4a). Under high atmospheric pressure, populations from low-altitude environments produced significantly more TA-G than populations from high-altitude environments (Figure 4a). With decreasing atmospheric pressure, this difference disappeared, leading to similar TA-G concentrations at 660 hPa. A similar pattern, albeit with more variability, was observed for di-PIEs (LMEM: climPCA1 × PStation, p = 0.012, Table 1, Figure 4b). These results are suggestive of adaptation of root defence expression of low-altitude populations to high atmospheric pressure.

3.5 | The performance of triploid plants is not constrained by resistance to atmospheric pressure

To test whether plant cytotype influences the capacity of T. officinale to grow at different atmospheric pressures, we tested for interactions between cytotype and atmospheric pressure at the different experimental sites. Across experimental sites, we did not detect any significant differences between diploid and triploid plants for any of the measured parameters (LMEMs: Ploidy, p > 0.05, Table 1, Figure 5). We also found no significant interactions between cytotype and atmospheric pressure (LMEMs:
Thus, the hypothesis that low atmospheric pressure restricts the performance of triploids at high altitudes is not supported by our data.

4 | DISCUSSION

Plants can adjust to the demands of changing environmental conditions through genetic differentiation and environmental plasticity. Whether and how atmospheric pressure as an environmental factor shapes plant ecology and evolution is poorly understood. By growing several T. officinale populations in controlled environments, our study provides evidence that atmospheric pressure influences plant root growth and chemical defence irrespective of the cytotype level of the plants and shows that the natural habitats of the populations shape the potential for phenotypic variability in response to varying atmospheric pressure. Here, we discuss our findings in an eco-evolutionary and plant-physiological context.

4.1 | Impact of reduced atmospheric pressure on plant biomass

Although many plants seem to be capable to grow vegetatively at pressures even below 25 kPa (Richards et al., 2006), they suffer from stress associated with hypoxia and desiccation and in turn develop responses and adaptations, which are often organ-specific (Zhou et al., 2017). In environments with reduced O₂ and CO₂ availability, roots may react more strongly to oxygen deficiency because, as heterotrophic organs, they are highly dependent on oxygen for mitochondrial energy production, while shoots, as autotrophic plant organs, may be restricted in photosynthesis due to the reduced CO₂ diffusion rate (Mastroph et al., 2014). Our results show that root but not shoot growth of T. officinale is reduced in low atmospheric pressure environments, resulting in a decrease of the root:shoot ratio. In our experimental stations, the reduction in atmospheric pressure is associated with naturally decreased levels of atmospheric gases including a reduced partial pressure of O₂, and we assume that the resulting mild hypoxia restricts root growth of T. officinale. This finding is in line with previous studies showing that roots are particularly sensitive to reduced O₂ partial pressure (He et al., 2007; Tang et al., 2015), whereas several studies with plants cultivated at reduced atmospheric pressure, but with a partial pressure of O₂ experimentally maintained at the same level as at ambient pressure, found no reduction in root biomass (Iwabuchi et al., 1996; Levine et al., 2008; Spanarkel & Drew, 2002).

4.2 | Impact of reduced atmospheric pressure on the sesquiterpene lactone TA-G

Exposure to low atmospheric pressure also affects plant chemistry (He et al., 2013; Levine et al., 2008; Zhou et al., 2017). In our study we
detected a decline in the concentration of the secondary metabolite TA-G in the latex of *T. officinale* with decreasing atmospheric pressure. TA-G is constitutively produced by the plant, acts repellent against root feeders and therefore defends the plant against herbivores (Bont et al., 2017; Huber, Epping, et al., 2016). To reduce the costs associated with constitutively produced chemical defences (Neilson et al., 2013), plants may use abiotic conditions as external stimuli to adjust the level of defensive secondary metabolites to the expected herbivore pressure. Temperature, for instance, is assumed to be a good indicator for herbivore attack in the field, because herbivore appearance and activity is often modulated by temperature (Bale et al., 2002). In our previous work, we found evidence that *T. officinale* can use seasonal temperature variation to synchronize deployment of chemical defences with expected herbivore attack intensity in the field, indicating an important role of abiotic conditions in fine-tuning the level of constitutively produced defensive metabolites (Huang et al., 2019).

With increasing altitude, it is often assumed that herbivore pressure decreases (Moreira et al., 2018). *T. officinale* might thus use low atmospheric pressure as an indicator of high-altitude growth conditions and associated expected lower herbivore pressure, and the observed decrease in TA-G production might be a fine-tuning of defence deployment to reduce costs and maximize plant fitness. The seeds of *T. officinale* are characterized by excellent flight abilities (Cummins et al., 2018) and occasionally disperse over distances of several kilometres using wind and convective updrafts (Tackenberg et al., 2003). Hence, along altitudinal gradients, *T. officinale* is expected to regularly colonize novel territories with different environmental requirements, which could promote the evolution of adaptive plastic responses that use atmospheric pressure as an abiotic signalling factor for local conditions, as suggested. However, these are highly speculative conclusions and require further investigations.

Our previous work has not only shown that TA-G is involved in herbivore defence in *T. officinale*, but it has also shown an association of TA-G with the climatic history of the plants’ natural habitats, suggesting an additional role of TA-G in abiotic stress management (Bont et al., 2020). The climatic parameters associated with TA-G production included sun

**Figure 5** Visualization of influence of ploidy on plant parameters depending on atmospheric pressure of experimental station. Predicted mean values with standard errors from mixed-effects models are shown. None of the plant parameters differ statistically significantly between diploid and triploid plants (n.s.: *p* > 0.05). 2n: diploid; 3n: triploid; TA-G: taraxinic acid ß-D-glucopyranosyl ester; di-PIEs: di-4-hydroxyphenylacetate inositol esters; tri-PIEs: tri-4-hydroxyphenylacetate inositol esters.
and rain intensity and were related to the latitude but not to the altitude of the natural populations—plants from the rainier North of Switzerland produced more TA-G than plants from the sun-intense regions in the South of Switzerland (Bont et al., 2020), leading to the hypothesis that TA-G may passively or actively be involved in moisture regulation. At low atmospheric pressure, water evaporation is increased, and evidence exists that when plants are grown under such conditions, their perceptual mechanisms of water movement are altered even when the plants are fully hydrated and do not experience actual desiccation (Paul et al., 2004; Zhou et al., 2017). The decline in TA-G concentration with decreasing atmospheric pressure observed in our study could therefore be a direct or indirect consequence of increased evaportranspiration and would then support an involvement of TA-G in moisture regulation.

### 4.3 | Heritable variation for plasticity in TA-G production indicates adaptive value

The capacity of phenotypic plasticity allows a plant with a given genotype to adjust its phenotype to the demands of contrasting environments (Nicotra et al., 2010; Sultan, 2000). In our experiments, the employed atmospheric pressure gradient was associated with measurable phenotypic plasticity, which again emphasizes the importance of atmospheric pressure for plant growth and development (Paul & Ferl, 2006). For the latex metabolites TA-G and di-PIEs, plastic responses differed among populations, indicating heritable within-species variation for plasticity in these traits, likely shaped by the climatic histories of the populations. In high atmospheric pressure environments, plants from low-altitude environments produced more of the metabolites than plants from high-altitude environments, whereas in low atmospheric pressure environments, the difference between the populations vanished. The capacity of low-altitude plants to increase production of TA-G and di-PIEs when grown under low-altitude atmospheric pressure could benefit these plants and may be an adaptive trait, as latex metabolites have defensive functions against root feeders (Bont et al., 2017; Huber, Bont, et al., 2016; Huber, Epping, et al., 2016) and herbivore pressure is often expected to increase with decreasing altitude (Moreira et al., 2018; Rasmann et al., 2014). These findings are consistent with previous studies showing that within species, constitutive defences often decrease with increasing altitude (Bakhtiari et al., 2019; Buckley et al., 2019; Meyer & Carlson, 2001; Pellissier et al., 2014). Our results further suggest that climatic conditions characterizing low-altitude environments select for genotypes with high plasticity in latex metabolite production, although whether the observed plasticity is a passive or active plastic response (van Kleunen & Fischer, 2005) and whether its expression increases plant fitness remain to be elucidated.

### 4.4 | Plant performance at reduced atmospheric pressure does not differ between cytotypes

Along altitudinal transects, diploid sexual _T. officinale_ predominate over triploid asexual individuals at higher altitudes (Calame & Felber, 2000), while along latitudinal gradients, triploids predominate over diploids at higher latitudes (Van Dijk et al., 2003). In our study, we tested the hypothesis that the performance of triploids at high altitudes is restricted by low atmospheric pressure. However, we found no evidence for a disadvantage of triploids at low atmospheric pressure, as diploids and triploids did not differ in any of the measured plant traits at different atmospheric pressures. The measured plant traits, however, estimate early plant performance and exclude traits related to dispersal, colonization and survival because the plants in this study were harvested before seed production. Therefore, although at reduced atmospheric pressure diploids and triploids exhibit similar growth and defence traits, we cannot rule out the possibility that the cytotypes differ in other fitness-determining traits that are crucial for long-term establishment. Further experiments on these traits would complement our results and provide a more comprehensive assessment of the role of atmospheric pressure in the geographic parthenogenesis of _T. officinale_.

### 4.5 | Atmospheric pressure should be considered in studies involving altitudinal gradients

Studies comparing different species have shown that plants growing in high-altitude environments have developed adaptations to cope with the challenges of these environments (Halbritter et al., 2018) and that some of these adaptations, for example changes in stomatal development, are directly related to atmospheric pressure (Kammer et al., 2015). Our study confirms that atmospheric pressure is an important abiotic factor that influences plastic responses in plant growth and development and shows that across _T. officinale_ populations, exposure to varying atmospheric pressures evokes heritable responses in growth and defence traits—responses that are shaped by the climatic conditions of the natural habitats of the populations. Given the non-negligible impact of atmospheric pressure on the expression and likely also on the evolution of plant traits, we suggest that atmospheric pressure should be included by default as an abiotic factor when studying plant variation along altitudinal gradients to prevent results from being obscured by possibly unrecognized effects of atmospheric pressure variation. This may be particularly important when altitudinal gradients are used to surrogate climate change (Carlyle et al., 2014; Frei et al., 2014; Michalet et al., 2014), since atmospheric pressure, unlike temperature or precipitation, is not expected to change under global warming.

### 4.6 | Experimental considerations and conclusions

An underlying assumption of our work is that we successfully standardized or randomized environmental parameters other than atmospheric pressure across the different experimental stations, thus allowing us to infer effects of this factor. Indeed, our experiment allowed us to control and/or randomize all major abiotic and biotic environmental parameters, including soil structure and composition, water supply, humidity, temperature as well as light quality and quantity and air pollution. Although
temperature and relative humidity differed slightly among experimental stations, we found no significant interaction with the effect of the atmospheric pressure on plant growth and defence and conclude that our results are not confounded by these abiotic variables. Pests and pathogens were not observed on our plants. Another factor that depends on altitude and covaries with atmospheric pressure is gravity. Gravity decreases with altitude, resulting in a delta of 0.01 m/s² between Bern and the Jungfraujoch. Whether such a small change in gravity has any measurable impact on plants is unknown. Plant gravitropism depends on sensing inclination rather than gravitational force (Chauvet et al., 2016), and effects of reduced gravitational force are typically only observed below 0.3 g (Kiss et al., 2019). Thus, we infer that the effects observed in our study are the result of changes in atmospheric pressure rather than other environmental factors. Further experiments with pressure chambers (Paul et al., 2004) could be used as a future approach to confirm the patterns observed in our study.

Our results emphasize that within species, plants from different populations respond differently to varying atmospheric pressure, especially in the production of secondary metabolites. For successful cultivation of plants in extraterrestrial habitats as food source, it therefore may be worthwhile to screen various populations of the species of interest under low atmospheric pressure to detect resistant populations with secondary metabolite profiles suitable for human nutrition.

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CONFLICTS OF INTEREST
The authors have no conflict of interest to declare.

AUTHORS’ CONTRIBUTIONS
M.E., Z.B. and C.C.M.A. designed the study; C.C.M.A., R.A.R.M. and Z.B. collected data; Z.B., C.C.M.A. and P.F.C. analysed and interpreted the data; Z.B. and M.E. wrote the first draft of the manuscript. All authors contributed to the final version of the manuscript.

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