1	The ecological and genetic drivers of silicon accumulation in cereal crops
2	$_{ m by}$
3	Isaac Peetoom Heida
	A THESIS SUBMITTED IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE
4 5	DEGREE OF
3	DEGITED OF
6	MASTER OF SCIENCE
6 7	in
8	THE FACULTY OF GRADUATE AND POSTDOCTORAL STUDIES
9	(Plant Science)
10	THE UNIVERSITY OF BRITISH COLUMBIA
11	(Vancouver)
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# 1 Abstract

As global agricultural production strains under degrading soil fertility and increasing losses due to climate change, there is growing research interest in new avenues for production 53 improvement. New crop technologies must meet increasing public and regulatory demand for environmental sustainability, encouraging scientists to revisit overlooked or relatively unknown techniques that may unlock productivity gains. One of the promising developments to arise over the past 20 years is the potential of silicon to improve crop plant performance. With benefits to multiple dimensions of crop performance, silicon may be a key tool to guard crop production against uncertain future growing conditions. Our ability to mobilize silicon-based cropping strategies is dependent on a thorough understanding of the ultimate and proximate causes of silicon accumulation, including both the ecological and genetic 61 interactions that can trigger increased uptake. In this thesis, I extend recent advances in our understanding of silicon ecology in cereal crops, testing for the presence of rapid silicification in common canadian crops, as well as using a genome-wide association study to identify genetic markers associated with high silicon content. 65

[to add: results, conclusion]

# <sub>67</sub> 2 Lay Summary

Silicon provides tremendous benefits to plant health, but is not widely utilized in agriculture,
despite it being highly abundance and naturally occurring in most soils. One of the main
factors limiting it's application in agriculture is our poor understanding of the exact dynamics
of how plants absorb and use silicon from the soil. I identified genetic traits associated with
high silicon content in Aegilops tauschii, a relative of bread wheat, as well as demonstrated
that cereal crops (e.g. wheat, barley, oats) have the ability to rapidly uptake silicon from
the soil. This rapid uptake means that silicon may be a highly effective defence against
insect pests. Combining these results with the genetic data, future research can aim towards
creating breeding programs to develop cereal crops that can withstand insect damage based
on their silicon content. This development could provide an environmentally friendly strategy
to maintain output to feed a growing human population.

### <sup>9</sup> 3 Preface

(This was taken nearly verbatim from Matt's thesis, so need to go back through to make sure I am not plagarizing)

The research presented in this thesis is original and unpublished. Isaac Peetoom Heida and Dr. Juli Carrillo, with assistance from Dr. Jean-Thomas Cornelis, conceptualized and developed the experiment presented in Chapter Two. Isaac Peetoom Heida, Dr. Juli Carrillo and Dr. Gurcharn Singh Brar, with input from Dr. Jean-Thomas Cornelis, conceptualized and developed the experiment presented in Chapter Three. Isaac Peetoom Heida developed the question and methodology for Chapter Two. Dr. Aaron Beattie, Dr. Mazen Aljarrah, and Dr. Gurcharn Brar provided seeds for the experiment. Isaac Peetoom Heida designed and set up the experiment, processed and analysed the samples, and performed the statistical analysis. Dr. Shaun Barker and the Mineral Deposit Research Unit of the University of British Columbia provided facilities and expertise for the XRF analysis of the tissue samples. For Chapter Three, Isaac Peetoom Heida led plot set up and maintenance, with assistance from Grace Wang, Vincent Fetterley, Sara Salad, Katherine Buchanan, Martina Clausen, and Paul Fisher, and Matt Tsuruda. Isaac Peetoom Heida led the sample harvest, processing

and analysis. Kelly Wang, Grace Wang, and Chelsea Gowton assisted with the sample harvesting. Dr. Daria Reshetniak, Paul Fisher, Lucas Friesen, Katie Pryer, Dr. Kinga Treder, Carly MacGregor, and Grace Wang all provided invaluable assistance with sample preparation. Chapters Two and Three of this thesis will be submitted to peer-reviewed journals for publication. For the purposes of this manuscript, actions are depicted in the singular first person.

### $_{\tiny ext{.01}}$ 4 Introduction

#### <sup>02</sup> 4.1 A case for silicon in agriculture

As global agricultural production strains under degrading soil fertility and increasing losses due to climate change, researchers are leaving no stone unturned in the search for new technologies for sustainable improvement in crop production. New crop technologies must 105 meet increasing public and regulatory demand for environmental sustainability, encouraging 106 scientists to revisit overlooked or relatively unknown techniques that may unlock productivity 107 gains. Over the past 20 years, plant-silicon relations has emerged as a promising field that 108 may safeguard crop performance and security within a changing biosphere. With benefits to 109 multiple dimensions of crop performance, silicon may be a key tool to guard crop production 110 against uncertain future growing conditions. 111

#### $_{112}$ 4.2 Silicon in nature

Silicon abounds in the earth's crust, with various silicates, such as silicon dioxide (SiO<sub>2</sub>), 113 comprising about 60% of the crust by mass. Nearly all terrestrial plants grow in soils containing silicon, and thus absorb nominal amounts through passive transport as water is 115 absorbed into the plant. Silicates are found in a variety of forms, and vary in their plant 116 availability. Crystalline forms, such as quartz, are highly resistant to weathering, and are 117 poor sources of plant-available silicon, while amorphous forms of SiO<sub>2</sub> are more available 118 Fraysse et al. 2009. As soils age, an increasing share of the plant-available silicon is derived 119 from biogenic silicates, such as diatom testes or plant phytoliths F. de Tombeur et al. 2020. 120 Uptake and usage of silicon is not uniform throughout the vascular plants. Plant clades vary

in their tissues. Plant phytoliths are amorphous masses of silica, chemically similar to opal, 123 that are found throughout the plant body and have highly variable geometries (Piperno 124 2006). Phytolith morphologies are generally conserved within taxonomic groups, allowing 125 their use as a tool for paleobotanical investigations. The highly variable morphology among 126 clades, with the conservation of morphology within clades suggests active selection upon 127 the structure of phytoliths. The morphology of phytoliths also varies between organs in the 128 plant. Stem phytoliths tend to be more elongate, with putative structural function, while leaf 129 phytoliths are typically more orbicular, and are likely an adaptation to counter herbivory. 130 Silicon deposition may create apoplastic barriers that seal and toughen the plant tissue, 131 which may benefit the plant in a multitude of ways (Coskun et al. 2019). Under this hypoth-132 esis, silicon deposits reduce water loss and radiation/temperature damage, and also limit the 133 spread of effector compounds, dampening the effects of fungal pathogen and herbivore excre-134 tions designed to interfere with plant defensive physiology (Coskun et al. 2019). Empirical 135 evidence shows that silicon is effective at limiting the growth and damage of insect and fungal 136 pests (Fauteux et al. 2005; F. P. Massey et al. 2007). The toughness of the depositions also 137 serves a more direct mechanical role, as the hardened granules of silicon interrupt the chewing 138 motions of herbivores, wearing down mandibles and teeth (Strömberg et al. 2016; Waterman, 139 Cibils-Stewart, et al. 2021), and reduce the digestive efficiencies of herbivores (Johnson et al. 140 2021). Indeed, when supplemented with silicon plants are generally more resistant to a wide 141 spectrum of stressors, including soil salinity, soil metal toxicity, cold and heat stress, UV 142 stress, water deficits, and phosphorus deficiencies (Cooke and Leishman 2016). Continuing 143 to untangle the various mechanisms through which silicon delivers beneficial effects to plants 144 is key to fully realizing the potential of silicon in sustainable agriculture. 145

in their expression of silicon transporter proteins and in the relative abundance of phyotliths

#### 146 4.3 Silicon in soils and roots

122

Plants interact with silicon on a variety of levels, mobilizing it from soil aggregates, transporting it into and throughout their bodies, and finally precipitating it out of their xylem into
solid masses in the leaves and stems. Within the soil environment, silicon commonly exists
in both crystalline (geologic) and amorphous (biogenic) forms (Haynes 2014). Amorphous

silicates can derive from previous plant material that has decayed in the soil, but also from 151 marine and aquatic organisms such as diatoms. Globally, the silicon cycle involves silicates 152 weathering out of terrestrial sediments, moving along water courses, and eventually being 153 deposited in the sea, where it is incorporated into various plankton species, and eventually 154 deposited in seafloor sediments. The continual exodus of silicon from terrestrial sediments 155 over geologic timescales means as ecosystems age, plants become more and more central in 156 the local silicon cycle, with much of the silicon in living plant tissue being recycled from 157 previous plant material decaying in the soil (F. de Tombeur et al. 2020). In highly weathered 158 soils with low nutrient availabilities, plants take a more active role in liberating nutrients, 159 including silicon, for uptake. Organic acids and chelating agents, exuded from plant roots, pry tightly bound nutrients such as phosphorus and silicon from soil aggregates, increasing their availabilities for uptake into the root system (Félix de Tombeur, Cornelis, et al. 2021). This active scavenging for silicon remains poorly understood, but it may be an important 163 mechanism in plant defence (allowing for increased uptake during a defensive response) and 164 breeding for increased root exudation may improve crop plant performance and nutrient use 165 efficiency (Félix de Tombeur, Roux, et al. 2021). 166

## 167 4.4 Silicon transporters

One of the most important advances in plant-silicon research was understanding the mech-168 anisms through which silicon is acquired and transported into the plant. Silicon's most 169 common form in soil solution is silicic acid (H2SiO4), which has a maximum solubility of around 2 mM (Haynes 2014). While there is some evidence that small amounts of silicic acid can be transported during water uptake, this method of transport is insufficient to explain the larger amounts of silicon found in some plant families. Research in rice has identified 173 four gene products that transport silicon into and through the plant body. Two of these 174 (LSi1, LSi2) transport silicic acid from the soil into the roots, while the other two (LSi3, 175 LSi6) act to unload silicic acid from the xylem into leaves and inflorescences (Yamaji, Saku-176 rai, et al. 2015). Orthologs of these proteins have been identified in other cereal crops, and 177 additional analogous silicon transporter proteins have been discovered in the Cucurbitaceae 178 (Reynolds et al. 2016). Though not identified, there is a hypothesized fifth protein responsi-

ble for loading silicic acid into the xylem (Farooq and Dietz 2015). The expression of these 180 genes, or lack-there-of, can not only influence the total amount of silicon accumulated by 181 the plant, but also its relative distribution, as knockout of LSi6 increases leaf silicon con-182 tent while decreasing the silicon content of seed husks in rice Yamaji, Mitatni, et al. 2008. 183 Breeding for silicon content and use-efficiency in crop plants may be crucial to improving 184 crop performance under a changing climate Christian et al. 2022. However, we still have a 185 relatively poor understanding surrounding the how genetics influence the silicon phenotype 186 of a plant. Further investigations into how genotypic variation is reflected in the silicon 187 content of plants can aid in the discovery of new genes involved in silicon accumulation, and 188 may provide targets for silicon breeding programs.

#### 190 4.5 Silicon in leaves

Once inside the plant, silicon is deposited in specialized silica cells, forming phytoliths (Wa-191 terman, Hall, et al. 2021). Silicon deposits show consistent and taxa-specific morphologies, 192 suggesting evolutionary pressure selecting for these structures to yield certain functions to 193 the plant (Piperno 2006). In stems, these phytoliths are often long and narrow, oriented 194 parallel with the shoot, and seem to increase structural rigidity (Strömberg et al. 2016). The 195 use of silicon as a structural component represents a highly energetically efficient strategy, 196 as silicon is 10x cheaper on an energy unit basis to produce than lignin (Strömberg et al. 197 2016). Stem silicification has been investigated as it relates to lodging resistance in cereal 198 crops, and silicon supplementation has been found to reduce the prevalence of lodging in rice and wheat (Dorairaj et al. 2017; Muszynska et al. 2021). In leaves, phytoliths are typically more stout, though they still increase the mechanical toughness of the leaf (Simpson et al. 2017). This overall toughness, and abundance of phytoliths in leaves likely evolved to 202 limit herbivore damage, rather than improve the growth characteristics as in stem phytoliths 203 (Strömberg et al. 2016). Interestingly, even in the absence of silicon, plants develop silica 204 cells, and rapidly fill them when silicon becomes available (Waterman, Cibils-Stewart, et al. 205 2021). As phytoliths develop in the leaves, polymerization of monosilicic acid is aided by 206 interactions with proteins in the cell wall, which act as sites of nucleation (Nawaz et al. 207 2019). Silicon deposition in the leaves can happen on relatively short time scales, outpacing the accumulation of other defensive compounds such as phenolics (Waterman, Hall, et al. 2021). Thus, silicon-based defences in crop plants may be one of the first lines of active anti-herbivore defence, providing rapid and sensitive responses to herbivory.

## 12 4.6 Looking back and looking forward

Much of today's plant silicon work is indebted to the pioneering work of Jones and Handreck (1967) and the subsequent mapping of silicon across the plant kingdom by Takahashi et al. 214 (1990). Epstein's seminal 1999 paper provided a comprehensive review of the state of knowl-215 edge in plant silicon, and has spurred a generation of researchers to extend the preliminary 216 findings of the 20th century out across crop production systems and plant ecologies around 217 the world. Silicon is best studied in the grass family (Poaceae) due to the comparatively 218 high silicon content found in most members of the family (often over 1% of dry weight), 219 as well as the economic importance of domesticated species within the clade (Reynolds et 220 al. 2016). Rice, maize, wheat, and barley alone account for one-third of the worlds' total 221 cultivated land area (FAO 2022), and are all domesticated grass species. Silicon supplemen-222 tation as an agricultural practice has been extensively studied in rice and sugar cane, as 223 these crops tend to deplete soil silicon stocks, necessitating replenishment by application of 224 silicon-rich amendments (Haynes 2014; Meena et al. 2014). Due to the overall high silicon 225 content of soils globally, Si is rarely truly limiting in soils, though certain forms of silicon 226 are much more plant available than others (Fraysse et al. 2009). Thus, the applicability and importance of silicon supplementation is unlikely to be realized in more temperate dry-land 228 production systems, particularly in wheat and barley. This does not however nullify the utility of silicon research in these systems, as great work can still be done to improve the 230 manner and efficiency in which these temperate crops utilize the ample silicon available in 231 their soils. Our ability to integrate silicon as a tool for production improvement in crop 232 production is currently limited by a poor understanding of the genetic controls over silicon 233 accumulation, as well as a limited understanding about the extent to which crops utilize 234 silicon in pest-protection.

# 5 Chapter 1: Identifying rapid silicon accumulation in cereal crops

#### 238 5.1 Introduction

To address acute damage from herbivores, plants have developed a host of defensive strate-239 gies, ranging from changes to the body plan down to the development of novel compounds to 240 poison those that would try to eat the plant (Agrawal and Fishbein 2006). In the broadest of terms, plant defenses can be categorized as either tolerance of resistance strategies. Resistance strategies involve attempts to limit tissue loss through interference with herbivory, 243 while tolerance strategies result in increased regrowth after tissue loss. Due to the vastly different nature and ontogeny of various defensive strategies in plants, plant defences oper-245 ate across a range of intensities and time scales, from short-term temporary activation, to 246 long-lasting changes in the morphology of the plant (Agrawal and Fishbein 2006; Karban 247 and Myers 1989). In most scenarios, induced plant defences are activated in response to 248 an external cue, and build in intensity over time, with defensive hormone signals peaking 249 approximately five hours after the initial induction event (Schmelz et al. 2003). Despite this 250 rapid hormonal response, actual defensive phenotypes are slower to emerge, often operating 251 on the scale of days or generations (Karban and Myers 1989). Many defensive responses are 252 also context dependent, where the identity of the damaging actor, the severity of damage, 253 and a host of other factors interact to determine the final defensive response (Waterman, 254 Cazzonelli, et al. 2019). The most effective defensive strategies should be those that can ei-255 ther prevent herbivory outright, or can mount a rapid response to limit damage. These same 256 strategies are also the most promising for crop production, where pest damage represents 257 both an economic and food security cost. Integrating better natural plant defences into crop production systems may be key to reducing the environmental impact of agriculture, but hinges upon a thorough understanding of plant defensive physiology. 260

One of the most promising avenues for new crop defence is the harnessing of silicon (Reynolds et al. 2016). Silicon acts on multiple temporal and physiological scales, delivering broad spectrum resistance to pests, pathogens, and abiotic stressors (Cooke and Leishman 2016; Coskun et al. 2019). Soluble silicon taken up from the soil is deposited predominantly

in the leaf epidermis, where it forms solid granules that increase the toughness of the tissue, 265 reducing herbivore digestive efficiency (Cooke and Leishman 2011). Plant silicon is expressed 266 latently, but also increases in response to herbivory (Takahashi et al. 1990). Multiple studies 267 have demonstrated lasting elevated silicon in response to real and simulated herbivory (F. 268 Massey et al. 2008; Hartley and DeGabriel 2016), and recent evidence points to silicon 269 accumulation as being a relatively rapid response, even preempting some chemical defences 270 (Waterman, Hall, et al. 2021). This rapid action makes silicon accumulation a promising 271 trait for future crop development. Despite the novel results, this pattern has so far been 272 observed in just one species, and only under artificial herbivory via the application of methyl-273 jasmonate. Though a useful tool for herbivory research, methyl-jasmonate application fails to 274 reproduce a complete herbivory signal for the plant, thus observed changes to plant defence 275 may not be representative of a true herbivory scenario (Strauss et al. 2002). Testing for this rapid silicon accumulation across a variety of grain crops, and under both simulated (methyl-277 jasmonate) and real herbivory is a crucial first step towards integrating rapid silicification 278 into our understanding of plant defence and crop protection. 279

Plant silicon research has mostly focused on members of the grass family (Poaceae) 280 due to their exceptional silicon content within the plant kingdom, as well as the economic 281 importance of domesticated grass species (Reynolds et al. 2016). Domesticated crops differ 282 significantly from their wild relatives, due to effects of strong selective pressure imposed by 283 humans (Chen et al. 2015). Most domesticated crops show much lower genetic diversity 284 than their wild ancestors (Hafeez et al. 2021; Smith et al. 2019). Initial selection for a 285 few individuals with favourable traits creates a genetic bottleneck, and the majority of 286 allelic diversity is lost. Subsequent selection by humans for agronomically relevant traits 287 can result in concurrent losses of adaptations to natural environments, as the traits that 288 maximize human value (eg. yield, ease of harvest) can come at the cost of ecologically 280 relevant traits such as defence (Whitehead et al. 2017; Chen et al. 2015). Indeed, in the 290 context of silicon, we can detect clear signals of domestication across the Poaceae family, 291 where wild ancestors consistently have higher baseline silicon content than their domesticated 292 descendants (Simpson et al. 2017). Due to the effects of selection on plant defence it becomes crucial to test new developments in the silicon-defence literature in modern crop species, both

- to validate their utility towards agricultural production, and to gather further observations on the dynamics of silicon-based defences in the first hours after herbivory.
- In this study, I tested four globally important cereal crop species for rapid silicon accumulation under artificial and real herbivory. In a glasshouse environment, I grew bread wheat (*Triticum aestivum*), oats (*Avena sativa*), barley (*Hordeum vulgare*) and Triticale (× *Triticosecale*), and tested the following hypotheses:
- 1. Rapid silicon accumulation is a conserved trait in the Poaceae, and the tested species silicon content would show a significant increase in silicon content within 18 hours of the herbivory treatment applications.
- 2. Due to different phylogeny and domestication history, the tested species would vary in
  the strength of their silicon accumulation response to herbivory.
- 30. Due to the different cues involved when comparing true herbivory damage and methyljasmonate induced defensive induction, the tested species would show different patterns
  of short-term silicon accumulation in response to cricket (*Acheta domesticus*) herbivory
  and methyl-jasmonate application.
- This study is a thematic replication of Waterman et al.'s 2021 paper, but attempts to extend
  the findings to commercially important grain crops. The findings of this study will refine
  our understanding of the prevalence of rapid silicification in the Poaceae, and will help to
  inform the value of potential applications of silicon-based defences into grain crops.

#### 314 5.2 Methods

#### 5.2.1 Plant growth and experimental treatments

To test the prevalence of rapid silicon accumulation in canadian cereal crops, I selected three cultivars for each of oats, bread wheat, triticale, and barley. I selected cultivars on the basis of minimizing shared pedigree, and no cultivars shared more than one common ancestor within the last two crossing generations. At the start of the experiment, I germinated seeds in germination trays filled with moist sand. After four days, I transplanted germinated seedlings into 10cm pots filled with SunGro potting mix amended with [amount] of silicic

acid. Though potting mix and fresh water contain some amount of plant available silicon, 322 I added the silicic acid to ensure that there would be no silicon limitation to the plants. I 323 randomized the location of each pot within the growing space. A flood table bottom watered 324 the pots with nutrient solution. I assigned each plant to one of three herbivory treatments: 325 control, simulated herbivory, or true herbivory. I simulated herbivory by application of 1 326 mM methyl-jasmonate solution to the entire above-ground portion of the plant (Waterman 327 et al. 2021b), while crickets housed in water-pik tubes provided true herbivory. Prior to 328 introduction to the plants, I acclimated crickets by feeding them on the same species 329 used in this trial. Immediately preceding cricket application, I placed them in their tubes 330 and starved them for 24 hours, as this increased the likelihood of the insects initiating feeding 331 rapidly upon exposure to the test plants. Prior to harvest, I recorded whether the crickets had initiated feeding on the plants by visual inspection of the leaves for missing tissue. 333

#### 334 5.2.2 Sample harvest and preparation

18 hours after treatment application, I harvested three fully expanded upper leaves the
plants, and split the leaves in half along the midvein. I placed one half of the tissue into a
coin envelope, oven dried it for 4 days at 60°C, transferred it to a 2ml microcentrifuge tube
with three 3.2mm diameter steel beads, and ground it into a fine powder using a tisselyser
(60 seconds at 30 Hz) in preparation for XRF analysis. I placed the other half of the tissue
into a microcentrifugre tube, flash froze it in liquid nitrogen, and subsequently freeze dried
it. After freeze drying, the samples for phenolic processing were ground under the same
conditions as the other samples.

#### 343 5.2.3 Silicon analysis

To measure the silicon content of the leaf tissue, I followed a modified version of the benchtop XRF method (Reidinger et al. 2012). I pressed leaf powder in a hydraulic press at 11 tons
of pressure, using a 13mm die to create a pellet. I then placed the pellet in an Olymus Vanta
pXRF mounted in a benchtop stand, and used a 45 second scan time to quantify silicon.
After each use, I cleaned the pellet die and XRF analyzer to minimize contamination between
samples.

#### 350 5.2.4 Statistical analysis

Despite the starvation, some crickets did not initiate feeding during the exposure period. 351 I filtered out plants assigned to the insect induction treatment that recieved no damage, 352 to ensure that they would not confound the model. Prior to running our full model, I 353 first tested for an effect of biomass of silicon content, as other defensive pathways show 354 correlations between plant size and defense levels Carmona et al. 2011. We found a negative 355 correlation between plant size and silicon content ( $\beta = -0.067 \pm 0.017, p < 0.001$ ), and thus 356 included plant size as a covariate in our final model. To test all three of our hypotheses, I used linear mixed effects models and tested the effects of our species and induction treatments on measured leaf silicon using the following model formula: 359

$$Si \sim Species * Induction + Biomass + (1|Genotype)$$

#### 360 5.2.5 Software used

I compiled the final dataset using DataFrames.jl (Kamiński et al. 2023) in Julia 1.8.5 (Bezanson et al. 2017). I implemented the biomass regression using GLM.jl (Bates, Alday, et al. 2023). I tested the full mixed effects model in R 4.2.2 (R Core Team 2022) using lme4 (Bates, Mächler, et al. 2015), and performed a post-hoc tukey test using emmeans (Lenth 2023). I generated graphics in Julia using Plots.jl (Breloff 2023).

#### 366 5.3 Results

Among the various cultivars, average uninduced silicon content ranged from 0.26% to 0.91% (Figure 1). Amongst species, oats had the lowest average silicon content at  $0.34\pm0.02\%(\mu\pm$  SE), while wheat had the highest average silicon content at  $0.76\pm0.05\%(\mu\pm SE)$  (Figure 1). Counter to my predictions, I failed to find strong support for inducible increases in silicon content among the tested plant species. Despite a small p-value in the ANOVA table of the model (p=0.046, F=3.14, df=2,185.9), the model showed only moderate support for an effect of methyl-jasmonate application ( $\beta=0.079\pm0.043, p=0.070$ ), and no support for an effect of cricket exposure on silicon content ( $\beta=0.038\pm0.050, p=0.453$ ). I found a

significant interaction between Species and Induction treatment (p = 0.040, F = 2.52, df = 6, 185.9). Post-hoc tukey tests revealed that this was driven primarily by wheat's response to my induction treatments, where both induction treatments were associated with decreased silicon content (methyl-jasmonate: p = 0.052, t = -1.95, df = 186, Cricket: p = 0.00184, t = -3.16, df = 186) (Figure 2).

#### 380 5.4 Discussion

Recent research in inducible silicon plant defenses has focussed on the short-term dynamics 381 of silicon uptake (Waterman, Hall, et al. 2021; Waterman, Cibils-Stewart, et al. 2021). The 382 promising results of this work has been highlighted for its potential applications in agri-383 culture, where sensitive and rapid defensive phenotypes could improve plant performance 384 and reduce reliance on more intensive pest-control measures. In this study, I attempted 385 to demonstrate rapid silicification in four cereal crops. I failed to find evidence of rapidly 386 induced silicon uptake in response to either methly-jasmonate application or herbivore expo-387 sure. In our study, the variation among our cultivars within species was similar in magnitude 388 to variation among species (Figure 1), possibly obscuring the effects of our treatments. This 389 study also differed from previous studies demonstrating rapid silicon uptake in a number 390 of ways. The two previous studies both grew plants in liquid nutrient solution, carefully 391 standardized to maintain consistently high silicon availability. In my study, I grew plants 392 in potting soil amended with solid silicic acid. In natural soil environments, the majority of plant-available silicon is derived from mineral or biogenic sources, and thus requires dissolution into the soil solution. The theoretical maximum concentration of silicic acid in soil solution is 2 mM, however observed concentrations can be much below that. In my growing 396 conditions, I applied silicic acid in excess of the average availability of phytoliths (a major 397 source of plant-availabe silicon), so as to avoid soil conditions with low silicon presense. De-398 spite this, silicon availability in the soil solution, and dissolution rates from solid to aqueous 399 forms, may have been too low to meet the demands of rapid silicon uptake. 400

# 401 5.5 Acknowledgements

# 5.6 Data Availability

# 5.7 Figures and Tables

Table 1: Credible Interval and Parameter estimates for the hierarchical model. Parameters are estimated against a baseline of Induction: None and Species: Barley.

Parameter	Credible Interval	Parameter Estimate
Insect	-4.2004.100	-4.108
Methyl Jasmonate	-4.204.15	-4.17
Oats	-5.004.95	-4.97
Wheat	-4.604.5	-4.51
Triticale	-4.134.08	-4.11

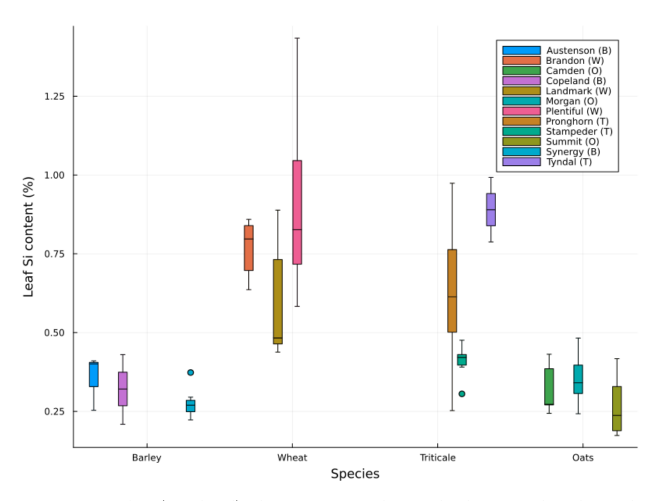


Figure 1: Baseline (uninduced) silicon content in the cereal cultivars used in this study. Cultivar species is notated in parentheses in the legend.

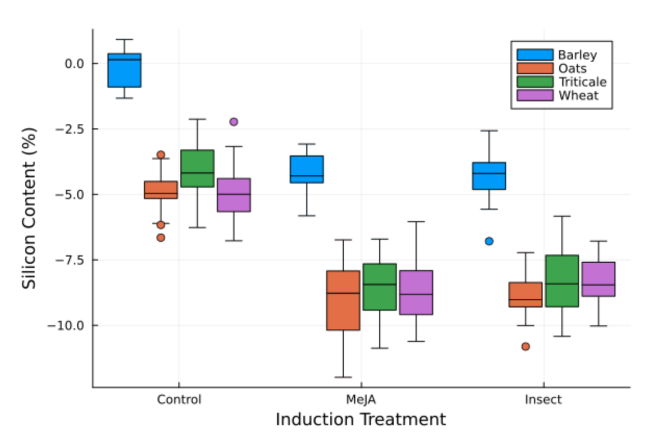


Figure 2: The effects of crop species and induction treatment on leaf silicon content. Plants were treated either with a 1mM methyl jasmonate spray, or exposure to house crickets Acheta domesticus. Leaves were sampled 24 hours after treatment, and were analyzed using XRF.

# 6 Chapter 2: Genetic drivers of silicon accumulation in a wild ancestor of wheat

#### <sub>06</sub> 6.1 Introduction

With a growing global population, and an increasingly imperiled biosphere, the quest for 407 simultaneous increases in both the output and sustainability of agriculture has spurred development and research into new techniques that can help to feed the world and reduce the negative ecological impacts of large scale agricultural production. Over the past thirty 410 years, research momentum has gathered around plant silicon as a potential tool to effect 411 sustainable increases in crop production, with particular applicability in the cereal crops 412 (Reynolds et al. 2016; Christian et al. 2022). Cereal crops are globally important, covering 413 over one-third of the world's arable land, making up over 50% of the daily caloric intake 414 for most people (FAO 2022; Rudel et al. 2009; Awika 2011). Cereals are members of the 415 grass family (Poaceae) and typically have relatively high plant silicon content (0.75% total 416 dry weight) (Reynolds et al. 2016). Silicon is highly abundant in many soils globally, and 417 is the second most abundant element in the earth's crust, behind only oxygen (Ma 2003). 418 It's high expression in cereals, high abundance in many soils, and incredible broad spectrum 419 effects on plant vigor and stress tolerance have make it a tantalizing target for improvements 420 in agricultural yield and sustainability. Though plants can complete their life cycle in the 421 absence of silicon, its influence on such a diverse range of plant physiological functions has 422 caused researchers to emphasize its importance relative to other non-essential nutrients. 423

Silicon underpins a variety of physiological and developmental strategies that plants use to cope with stress. For biotic stressors, silicon can reduce the damage plants experience from herbivory, increase resistance to fungal pathogens, and improve competitive ability with other organisms (Fauteux et al. 2005; Katz 2019). On the abiotic side, silicon supplementation improves plant resistance to soil salinity and heavy metal contamination, improves performance against temperature extremes and high irradiation, and helps plants to cope with drought stress (Cooke and Leishman 2016). In comparing stressed plants grown in the absence or presence of silicon, Si+ plants showed a transcriptome profile similar to unstressed plants (Coskun et al. 2019). A current hypothesis explaining the broad-spectrum activity of

silicon is presented in Coskun et al. (2019), where the authors suggest that silicon deposited in the apoplast of plant tissues where it modulates biological functions of the plant, and ecological interaction with natural enemies, yielding net positive increases in plant performance [I could be more specific if needed]. Realizing these beneficial effects depends on the plant's ability to efficiently source silicon from the soil and uptake it in sufficient amounts. Finding ways to improve crops towards increased silicon use efficiency is key to harnessing the benefits that plant silicon can confer.

Plants gather silicon from the soil solution, using a suite of transporter proteins to pump 440 it into their vascular systems and then transport it throughout the body (Reynolds et al. 441 2016). Variation in the relative expression of these transporters, as well as differences in the development of the end points for silicon deposition (silica cells), may drive phenotypic variation among individuals. Additionally, individuals may vary in their ability to scavenge silicon from the soil. The soluble form of silicon, silicic acid (SiOH4) has a maximum 445 solubility in water of around 2 mM, though typical soil concentrations range from 0.1 mM to 446 0.6 mM (Epstein 1994). Soluble silicon in the soil is derived primarily from the weathering of silicate minerals, and secondarily from the remobilization of silicon in decaying plant material (Félix de Tombeur, Cornelis, et al. 2021). Weathering of silicates releases a host 449 of plant nutrients including Al, Si, Fe, and P (Félix de Tombeur, Cornelis, et al. 2021). 450 Soil biota can drive weathering, using organic acids and other molecules to complex metal 451 ions off of soil aggregates, making them available for uptake by organisms. Plant roots can 452 release carboxylates and phytosiderophores to weather P and Si out of soil minerals. Along 453 with Si and P mobilization, Mn is often released, and taken up by plants roots. Previous 454 research has used leaf Mn content to proxy for the carboxylate releasing activity of plants 455 (Lambers et al. 2015), yet so far I are unaware of any studies looking for quantitative variation 456 among genotypes of leaf Mn. If we could identify regions of the plant genome associated 457 with variation in root weathering activity, we may be able to target this trait in breeding 458 programs that improve nutrient use efficiency, ultimately easing our dependence on external inputs to agricultural fields.

The use of x-ray fluorescence (XRF) to quantify plant silicon has greatly reduced the costs, danger, and processing time of for studies focusing on this topic (Reidinger et al.

2012). XRF works by using low-power x-rays to excite elements in the sample, and measures 463 the resulting emitted light. One of the most exciting features of XRF is the fact that it can 464 analyse multiple elements at once, allowing for broad characterization of the sample for most 465 elements heavier than aluminum. Though XRF is an established technique to measure plant 466 Si, its may also be used to measure other metals of interest, including manganese. In this 467 study I use XRF to quantify variation in Si and Mn content among a diversity panel of a 468 wild ancestor of bread wheat, Aegilops tauschii. This panel has publicly available sequence 460 data, allowing us to perform a genome-wide association sutdy to link Si and Mn variation 470 to genotypic variation, laying the groundwork for future, more targetted, explorations of the 471 genome to identify genetic controls over these traits, and hopefully develop breeding targets 472 to improve plant performance and safeguard yields against a destabilizing climate.

#### 474 6.2 Methods

#### 475 6.2.1 Plant growing conditions

For this experiment, I used a the L2 panel of Aegilops tauschii from (Gaurav et al. 2021) 476 grown at three different sites. Two of the sites were outdoors on the University of British 477 Columbia campus, with planting occurring in the fall, while the third site was a glasshouse, 478 where I vernalized seedlings in growth chambers prior to transplanting into the glasshouse 479 environment. For full site details see Supplementary Table S1. Using 151 accessions, I 480 started trays of seedlings in glasshouse or growth chamber environments. At approximately 481 eight weeks after germination, seedlings were transplanted to their field sites. For each en-482 vironment, I started four replicates of each accession. I planted the plants in a randomized 483 block design, to minimize the effects of soil heterogeneity on our phenotype measurements. Each outdoor block was a 16 m<sup>2</sup> square, with plants arranged ~35 cm apart. Shortly after transplanting to the field sites, I applied water-soluble fertilizer to improve transplant survival, as well as slow-release fertilizer pellets. Field transplantation took place on the 15th 487 of October 2022 and the 16th of December 2022. For the glasshouse environment, I started 488 seedlings in growth chambers in January 2022. After 12 weeks, I moved the seedlings to ver-489 nalization chambers (4°C, 8:16h light:dark) for eight weeks. I then transplanted these plants 490 into 10cm square pots filled with SunGro potting mix and amended with [amount] of silicic acid (Tixosil 68B, Solvay). Pots were arranged using the same randomized block design,
adapted to fit on two flood tables. To ensure a comparable life stage accross environments
at time of harvest, these plants grew for three months (mid June – mid September 2022),
until they had mature flower heads.

#### 496 6.2.2 Plant harvest and sample preparation

When the plants had reached maturity, I harvested the entire above-ground portion of each plant. For the outdoor sites, harvest occurred between the 1st and 5th of July 2022, while I harvested the glasshouse plants between the 19th and 21st of September 2022. I placed harvested material in labelled paper bags, and dried it in drying ovens at 60°C for 48 hours. 500 To harvest leaf material for analysis, I selected stems with flower heads, and removed the 501 three leaves closest to the flowers. Since portions of the plant body have different silicon 502 contents (Dai et al. 2005), I chose a consistent set of leaves to minimize introduced variation. 503 I picked leaves until approximately 200mg of dry leaf was collected. I then washed leaves in 504 distilled water to remove any soil residues which might introduce silicon, and re-dried the 505 samples at 60°C for 48 hours. Dried, clean leaves were then packed into 2 ml microcentrifuge 506 tubes with zircon grinding pellets, flash frozen by immersion in liquid nitrogen, and ground 507 in a tissuelyser ball mill for 30 seconds at 60 Hz. The resulting leaf powder was stored sealed 508 until XRF analysis. 500

#### 510 6.2.3 Sample analysis

To analyse the silicon and manganese content of the accessions, I followed the XRF procedure presented in Reidinger et al. (2012). In short, I pressed leaf powder into 13mm diameter pellets at 11 tonnes of pressure and analysed the resulting pellets in an Olympus Vanta p-XRF device mounted in a bench stand. I used a read time of 30s to ensure accurate measurements. To minimize cross-contamination between samples, I wiped clean the pellet press and XRF device after each sample.

#### 517 6.2.4 Statistical analysis

To perfrom the GWAS analysis, I followed the methodology and code published in Gaurav 518 et al. Nov. 1, 2021. For brevity, this methodology only describes the steps I took using the 519 data generated from Gaurav et al. Nov. 1, 2021. For full details on how they generated the 520 sequence data and prepared the final data sets refer to their manuscript. Prior to the linear 521 regression to assess genetic associations. I first tested my phenotypic data for block effects 522 within sites. For each site, we tested a linear model to see if measured leaf silicon content 523 was significantly associated with block. I tested the model  $Si \sim Block + Accession$ . When 524 I validated that block had a non-significant effect (p > 0.1), I reparameterized leaf silicon 525 measurements at the site level; I converted the raw %Si to be expressed as the number of 526 standard deviations the sample deviated from the site mean of %Si. I did this to eliminate a possible confounding effect of inter-site differences that might reduce the power of the GWAS linear regression. 529

As per Gaurav et al. Nov. 1, 2021, to reduce the computational intensity of my analysis,
I prefiltered the total k-mer matrix to remove k-mers with a low chance of being informative
[word choice?].

#### 533 6.3 Results

Of the approximately 1700 plants planted, [1300] produced enough leaf material for analysis. 534 Silicon content in Aegilops tuaschii ranged from [value]% to [value%]. Site accounted for 535 a large amount of variation in silicon content, suggesting a strong environmental effect on 536 the silicon phenotype. Overall, my analysis revealed [four] regions of the Aegilops tauschii genome that has significant associations with silicon content (Figure 3). One of these genomes 538 was on chromosome 4S, near a known gene analogue to Lsi1, a silicon transporter protein. 539 My results for manganese content are less clear. I detected no regions that met the threshold for significance, though there were three that had pronounced peaks relative to the average 541 response (Figure 4). Within the plants, silicon and manganese content were correlated ( $R^2$ 542 = 0.15, p = 0.049) (Figure 3). 543

- 6.4 Discussion
- 6.5 Acknowledgements
- 546 6.6 Data Availability

# <sup>547</sup> 6.7 Tables and Figures

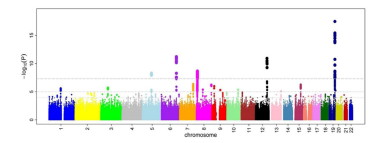


Figure 3: This is an example Manhattan Plot from the GWAS output. The real figure will show associations with silicon content

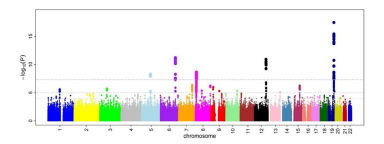


Figure 4: This is another Manhattan Plot, this time showing associations with manganese content

# Reported happiness as a function of income

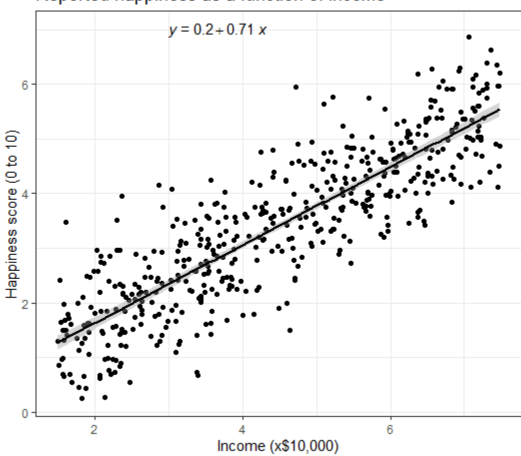


Figure 5: This is the regression comparing Si to Mn content in the leaf tissue

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