1	Short title: Silencing of <i>TaCKX1</i> and grain yield
2	Corresponding author: Anna Nadolska-Orczyk. E-mail: a.orczyk@ihar.edu.pl
3	
4	Title: Silencing of TaCKX1 mediates expression of other TaCKX genes to
5	increase grain yield in wheat
6	
7	Author names and affiliations:
8	Bartosz Jabłoński ¹ , Hanna Ogonowska ¹ , Karolina Szala ¹ , Andrzej Bajguz ³ , Wacław Orczyk ²
9	and Anna Nadolska-Orczyk ^{1c} (corresponding author: a.orczyk@ihar.edu.pl)
10	
11	¹ Department of Functional Genomics, ² Department of Genetic Engineering, Plant Breeding
12	and Acclimatization Institute - National Research Institute, Radzikow, 05-870 Blonie, Poland
13	³ Laboratory of Plant Biochemistry, Institute of Biology, University of Bialystok,
14	Ciolkowskiego 1J, 15-245 Bialystok, Poland
15	
16	ORCID IDs:
17	A.N-O.: 0000-0001-6127-3860
18	B.J.: 0000-0002-3386-5049
19	H.O.: 0000-0001-7385-1592
20	K.S.: 0000-0002-2951-8634
21	A.B.: 0000-0003-4275-0881
22	W.O.: 0000-0003-4735-0000
23	
24	One-sentence summary: Different levels of <i>TaCKX1</i> silencing influence various models of
25	coordinated expression of TaCKX genes and phytohormone levels in 7 DAP spikes, as well as
26	yield parameters.
27	
28	Author contributions: A.N-O. conceived and designed the original research plan and wrote
29	the manuscript; B.J. performed most of the experimental work; H.O. prepared the figures and
30	take part in data analysis; K.S. designed primers and performed comparative analysis with

31 databases, took part in selected experiments; A.B. performed measurements of

- 32 phytohormones; WO took part in planning and discussing results and helped with performing
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38 Abstract

- 39 TaCKX family genes influence development of wheat plants by specific regulation of
- 40 cytokinin content in different organs. However, their detailed role is not known. The
- 41 *TaCKX1*, highly and specifically expressed in developing spikes and in seedling roots, was
- 42 silenced by RNAi-mediated gene silencing via *Agrobacterium* and the effect of silencing was
- 43 investigated in 7 DAP spikes of T_1 and T_2 generations. Various levels of *TaCKX1* silencing in
- 44 both generations influence different models of co-expression with other *TaCKX* genes and
- 45 parameters of yield-related traits. Only a high level of silencing in T₂ resulted in strong down-
- 46 regulation of *TaCKX11 (3)*, up-regulation of *TaCKX2.1, 2.2, 5* and *9 (10)*, and a high yielding
- 47 phenotype. This phenotype is characterised by higher spike number, grain number and grain
- 48 yield, as well as slightly higher mass of seedling roots, but lower thousand grain weight
- 49 (TGW) and slightly lower spike length. Content of most of cytokinin forms in 7 DAP spikes
- 50 of silenced T_2 lines increased from 40 to 76% compared to the non-silenced control. The CKs
- 51 cross talk with other phytohormones.
- 52 Each of the tested yield-related traits is regulated by various up- or down-regulated TaCKX
- 53 genes and phytohormones. Unexpectedly, increased expression of *TaCKX2.1* in silent for
- 54 *TaCKX1* T₂ plants up-regulated trans- and cis-zeatin and trans-zeatin glucosides, determining
- 55 lower TGW and chlorophyll content in flag leaves but higher grain yield. The coordinated
- 56 effect of *TaCKX1* silencing on expression of other *TaCKX* genes, phytohormone levels in 7
- 57 DAP spikes and yield-related traits in silenced T_2 lines is presented.

58 Introduction

59 Wheat (*Triticum aestivum* L.) is the third economically most important crop in the world after 60 corn and rice and probably the most important in moderate climates. It provides

61 approximately 20% of human calories and protein (Reynolds and Braun, 2019). The large

62 genome of this high-yielding species, composed of three (AABBDD) genomes, has been very

63 challenging for improving traits (Borrill et al., 2019). However, it might be a great reservoir

64 for further increase of grain productivity (Nadolska-Orczyk et al., 2017). Continuous increase

- of wheat production is necessary to feed the rapidly growing world population (Foley et al.,
- 66 2011) http://iwyp.org/). Biotechnological tools implemented in the process of increasing
- 67 wheat productivity are expected to be beneficial.
- 68 Cytokinins (CKs) are important regulators of plant growth and development, influencing

69 many agriculturally important processes (reviewed in Kieber and Schaller, 2018). Most of the

- 70 CKs positively regulate cell division and organization of shoot stem cell centres as well as
- 71 stimulating the endocycle in roots by auxin-dependent or auxin-independent mechanisms

72 (Schaller et al., 2014). CKs are also involved in regulation of various developmental and

73 physiological processes including size and structure during leaf development (Skalak et al.,

- 2019), delay of senescence (Gan and Amasino, 1997; Lara et al., 2004), apical dominance
- 75 (Tanaka et al., 2006), root proliferation (Werner et al., 2001; Werner et al., 2003). This
- regulation might occur at the posttranscriptional and/or posttranslational level (Cerny et al.,

2011; Kim et al., 2012) or by modulation of context-dependent chromatin accessibility (Potter

et al., 2018). CKs modulate expression of other genes involved in the control of various

79 processes including meristem activity, hormonal cross talk, nutrient acquisition, and various

80 stress responses (Brenner et al., 2012). There is growing evidence on their key role in seed

81 yield regulation (reviewed by Jameson and Song, 2016). In cereals and grasses an increased

82 content of CKs has been reported to positively affect sink potential in developing grains (Liu

et al., 2013), maintain leaf chlorophyll status during plant senescence (Zhang et al., 2016) and

84 grain filling (Panda et al., 2018). Moreover, in wheat the CKs take part in regulation of seed

85 dormancy (Chitnis et al., 2014).

- 86 The majority of naturally occurring CKs in plants belong to isoprenoid cytokinins grouping
- N^{6} -(12-isopentenyl) adenine (iP), *trans*-zeatin (tZ), *cis*-zeatin (cZ), and dihydrozeatin (DZ)
- 88 derived from tRNA degradation or from isopentenylation of free adenine nucleosides
- 89 catalysed by isopentenyltransferase (IPT) or tRNA-IPT. The second, smaller group comprise
- 90 N6-aromatic CKs, represented by benzyladenine (BA) (Sakakibara, 2006). To better

- 91 characterize their physiological role, CKs are classified into free-base active forms as tZ, cZ
- 92 and iP, translocation forms (nucleosides) as tZ-ribosides (tZR), which exhibit a low level of
- 93 activity, and sugar conjugates (O-glucosides), which are storage and inactivated forms
- 94 (Sakakibara, 2006; Bajguz and Piotrowska, 2009).

95 CKs function as local or long-distance regulatory signals, but the mechanisms of their precise

- 96 spatial and temporal control are still largely unknown (Brandizzi, 2019). They are produced in
- 97 roots as well as in various sites of aerial part of plants (Kudo et al., 2010). The level of CKs in
- 98 respective cells and tissues is dependent on many processes including biosynthesis,
- 99 metabolism, activation, transport, and signal transduction. Active CKs can be metabolized via
- 100 oxidation by cytokinin oxidase/dehydrogenase (CKX) or by activity of glycosyltransferases.
- 101 Many reports have demonstrated that the irreversible degradation step by CKX enzyme plays
- 102 an important role in regulation of cytokinin level in some cereals: maize (Brugie` re N, 2003),
- 103 rice (Ashikari et al., 2005), barley (Zalewski et al., 2010; Zalewski et al., 2014) and wheat
- 104 (Song et al., 2012; Zhang et al., 2012; Ogonowska et al., 2019).
- 105 The level of CKs might be also regulated by transport through the plant. tZ-type CKs are
- 106 mainly synthesized in roots and transported apoplastically to shoots, which promote the
- 107 growth of the above-ground parts of the plant (Beveridge et al., 1997; Hirose et al., 2008). In
- 108 contrast, the iP and cZ type CKs are the major forms found in phloem and are translocated
- 109 from shoots to roots (Corbesier et al., 2003; Hirose et al., 2008).
- 110 The *CKX* gene families in plants show different numbers of genes and various expression
- 111 patterns, which are tissue- and organ-specific, suggesting gene-specific functions. Specificity
- 112 of expression of 11 *TaCKX* in developing wheat plants were assigned to four groups: highly
- specific to leaves, specific to developing spikes and inflorescences, highly specific to roots
- and expressed in various levels through all the organs tested (Ogonowska et al., 2019). The
- 115 *TaCKX* genes co-operated inside and among organs. Their role in plant productivity has been
- 116 described in many plants including model plants and some cereals. Knock-out mutation or
- silencing by RNAi of OsCKX2 in rice significantly increased grain number (Ashikari et al.,
- 118 2005). The same effect of elevated grain number, spike number and yield was reported for
- 119 RNAi-silenced HvCKX1 in barley (Zalewski et al., 2010; Zalewski et al., 2012; Zalewski et
- 120 al., 2014) and repeated for the same gene under field conditions (Holubova et al., 2018).
- 121 Moreover, significantly increased grain number per spike was found as the effect of the
- 122 *TaCKX2.4* gene silenced by RNAi (Li et al., 2018). Knock-out mutation of *HvCKX1* by
- 123 CRISPR/Cas9 editing had a limited effect on yield productivity (Gasparis et al., 2019) or no

vield data were supplied (Holubova et al., 2018). However, the results were contradictory for 124 125 root growth. In the first report, by Gasparis et al. (2019), knock-out ckx1 mutants showed 126 significantly decreased CKX enzyme activity in young spikes and 10-day old roots, which 127 corresponded to greater root length, increased surface area, and greater numbers of root hairs. 128 In the second report, by Holubova et al. (2018), root growth in *Hvckx1* mutant plants was 129 reduced, which was similar to knock-out mutants of ckx3 showing smaller roots (Gasparis et 130 al., 2019). The role of other TaCKX genes in wheat was analysed based on natural TaCKX 131 variation. Haplotype variants of *TaCKX6a02* and *TaCKX6-D1* were related to higher filling 132 rate and grain size (Zhang et al., 2012; Lu et al., 2015). QTL found in recombinant inbred 133 lines containing a higher copy number of *TaCKX4* was associated with higher chlorophyll 134 content and grain size (Chang et al., 2015). To arrange the numbering of *TaCKX* family 135 genes, a new annotation for the first two was suggested by Ogonowska et al. (2019) based on 136 the Ensembl Plants database (Kersey et al., 2018) and phylogenetic analysis. *TaCKX6a02* was 137 annotated as TaCKX2.1, TaCKX6-D1 (JQ797673) was annotated as TaCKX2.2 and TaCKX2.4 138 was annotated as *TaCKX2.2*. Annotations for these genes were maintained in the recently 139 published review on the TaCKX (Chen et al. 2019), however tested in this research TaCKX10 140 was renamed as TaCKX9 and TaCKX3 was renamed as TaCKX11. Newly revised by Chen et 141 al. (2019) naming is applied and former names are in brackets. 142 Due to the size and complexity of the wheat genomes the knowledge about the role of *TaCKX* 143 genes, containing homoeologues from three genomes, is more difficult to obtain, because of

144 the limited number of natural mutants. Most homoeologues genes are expected to have

145 overlapping functions (Uauy, 2006) therefore the effect of gene mutations might be masked

146 by the other genomes. On average, ~70% of wheat homoeologue triads/triplets (composed of

147 A, B, and D genome copies) showed balanced expression patterns/the same co-expression

148 modules (Ramirez-Gonzalez et al., 2018; Takahagi et al., 2018). One solution to silence all of

them is to apply RNAi-mediated gene silencing, which allowed silencing all of the

150 homoeologues. Moreover, this tool made it possible to obtain a number of lines with different

151 levels of silencing, which in the case of genes coding proteins of key importance for life

152 (when lack of protein was correlated with lethality) gave a possibility to regenerate plants for

analysis (Travella et al., 2006). Introduction of a silencing cassette by stable transformation

results in a stable, and inherited to T₄, effect of silencing (Gasparis et al., 2011; Zalewski et

al., 2014). The applicability of *Agrobacterium*-mediated transformation compared to a

156 biolistic one for gene silencing of the developmentally regulated gene *HvCKX10 (2)* was

157 proved to be reliable (Zalewski et al., 2012).

- 158 We present the first report on the role of *TaCKX1* in co-regulation of expression of other
- 159 TaCKX genes, phytohormone content and their joint participation in regulation of yield-
- 160 related traits in wheat. *TaCKX1*, silenced by a hpRNA type of vector via *Agrobacterium*,
- 161 showed different levels of silencing in T_1 and T_2 , which has been related to different models
- 162 of gene action. The coordinated effect of *TaCKX1* silencing in 7 DAP spikes of T_2 plants on
- 163 expression of other *TaCKX* genes, phytohormone levels as well as phenotype based on ratio
- 164 indicators is presented. Moreover, models of regulation of phytohormone levels and
- 165 phenotypic traits by coordinated expression of *TaCKX* genes based on correlation coefficients
- 166 (cc) in non-silenced and silenced wheat plants are proposed.
- 167

168 **Results**

169 Expression levels of silenced *TaCKX1* in segregating T₁ and T₂ plants.

- 170 Expression levels of *TaCKX1* were measured in 44 segregating T₁ plants from 8 T₀ PCR+
- 171 lines. In 14 T_1 plants relative expression (related to the control = 1.00) ranged from 0.39 to
- 172 0.88 with the mean of 0.67 (\pm 0.14). In 30 T₁ plants relative expression ranged from 0.90 to
- 173 1.52 with the mean of $1.16 (\pm 0.18)$ (Fig. 1 A). The proportion of silenced to non-silenced
- 174 plants changed in the T₂ generation. There were 42 silenced from 0.24 to 0.88 plants with the
- mean of $0.54 (\pm 0.14)$ and 20 non-silenced plants. Eight of them, with low relative expression
- 176 ranging from 0.24 to 0.40 (mean 0.33 \pm 0.14), and representing different T₁ lines, were
- 177 selected for further analysis.

178 **Co-expression of silenced** *TaCKX1* with other *TaCKX* genes in T_1 and T_2 and CKX

- 179 enzyme activity.
- 180 Mean relative expression of *TaCKX1* in the selected 8 lines was 0.67 in T_1 and was decreased
- to 0.33 in T_2 (Fig. 2). Similarly, in the case of *TaCKX11* (3) related gene expression was 0.81
- 182 in T_1 and was decreased to 0.34 in T_2 . Relative expression levels of *TaCKX2.2* and *TaCKX9*
- 183 (10) were decreased in T_1 to 0.51 and 0.39 and increased in T_2 slightly above the control
- 184 level, to 1.08 and to 1.10 respectively. Mean relative values for *TaCKX2.1* were similar to
- 185 control in T_1 (1.05) and slightly increased in T_2 (1.17). Relative expression of *TaCKX5*, which
- 186 was in T_1 below the control level (0.84), was significantly increased to 1.82 in T_2 . The relative
- 187 values of CKX enzyme activity in both generations were around the control, 1.00.
- 188 The effect of *TaCKX1* silencing on the levels of expression of selected *TaCKX* genes is
- 189 presented by the expression ratio indicator (Table 1), which is a quotient of the mean relative
- 190 value in silent per mean relative value in non-silent, control plants. In the case of *TaCKX1* and

- 191 TaCKX11 (3), the ratio indicator, significantly decreased in T₁, was strongly decreased in T₂.
- 192 The value of the ratio indicator for *TaCKX2.2* was not changed in T_1 compared to the control
- and was only slightly decreased in T_2 . The expression ratio indicator of *TaCKX9* (10),
- 194 strongly decreased to 0.59 in T_1 , rose above the control level (1.15) in T_2 . Already high in T_1 ,
- 195 the expression ratio indicator for *TaCKX2.1* (1.22) increased to 1.32 in T_2 .
- 196 Table 1. Effect of *TaCKX1* silencing on expression levels of selected *TaCKX* genes presented
- 197 by expression ratio indicator (mean value in silent/mean value in non-silent, control plants) in
- 198 T_1 and T_2 generations.

Expression ratio	T ₁ (SD)	T ₂ (SD)	Effect of <i>TaCKX1</i> silencing
indicator			T ₁ / T ₂
TaCKX1*	0.58 (0.12)	0.28 (0.05)	decreased / strongly decreased
ТаСКХ11 (3)	0.80	0.36	decreased / strongly decreased
TaCKX2.2	1.08	0.98	slightly increased / similar
ТаСКХ9 (10)	0.59	1.15	strongly decreased / slightly increased
TaCKX2.1	1.22	1.32	increased / increased
TaCKX5	1.00	1.08	the same / similar
CKX activity	1.01	0.99	the same / the same

199

200 Phenotype ratio indicator (mean value in silent/mean value in non-silent, control plants) for

201 CKX enzyme activity was 1.01 in T_1 and 0.99 in T_2 . Enzyme activity was not affected by the

202 level of *TaCKX1* expression/silencing.

203 In T₁ segregating plants CKX enzyme activity significantly correlated with spike length (0.51;

n=16) and grain weight (0.50; n=16), but in T₂ these correlations were not significant.

205 Influence of *TaCKX1* silencing on phenotypic traits and chlorophyll content in flag

206 leaves of T_1 and T_2 plants.

207 The values of phenotypic traits in T₁ plants with slightly decreased relative expression of

208 *TaCKX1* (0.67 \pm 0.14) compared to control plants (1.00) were on the same level in the case of

209 plant height and lower for number of spikes, spike length, grain number, and grain yield

- 210 (Table S2). Higher values were obtained for TGW. Data for chlorophyll content measured by
- 211 SPAD in the first spike and the next spikes were similar. All these differences were not
- 212 significant. Opposite results were obtained for some traits in T₂ plants with highly silent
- 213 *TaCKX1* (0.33 \pm 0.06) compared to the control (1.00) (Table S3). Silent T₂ plants were
- substantially smaller, had a higher number of spikes, number of grains, grain yield, seedling

215 root weight and SPAD values for the next spikes. TGW and spike length were significantly

216 lower than in control plants.

217 These differences between the slightly silenced T_1 and highly silent T_2 generation are

218 expressed by comparison of ratio indicators of phenotypic traits in both generations (Fig. 3).

219 There were no changes in plant height (cm), TGW or spike length in T_1 plants compared to

220 the control; however, these values were respectively 7%, 10% and 25% lower in T_2 plants.

221 Opposite phenotype ratio indicators for number of spikes per plant and number of grains per

222 plant were about 21% and 30% lower in T_1 and 57% and 29% higher in T_2 . These differences

223 for spike number, grain number and TGW were significant.

224 The levels of expression of *TaCKX1* in 7 DAP spikes of all T₁ significantly correlated with

number of grains, grain weight, spike length and spike number (0.47, 0.39, 0.42 and 0.33

respectively; n=42) and grain weight correlated with enzyme activity (0.33; n=42). The

227 *TaCKX9 (10)* expression level significantly correlated with grain number (0.51; n=16).

228 Correlation coefficients among expression of all tested *TaCKX* genes and enzyme activity,

and phenotypic traits in non-silent and highly silent T_2 are included in Table S4 A, B. All

these correlations are graphically presented in Fig. 6 A-H and described in section 6 of

results.

232 Phytohormone content in 7 DAP spikes of T₂.

tZOGs, which were mainly composed of tZ-O-glucoside (tZOG) and tZ-9-glucoside-O-

glucoside (tZ9GOG), were the most abundant cytokinin group in 7 DAP spikes (Fig. 4 A).

Their mean content in control plants was 861.18 and 452.43 ng/g biomass and in silent T₂ was

236 953.86 and 974.1 ng/g biomass respectively. The second most abundant was tZ with the level

of 119 ng/g biomass in the control and about twice as high in silent T_2 (230). The next was

238 *cis*-zeatin *O*-glucoside cZOG, which was more abundant in the control than the groups of

silent plants, and the content was 63.48 and 28.68 ng/g biomass respectively. Opposite to

cZOG but similar to tZ, cZ was less abundant in control (29.03 ng/g biomass) but more than

241 three times more abundant in silent (51.03 ng/g biomass) T_2 plants. Concentration of DZGs

242 (sum of DZ7G, DZOG and DZOGR) was higher in silent (29 ng/g biomass) than in control

243 plants (21 ng/g biomass). Very low concentrations (below 0.5 ng/g biomass) were measured

for iP and BA. Concentration of IAA was comparable in control and in silent plants (14.26

and 14.88 ng/g biomass respectively). In the case of ABA, the low concentration in the

control was slightly decreased in silent plants (2.61 and 2.29 ng/g biomass respectively).

247 Concentration of GA was increased from 2.81 ng/g biomass in the control to 29.26 ng/g

biomass in silent plants, which was more than a 10-fold increase.

- 249 Most of the phytohormone ratio indicators in the group of six silent T₂ plants (Fig. 4 B) were
- 250 much higher than in control plants. There were the following cytokinins: tZ (1.53), tZ9GOG
- 251 (2.15), tZOG (1.11), cZ (1.76), sum of DZGs (1.40) and iP (1.32). The ratio indicators for
- some of them were significantly lower, as in the case of BA (0.27) and cZOG (0.45). Similar
- 253 values were observed for IAA, slightly lower for ABA (0.88), but much higher for GA
- 254 (10.42).

255 Coordinated effect of *TaCKX1* silencing on expression of other *TaCKX* genes and 256 phytohormone level in 7 DAP spikes as well as phenotype in T₂.

- 257 A graphic presentation of the coordinated effect of *TaCKX1* silencing on expression of other
- 258 TaCKX genes and phytohormone levels in 7 DAP spikes as well as the phenotype of T₂ plants
- is presented in Fig. 5. The significant decrease of expression of *TaCKX1* was coordinated
- 260 with the significant decrease of *TaCKX11 (3)*, which presumably resulted in a significant
- 261 increase of most CKs: tZ, tZGs, cZ, DZGs, iP as well as GA. The increased phytohormone
- 262 level in the first 7 DAP spike positively influenced traits such as spike number and grain
- number, reaching the ratio indicators 1.57 and 1.29, respectively, and negatively TGW (0.78),
- spike length (0.86), plant height (0.93) and flag leaf senescence (0.95). Opposite data were
- 265 obtained for TaCKX2.1 and TaCKX9 (10), which showed increased expression in silenced, 7
- 266 DAP spikes (1.32 and 1.15 respectively). This might have influenced the decreased ratio
- indicators for phytohormones cZOG (0.45), BA (0.27) and ABA content (0.88) and
- slightly increased ratio indicators for yield-related traits: root weight and grain yield (1.07 and
- 269 1.03 respectively). Expression ratio indicators for *TaCKX5* and *TaCKX2.2* were both close to
- 1.00, but their expression significantly increased compared to T_1 and positively correlated
- 271 with the expression of *TaCKX2.1* and *TaCKX9 (10)* respectively.

272 Models of co-regulation of phytohormone levels and phenotype traits by coordinated

- 273 expression of *TaCKX* genes in non-silenced and silenced T₂ plants
- 274 Two different models of co-regulation of TaCKX expression and phenotypic traits in non-
- silenced and silenced plants of the T₂ generation are proposed (Fig. 6 A-H) based on
- 276 correlation coefficients (Table S4 A, B).
- 277 Plant height (Fig. 6 A). There was no correlation between plant height and expression values
- 278 of any TaCKX expressed in 7 DAP spikes of non-silent as well as silent plants. In the first
- 279 group of plants this trait negatively correlated with BA and positively with IAA and GA

280 content. By contrast, in silent plants the values of plant height were negatively correlated with

growing concentration of tZ and tZGs, which resulted in a smaller plant phenotype.

282 Spike length (Fig. 6 B) in non-silent plants was positively correlated with cZOG, and

283 negatively with ABA content. These correlations determined longer spikes and the trait

284 negatively correlated with spike number and grain number. A strong positive correlation

between CKX activity and spike length was noted in silent plants. The values of enzyme

activity correlated positively with slightly increased *TaCKX5* expression, which negatively

correlated with increasing content of cZ. Spike length in silent plants was positively

correlated with grain yield.

289 TGW (Fig. 6 C). There was no correlation of TGW with expression of any *TaCKX* expressed

290 in 7 DAP spikes of non-silent plants; however, the trait was strongly negatively correlated

291 with tZ and cZ content and positively with GA. The grains in this group of plants were larger

and TGW higher. By contrast, in silent plants there was a strong negative correlation of the

trait with growing expression of *TaCKX2.1*, which positively regulated tZ, tZGs, cZ and iP,

but negatively GA content. Moreover, the values of expression of down-regulated *TaCKX11*

295 (3) negatively correlated with growing content of tZGs and iP and positively with the trait.

296 Altogether it resulted in lower TGW compared to non-silenced plants. The trait in silent

297 plants was strongly and positively correlated with grain yield (0.82) and root weight (0.77).

298 Grain yield (Fig. 6 D). Expression levels of *TaCKX1*, *TaCKX2.2* and *TaCKX5* in non-silent

299 plants positively correlated with iP and negatively with BA content. However, expression of

300 *TaCKX11 (3)* and *TaCKX2.1* negatively regulates tZGs. Altogether it resulted in lower grain

301 yield comparing to silenced plants, and the trait was strongly positively correlated with spike

302 number (0.93) and grain number (0.99). Increasing expression of *TaCKX2.1* positively

303 correlated with growing content of tZGs and cZ and negatively with the trait in silent plants.

304 Decreasing expression of *TaCKX11 (3)*, which was positively correlated with decreased

305 cZOG content and negatively with GA content, positively correlated with the trait. Moreover,

306 a positive correlation was observed between CKX activity and grain yield in this group of

307 plants, which was higher than in non-silent plants. The trait was strongly correlated with

308 TGW (0.82) and root weight (0.66).

309 Spike number (Fig. 6 E) and grain number (Fig. 6 F) in non-silenced plants were positively

310 regulated by *TaCKX1*, *TaCKX2.2* and *TaCKX5*, and their expression was positively correlated

311 with iP and negatively with BA. On the other hand, expression levels of *TaCKX2.1* plus

312 *TaCKX11 (3)* were negatively correlated with the traits as well as with iP and positively with

313 BA. Both groups of genes finally affected lower spike and grain number in non-silent

- 314 comparing to silent plants and were strongly and positively correlated with each other (0.91)
- and grain yield (0.93 and 0.99 respectively). In silent plants decreasing expression of *TaCKX1*
- 316 is negatively correlated with both spike and grain number and the gene negatively regulates
- 317 decreasing BA content. In the case of grain number, the main player positively correlated with
- 318 the trait is *TaCKX5*, increased expression of which was correlated with slightly higher IAA
- 319 content, which resulted in higher grain number. Spike number is also positively regulated by
- 320 *TaCKX5* co-expressed with *TaCKX2.1*, and both genes were positively correlated with
- 321 growing CKs, DZGs and iP as well as GA, determining higher spike number. Both traits are
- 322 highly correlated (0.88) with each other.
- 323 Seedling root weight (Fig. 6 G). There was strong, positive correlation between *TaCKX9* (10)
- 324 expression in 7 DAP spikes and seedling root weight in non-silenced plants, although the
- 325 level of expression of the gene did not correlate with any phytohormone content. Moreover,
- 326 CKX activity negatively correlated with the trait, which finally resulted in lower root weight.
- 327 Decreasing expression of *TaCKX11 (3)* in the case of silent plants was positively correlated
- 328 with decreasing content of cZOG and strongly positively correlated with the trait. Increasing
- 329 expression levels of *TaCKX9* (10) plus *TaCKX2.2* negatively correlated with decreasing
- 330 content of cZOG and root weight.
- 331 Chlorophyll content measured by SPAD in flag leaves of first spikes (Fig. 6 H). There was no
- 332 correlation between expression level of any *TaCKX* measured in 7 DAP spikes of non-silent
- 333 plants and the trait. The only correlations were between phytohormone content and the trait,
- 334 positive for tZ and cZ and negative for GA, which resulted in higher SPAD values
- 335 (chlorophyll content). Increasing expression of *TaCKX2.1* was strongly positively correlated
- with growing values of tZ, tZGs, cZ and DZGs as well as GA in silent plants. A strong
- negative correlation was observed between the gene expression and chlorophyll content,
- 338 which means that increasing expression of *TaCKX2.1* in 7 DAP spikes results in lower
- 339 chlorophyll content in silent plants.
- 340

341 Discussion

342 First, 7 DAP spike was chosen as a research objective in wheat since decreased *HvCKX1*

343 expression at this stage in barley resulted in higher yield due to the higher spike and grain

- number (Zalewski et al., 2010; Zalewski et al., 2014). The samples were taken from the
- 345 middle part of the spikes, when anthesis starts, in order to ensure a similar developmental

- 346 stage of spikelets for research. The 7 DAP spikes of wheat represent the middle of cell
- 347 division/cell expansion stage (Gao et al., 1992; Hess et al., 2002). The 7-day old embryo is in
- 348 the early stage of development, beginning to change from globular to torpedo shape, with the
- root pole starting to differentiate and endosperm starting to degenerate (http://bio-
- 350 gromit.bio.bris.ac.uk/cerealgenomics/cgi-bin/grain3.pl).
- 351 The *TaCKX1* gene is an orthologue of *HvCKX1* and both genes are specifically expressed in
- developing spikes (Ogonowska et al., 2019). In our earlier research with barley we
- 353 hypothesized and continue here to address the hypothesis that the level and pattern of
- 354 expression of a defined *CKX* family gene might determine the specific phenotype and indicate
- 355 its function (Zalewski et al., 2014).

356 Various levels of *TaCKX1* silencing influence different models of co-expression with

357 other *TaCKX* genes and parameters of yield-related traits

- 358 One third of segregating T_1 plants showed a decreased expression level of *TaCKX1* between
- 12% and 61%. Further selection of T₂ led to obtaining plants with much higher, exceeding
- 360 60%, silencing of the gene. Different levels of silencing of TaCKXI in T_1 and T_2 generate
- 361 various results of co-expression with other *TaCKX* genes and plant phenotype. Slightly
- decreased expression of TaCKX1 in T₁ was correlated with slightly decreased expression of
- 363 *TaCKX11 (3)* and *TaCKX5* and significantly decreased *TaCKX2.2* and *TaCKX9 (10)*. Highly
- decreased expression of TaCKXI in T₂ was correlated with highly decreased expression of
- 365 *TaCKX11 (3)* and highly increased *TaCKX5*, *TaCKX2.2* and *TaCKX9 (10)*, and expression of
- 366 *TaCKX2.1* was on a similar level to the control in T_1 and slightly increased in T_2 . Expression
- of TaCKX9 (10) was highly and significantly correlated with TaCKX1 only in T₁. However, a
- new and strong positive correlation between *TaCKX9* (10) and *TaCKX2.2* in highly silenced
- 369 T_2 was observed. Slightly decreased co-expression of silenced *TaCKX1* together with
- 370 TaCKX11 (3) in T₁ and stronger in T₂ indicate their positive co-regulation. It should be
- 371 underlined that there is no homology between the sequence of *TaCKX1* used for silencing and
- 372 sequences of other *TaCKX* genes tested; therefore the process of RNAi silencing was
- 373 specifically addressed to *TaCKX1* silencing. It indicates that the level of silencing of the
- 374 modified gene affected variable levels of expression of the other *TaCKX* genes in a co-
- 375 operative process maintaining homeostasis of CKX enzyme in the research object.
- 376 Transcriptome analysis of knock-out *HvCKX1* in barley (Gasparis et al., 2019) was associated
- 377 with down- or up-regulation of other *HvCKX* family genes as well. Besides, models of co-

378 regulation of other *CKX* by highly silenced *TaCKX1* and knock-out *HvCKX1* differ between

these species.

- 380 The differences in the levels of expression of *TaCKX1* and various co-expression of other
- 381 *TaCKX* genes in T_1 and T_2 resulted in opposite phenotypic effects. Since spike number, grain
- number and grain yield were reduced in T_1 , representing slightly decreased *TaCKX1*
- 383 expression, the same yield-related traits were significantly higher in highly silenced T₂ plants.
- 384 High-yielding phenotype occurred when highly silenced *TaCKX1* co-operated with down-
- regulated *TaCKX11 (3)* but up-regulated *TaCKX5*, *TaCKX2.2*, *TaCKX2.1* and *TaCKX9 (10)*.
- 386 These differences showed that both levels of silencing might be helpful to better understand
- 387 the function of developmentally regulated genes. Unexpectedly, changes in expression levels
- 388 of co-working *TaCKX* did not result in different enzyme activity, even in highly silenced T_2
- 389 plants. It might be explained that down-regulation of *TaCKX1* and *TaCKX11(3)* is
- 390 compensated by up-regulation of *TaCKX2.2*, *TaCKX5* and *TaCKX9* (10), and therefore the
- 391 contribution of isozymes encoded by the genes in the general pool of CKX enzyme activity is
- 392 the same. Since CKX enzymes indicate different specificities for the particular cytokinin
- 393 hormone(Gajdosova et al., 2011), the cytokinin contribution and phenotypic traits of modified
- 394 plants were changed accordingly, with consequent differences in the active pool of CKs
- influencing phenotype.
- 396 Schaller et al. (Schaller et al., 2014) listed three possible explanations of differing, positive or
- 397 negative regulatory roles of CKs in the process of cell division. The first two are the presence
- 398 of additional specific regulators or other signals, which have also been reported by others
- 399 (Reid et al., 2016; Mao et al., 2019), and the third possibility is the level of cytokinin activity.
- 400 Involvement of additional specific regulators in differentiating the regulatory roles of CKs is
- 401 also plausible in our case. However, the most convincing explanation is changed levels of CK
 402 activity (concentrations), which was proved to be significantly increased in silent plants and is
 403 further discussed below.

404 **Co-operating effect of** *TaCKX* **on the level of active CKs in silenced plants**

- 405 Since *TaCKX* family genes encode CKX isozymes, which specifically degrade CKs, we might
- 406 expect that the consequence of decreased expression of *TaCKX* genes is a decrease of
- 407 isozyme activity in the general pool of CKX enzymes, and increased content of active
- 408 cytokinins in the respective organ. Therefore according to our diagram (Fig. 5), highly
- 409 decreased expression of *TaCKX1* and *TaCKX11* (3) in 7 DAP spikes is expected to result in
- 410 the observed increase of most major, active forms of CKs. Already the highest content of

411 tZGs and tZ in non-silenced plants was increased by 45-53% in silenced plants. The third was 412 cZ, with about $\frac{1}{4}$ of the content of tZ in non-silenced and increased by 76% in silenced plants. 413 Conversely, the higher level of cZOG in non-silenced plants was decreased to 45% in 414 silenced plants. We documented that both tZ and cZ, which are isomers of zeatin, together 415 with their derivatives are a major group of isoprenoid CKs in 7 DAP spikes. It has already 416 been shown that trans-zeatin is the predominant form after anthesis (Morris et al. 1993, Hess 417 et al. 2002), but comprehensive analysis of cytokinins during spike, spikelet, ovule and grain 418 development has not yet been reported for wheat using LC-MS/MS (Chen et al. 2019). 419 The content of DZGs increased by 40% in silent compared to non-silent wheat plants, 420 suggesting that this less known isoprenoid form of CKs might also play an important role in 421 plant productivity. Interestingly, isoprenoid iP was represented in 7-DAP spikes of non-silent 422 plants at very low quantities, but its content in 7 DAP spikes of silent plants was increased by 423 32%. A similar relationship between reduced expression of selected CKX family genes and 424 cytokinin accumulation in reproductive organs has been observed in other species including 425 A. thaliana (Bartrina et al., 2011), rice (Ashikari et al., 2005) and barley (Holubova et al., 426 2018), but detailed data are not comparable to our research in wheat. 427 The physiological significance of these isoprenoid forms is not very well known. In maize 428 plants iP and tZ are the most abundant and are susceptible to CKX enzyme (Bilyeu et al., 429 2001). DZ, which generally occurs in small quantities, is biologically stable and resistant to 430 CKX. cZ was found to be less active and more stable than tZ and iP because of its low affinity 431 to CKX (Bilyeu et al., 2001). Also many other researchers indicate tZ as a bioactive 432 cytokinin, whereas cZ was reported to have a weak biological impact and unknown biological 433 role (Schafer et al., 2015). According to Gajdosova et al. (2011), cis-Z-type CKs are frequent 434 in the developmental stages, where they are associated with limited growth, although the 435 metabolic fate of both cZ and tZ varied between different species. In addition, cZ levels 436 change significantly during development in maize grain, as well as in shoot and root tissues 437 (Saleem et al., 2010; Zalabak et al., 2014). The high levels of cZ at the first developmental 438 stage of barley spike observed by Powell et al. (Powell et al., 2013) might indicate a possible 439 role of this form in early barley embryo development. A significant increase of active cZ was 440 also observed in 7 DAP spikes representing early stages of embryo development in our 441 research and was negatively correlated with some yield-related traits such as higher grain 442 yield and root mass and shorter spike length and lower TGW (discussed further below).

- 443 The BA is represented in 7-DAP spikes of wheat at trace amounts but their content was
- 444 significantly decreased in silent plants. However, their correlations with the *TaCKX* genes as
- 445 well as yield-related traits of non-silenced plants indicate their importance (discussed in more
- 446 detail below). Interestingly, BA was found to participate in posttranscriptional and/or
- 447 posttranslational regulation of protein abundance in *Arabidopsis*, showing high specificity to
- 448 shoots and roots, and affected differential regulation of hormonal homeostasis (Zd'arska et al.,
- 449 2013).

450 Cross talk of CKs with other phytohormones

- 451 Negative correlations between ABA content and *TaCKX2.2* and *TaCKX9* (10) expression, and
- 452 positive with *TaCKX11 (3)*, were associated with a slight decrease of ABA content in 7 DAP

453 spikes of silenced plants. Moreover, ABA was strongly positively correlated with BA. The

454 main auxin, IAA, remained at the same level. A ten-fold increase of GA content in silenced

- 455 comparing to non-silenced plants was observed.
- 456 Such cross regulation of CKs and other plant hormones is documented in other species. In
- 457 maize kernels the *CKX1* gene is up-regulated by cytokinin and ABA, and abiotic stress
- 458 (Brugie` re N, 2003). In tobacco altered cytokinin metabolism affected cytokinin, auxin, and
- 459 ABA contents in leaves and chloroplasts (Polanska et al., 2007), which host the highest
- 460 proportion of CK-regulated proteins (Cerny et al., 2013). Moreover, auxin, ABA and
- 461 cytokinin are involved in the hormonal control of nitrogen acquisition and signalling (Kiba et
- 462 al., 2011), which often limits plant growth and development. All four phytohormones CKs,
- 463 GA, IAA and ABA were found to be involved in regulation of grain development in drought
- 464 conditions (Abid et al., 2017). Moreover, in shoots, BA up-regulated the abundance of
- 465 proteins involved in ABA biosynthesis and the ABA response, whereas in the roots, BA
- 466 strongly up-regulated the majority of proteins in the ethylene biosynthetic pathway (Zd'arska
- 467 et al., 2013). We proved that IAA, GA and ABA contents are also co-regulated by CKs in
- 468 non-silenced and silenced 7 DAP spikes. Up-regulation of major CKs and down-regulation of
- some minor ones in silent plants influence GA, ABA and IAA content in a similar manner as
- 470 in abiotic stress conditions.

471 Coordinated effect of *TaCKX* gene expression on the content of CKs, other

472 phytohormones and yield-related traits

- 473 Plant height in non-silenced plants is down-regulated by BA and up-regulated by IAA and
- 474 GA content in the first 7 DAP spikes, resulting in taller plants. Oppositely, increased content
- 475 of tZ and tZGs negatively correlated with the trait in silent plants, and therefore increase of

these main CKs in developing spikes negatively stimulated plant height. Since expression of
genes was measured in 7 DAP spikes, none of the individual genes was shown to regulate the
trait.

As it was proved by Brenner and Schmulling (Brenner and Schmulling, 2012) and similarly to
our results, plant height and root weight are regulated by CKs and IAA in opposite ways. It

481 may be dependent on basipetal auxin flow in the stem, which suppresses axillary bud

482 outgrowth, and similarly as in pea, auxin derived from a shoot apex suppresses the local level

483 of CKs in the nodal stem through the regulation of *CKX* or *IPT* genes (Shimizu-Sato et al.,

484 2009).

485 The main role in spike length seemed to be played by cZ and its derivatives. Increased content

486 of cZOG in non-silenced plants negatively correlated with ABA, resulting in longer spikes. In

487 silent plants the trait is positively regulated by *TaCKX2.2* together with *TaCKX5*, and the

488 latter is a positive regulator of enzyme activity and negative of cZ content. Consequently

489 higher content of cZ in 7 DAP spikes led to shorter spikes. cZOG found as positive regulators

490 of longer spikes are sugar conjugates of cZ-0-glucoside, which are inactivated forms of cZ,

491 showing metabolic stability against CKX activity (Sakakibara, 2010). The tZ-0-glucosides

492 hyper-accumulates in *IPT* overexpressing plants (Zubko et al., 2002). 0-glucosylation of cZ is

493 catalysed by a specific 0-glucotransferase, cisZOG1, discovered in maize (Martin et al.,

494 2001), and this form mainly functions in the early stages of seed development.

495 Knowledge of function of cZ degradation pathways via the CKX enzyme is limited.

496 Interestingly, two Arabidopsis genes, CKX1 and CKX7, expressed in stages of active growth,

497 were shown to have high preference for cZ (Gajdosova et al., 2011). Accordingly,

498 overexpression of CKX7 highly decreased levels of free cZ in Arabidopsis (Kollmer et al.,

499 2014). In our case relatively low expression of *TaCKX2.2* together with *TaCKX5* led to lower

500 CKX activity and higher cZ content.

501 None of the tested individual *TaCKX* genes was involved in high TGW in non-silenced

502 plants, but a negative correlation with tZ and cZ and positive with GA was found. A

significant negative correlation of *TaCKX2.1* and a positive correlation of *TaCKX11(3)* in

504 determining low TGW were observed in silenced plants. Unexpectedly increased expression

505 of the first one positively influenced tZ, tZGs, cZ and iP content and negatively GA content,

and the opposite was true for the second gene, resulting in lower TGW. Therefore both

507 TaCKX2.1 and TaCKX11 (3), acting in an opposite manner, maintain homeostasis of CKX

508 enzyme activity and co-regulate TGW in silenced plants.

509 Greater concentration of CKs, especially tZ, was observed during the grain filling stage of 510 high-yielding cultivars (Powell et al., 2013). We might suppose that the observed higher 511 concentrations of tZ and other CKs at the 7 DAP stage, which originally was a consequence 512 of *TaCKX1* silencing, might accelerate germination of the grains, which resulted in smaller

513 grains/lower TGW than in non-silenced plants. The silenced *TaCKX1* co-work with down-

regulated *TaCKX11 (3)* in increasing CK content as well as up-regulating *TaCKX2.1*, with

515 seems to play a regulatory role.

516 The involvement of GA in TGW and other traits demonstrated by us might be the effect of

517 co-regulation of *CKX* and other gibberellin-responsive genes. One of them is the dwarfing

518 gene *Rht12*, which can significantly reduce plant height without changing seedling vigour and

519 substantially increase ear fertility in bread wheat (Chen et al., 2018). Similar results were

520 obtained with allelic variants of *Ppd-D1* (chromosome 2D) and *Rht-D1* (chromosome 4D)

521 loci, which affect some plant growth traits, e.g. leaf area and spike length (Guo et al., 2018).

522 Fahy et al. (Fahy et al., 2018) in their research on starch accumulation suggested that final

523 grain weight might be largely determined by developmental processes prior to grain filling.

524 This is in agreement with our observations, in which yield-related traits are differently

525 regulated in two groups of plants, non-silent and silent. Therefore we might suppose that

526 coordinated co-regulation of expression of *TaCKX* genes and related CKs takes place during

527 whole plant and spike development and small seeds in silenced plants are determined at

528 earlier stages.

529 Grain yield, which is very strongly correlated with grain and spike number in non-silent

530 plants but with TGW in silent plants, is a more complex feature. Two groups of genes up-

531 regulating or down-regulating grain yield in non-silent plants have been found. The first one

532 includes *TaCKX1*, 2.2 and 5 positively regulating iP and tZGs content but negatively BA. The

533 second comprises *TaCKX11 (3)* (and 2.1) acting in the opposite way. We might expect that

these genes, which negatively correlate with the trait, are main determinants of lower grain

535 yield, affected by down-regulation of tZGs and BA. It is worth to mention that *TaCKX5* is

536 highly expressed in inflorescences and leaves. Higher grain yield was positively regulated by

537 enzyme activity and both *TaCKX11 (3)* as well as *TaCKX2.1* in silenced plants. Indeed,

538 correlation of *TaCKX2.1* expression with the trait was negative. However, the gene positively

539 regulated tZGs and cZ content just like for TGW, which is rather untypical for a gene

540 encoding a CKX enzyme degrading CKs. Therefore the positive regulation of the main CK

541 content by *TaCKX2.1* observed by us supports its role in regulation of expression of other542 genes rather than encoding the CKX isozyme.

543 As observed in barley cultivars, changes in cytokinin form and concentration in developing 544 kernels correspond with variation in yield (Powell et al., 2013). Interestingly, the authors 545 observed no peaks and no differences in CKX activity at the particular stages of spike 546 development, which is in agreement with homeostasis of the pool of isozymes in 7 DAP 547 spikes of wheat, as suggested by us, which is independent of the level of silencing of TaCKX1 548 but is rather a consequence of co-regulation of expression of other TaCKX genes. A similar 549 effect of increased grain yield, which was a consequence of higher spike and grain number, 550 was obtained in barley with silenced by RNAi HvCKX1, an orthologue of TaCKX1 (Zalewski 551 et al., 2010; Zalewski et al., 2014; Holubova et al., 2018). The three CKs measured, cZ, tZ 552 and iP, were differently regulated in earlier or later, but not precisely defined, stages of spike 553 development (Holubova et al., 2018); therefore the results of CK contents are not comparable. 554 Incomparable to the results obtained for RNAi silenced *TaCKX1* and *HvCKX1*, no changes in 555 yield parameters, including spike and seed number, were observed in mutant lines with 556 knock-out of *HvCKX1* (Gasparis et al., 2019). These essential phenotypic differences between 557 RNAi-silenced TaCKX1 and HvCKX1 or HvCKX1 knocked out by CRISPR-Cas9 might be 558 the result of different processes involved in inactivation of the gene. The first one is regulated 559 at the posttranscriptional and the second at the transcriptional level. Since CKs might regulate 560 various developmental and physiological processes at the posttranscriptional level (Cerny et 561 al., 2011; Kim et al., 2012) or by modulation of context-dependent chromatin accessibility 562 (Potter et al., 2018), the way of deactivating *TaCKX* function seemed to be important. Spike number and grain number are highly correlated in both non-silent and silent plants and 563 564 are regulated by the same groups of *TaCKX* genes as well as phytohormones. The first group 565 includes TaCKX1, 2.2 and 5 positively regulating iP but negatively BA. The second 566 comprises TaCKX11 (3) and 2.1 acting in the opposite way, and homeostasis of these 567 hormones in non-silenced plants maintains a lower spike number. The main role in controlling 568 higher spike and grain number in silent plants seemed to be played by TaCKX2.1 and 5. Their 569 increased expression determine higher spike and grain number. The higher spike number was 570 correlated with higher DZGs, iP and GA content. These correlations are not significant 571 because they were measured in a stage of plant development in which the number of spikes 572 and seed number have already been set.

As reported, the higher spike number was the consequence of a higher tiller number, which was positively correlated with the content of endogenous zeatin in the field-grown wheat after exogenous hormonal application (Cai et al., 2018). Moreover, the authors showed that IAA application inhibited the occurrence of tillers, by changing the ratios of IAA and ABA to zeatin. Shoot branching might also be dependent on acropetal transport of cytokinin

578 (Shimizu-Sato et al., 2009).

579 Root weight was positively correlated with TaCKX9 (10) expression in 7 DAP spikes of non-

580 silent plants and, conversely, negatively regulated by increased expression of this gene in

581 silenced plants. Therefore lower (compared to silent plants) expression of the gene in the

investigated organ determined lower root weight in the first group of plants, but higher in the

583 second. *TaCKX9 (10)*, which is highly and specifically expressed in leaves (Ogonowska et al.

584 2019) and showed increased expression in 7 DAP spikes of silent plants, down-regulated

585 cZOG. The same cZOG was up-regulated by *TaCKX11 (3)*, but expression of this gene in 7

586 DAP spikes of silent plants is strongly decreased. Both cZ and cZOG are involved in spike

587 length regulation as well as TGW and grain yield in the group of silenced plants.

588 Since our results are restricted to the 7 DAP spike but are correlated with weight of seedling

roots, we should take into consideration possible action of cytokinin transport and signalling

590 genes as well as other phytohormones, which take part in hormonal cross-talk to control

regulation of root growth (Pacifici et al., 2015). Accordingly cZ type CKs found as the major

592 forms in phloem and are translocated from shoots to roots (Corbesier et al., 2003; Hirose et

al., 2008). It is also worth underlining that only *TaCKX2.1* and *2.2* are specifically expressed

in developing spikes. Both *TaCKX9 (10)*, highly and specifically expressed in leaves, and
 TaCKX11 (3), expressed in all organs, seemed to regulate seedling roots as well, although in

the opposite manner.

597 The lower plant height and higher root weight observed in the group of silenced plants of 598 wheat is in agreement with opposed regulation of these traits by CKs and IAA mentioned

by above (Brenner and Schmulling 2012). CKs are considered as negative regulators of root

growth, so their reduction causes enhanced root growth (Werner et al., 2010; Mrizova et al.,

601 2013), while excess of CKs inhibits primary root growth in *Arabidopsis* (Riefler et al., 2006;

602 Dello Ioio et al., 2012). It was also demonstrated that long-distance, basipetal transport of

603 cytokinin controls polar auxin transport and maintains the vascular pattern in the root

604 meristem (Bishopp et al., 2011). Therefore CKs might operate differently in distinct parts of

605 the plant. The cZOG, represented mainly by inactive cZ-0-glucoside, down-regulated by

606 *TaCKX9* (10) in 7 DAP spikes, was probably up-regulated in seedling roots, resulting in

- 607 higher root weight. Content of cZ riboside was proved to be most abundant in the roots of
- maize (Veach et al., 2003). Opposite data, up-regulated content of active cZ in 7 DAP spikes,
- 609 might influence down-regulation of this CK in roots. It has been documented that such
- 610 suppressing cZ levels mediated by overexpression of AtCKX7 affected root development in
- 611 *Arabidopsis* (Kollmer et al., 2014). Higher weight of seedling root was also obtained by
- 612 silencing via RNAi or knock-out via CRISPR/Cas9 of HvCKX1 in barley plants, as in wheat,
- and the trait corresponded with decreased activity of CKX enzyme measured in roots
- 614 (Zalewski et al. 2010, Gasparis et al. 2019).
- 615 Some *CKX* genes might be induced by transcription factors (Reid et al., 2016; Mao et al.,
- 616 2019). An important role of some NAC transcription factors in regulation of *TaCKX*
- 617 expression and subsequent organ development has also been documented in our not yet
- 618 published results. Possible mechanisms of co-regulation of all these genes in the developing
- 619 roots of silent and non-silent plants is being further investigated by us.
- 620 Leaf senescence was determined in the flag leaf of the first spike by measuring chlorophyll
- 621 content. The trait was down-regulated by increased *TaCKX2.1* expression in silent plants.
- 622 Opposite to the general role of *TaCKX* action, increased expression of the gene does not
- 623 down-regulate but up-regulates tZ, tZ derivatives and cZ content in 7 DAP spikes. *TaCKX2.1*
- 624 functions in a similar way, by up-regulating tZ, tZ derivatives and cZ in determining lower
- TGW and higher grain yield in silent T₂ plants. Lower content of active CKs in the first spikes
- of non-silent plants is expected to up-regulate these CKs in flag leaves of these spikes,
- 627 maintaining prolonged chlorophyll content. In contrast, higher content of tZ, tZGs, cZ and
- 628 DZGs as well as GA in 7 DAP spikes of silent plants is expected to down-regulate CKs in the
- 629 flag leaves, accelerating their senescence, which is documented by the results.
- 630 It was previously demonstrated that level of chlorophyll content in flag leaves is associated
- 631 with the senescence process, in which CKs suppress inhibition of senescence (Gan and
- Amasino, 1995). During this processes, proteins are degraded and nutrients are re-mobilised
- from senescing leaves especially to the developing grains (Gregersen et al., 2008). We might
- 634 suppose that slower spike ripening in non-silent plants, which is dependent on lower CK
- 635 content in the 7 DAP spike, causes slower flow of micronutrients as well as CKs from flag
- 636 leaf to spike. Therefore prolonged chlorophyll content in the flag leaf of the first spike
- 637 negatively correlated with TGW but positively with plant height. Opposite data were obtained
- 638 for flag leaves of silent plants, in which higher content of CKs in 7 DAP spikes might be the

result of faster flow accelerating leaf senescence. Reduced chlorophyll content in flag leavesof the first spike of silent plants positively correlated with grain yield.

641 The important role of tZ and less active cZ in suppression of senescence was proved in maize

642 leaves (Behr et al., 2012), and in an oat-leaf assay (Gajdosova et al., 2011). It was also

- 643 documented that delayed senescence of wheat stay-green mutant, tasg1, at the late filling
- 644 stage was related to high cytokinin and nitrogen contents (Wang et al., 2019).
- 645

646 CONCLUSION

- 647 Based on the 7 DAP spike as a research object, we have documented that silencing of
- 648 *TaCKX1* by RNAi strongly influenced up- or down-regulation of other *TaCKX* genes, as well
- as phytohormone levels and consequently phenotype. This co-regulation is dependent on the
- 650 level of silencing of the gene and is independent of cross-silencing of other *TaCKX*.
- 651 In the general model of regulation of yield-related traits, highly silenced *TaCKX1*, which is
- 652 specific for developing spikes, strongly down-regulates *TaCKX11 (3)*, expressed through
- different organs of the developing plant. Coordinated action of these genes is expected to lead
- to increased contents of tZ, tZGs, cZ and DZGs as well as GA, determining a high-yielding
- 655 phenotype. In contrast, up-regulated *TaCKX2.1*, which is specifically expressed in developing
- spikes, and *TaCKX9 (10)*, strongly expressed in leaves, might be down-regulators of cZOG.
- 657 However, detailed analysis revealed that each tested yield-related trait is regulated by various
- 658 up- or down-regulated *TaCKX* genes and phytohormones. Key genes involved in regulation of
- 659 grain yield, TGW or root weight in highly silenced plants are *TaCKX2.1* and *TaCKX11 (3)*
- acting antagonistically; increased expression of the first one determines growth of tZ, tZ
- derivatives and cZ, whereas decreased expression of the second down-regulates content of
- 662 cZOG. A key role in determination of the high-yielding phenotype seemed to be played by the
- growing content of tZ in 7 DAP spikes, which might accelerate maturation of immature grains
- by speeding up nutrient flow from flag leaves. This finally led to reduction of TGW but
- 665 enhancement of grain number and yield. The latter traits are the result of a higher spike
- number, which is determined in the early stages of plant development.
- 667
- 668 Materials and methods

669 Vector construction

- 670 The hpRNA type of silencing cassette was constructed in pBract207
- 671 (<u>https://www.jic.ac.uk/technologies/crop-transformation-bract/</u>). It contains the *Hpt* selection

- 672 gene under the 35S promoter and cloning sites for the cloning silencing cassette under the Ubi
- 673 promoter. The vector is compatible with the Gateway cloning system. For cloning purposes a
- 674 coding sequence of *TaCKX1* (NCBI JN128583) 378 codons long was used. In the first step,
- 675 the cassette was amplified using: EAC11-F: 5'-
- 676 TTGAATTCGACTTCGACCGCGGCGTTTT-3' and EAC12-R: 5'-
- 677 TTGAATTCATGTCTTGGCCAGGGGAGAG-3 and cloned into the entry vector
- 678 pCR8/GW/TOPO (Invitrogen). In the next step, the cassette was cloned to the destination
- 679 Bract7 vector in the Gateway reaction. The presence of the silencing cassette in the vector
- 680 was verified by restriction analysis and sequencing. The vector was electroporated into the
- 681 AGL1 strain of Agrobacterium tumefaciens and used for transformation.

682 Plant material, Agrobacterium-mediated transformation and in-vitro culture

- 683 The spring cultivar of common wheat (*Triticum aestivum* L.) Kontesa was used as a donor
- 684 plant for transformation experiments as well as transgenic plants. Seeds were germinated into
- 685 Petri dishes for one day at 4°C and then five days at room temperature in the dark. Six out of
- ten seedlings from each Petri dish were replanted into pots with soil. The plants were grown
- 687 in a growth chamber under controlled environmental conditions with 20°C/18°C day/night
- temperatures and a 16 h light/8 h dark photoperiod. The light intensity was 350 μ mol·s⁻¹·m⁻².
- 689 Agrobacterium-mediated transformation experiments were performed according to our
- 690 previously described protocols for wheat (Przetakiewicz et al., 2003; Przetakiewicz et al.,
- 691 2004). Putative transgenic plants were regenerated and selected on modified MS media
- 692 containing 25 mg l^{-1} of hygromycin as a selectable agent.
- 693 First, 7 days after pollination (DAP) spikes from T_1 , T_2 and control plants were collected for
- 694 RT-qPCR and phytohormone quantification. Only 1/3 of the middle part of each spike was
- 695 used for experiments (upper and lower parts were removed).

696 PCR analysis

- 697 Genomic DNA was isolated from well-developed leaves of 14-day plants according to the
- 698 modified CTAB procedure (Murray and Thompson, 1980) or by using the KAPA3G Plant
- 699 PCR Kit (Kapa Biosystems). The PCR for genomic DNA isolated by CTAB was carried out
- in a 25 ml reaction mixture using Platinum Taq DNA Polymerase (Invitrogen) and 120 ng of
- template DNA. The reaction was run using the following program: initial denaturation step at
- 702 94°C for 2 min, 35 cycles of amplification at 94°C for 30 s, 65°C for 30 s, 72°C for 30 s with
- a final extension step at 72°C for 5 min. The PCR for genomic DNA isolated by KAPA3G
- was carried out in a 50 µl reaction mixture using 1 U of KAPA3G Plant DNA Polymerase and

- a 0.5 x 0.5 mm leaf fragment. The reaction was run using the following program: initial
- denaturation step at 95°C for 3 min, 40 cycles of amplification at 95°C for 20 s, 68°C for 30 s,
- 707 72°C for 30s with a final extension step at 72°C for 2 min.
- 708 Putative transgenic T_0 and T_1 plants were tested with two pairs of specific primers amplifying
- a fragment of the *hpt* selection gene. The sequences of the primers for the first pair were:
- 710 hygF1 5'-ATGACGCACAATCCCACTATCCT-3' and hygR1 5'-
- 711 AGTTCGGTTTCAGGCAGGTCTT-3', and the amplified fragment was 405 bp. The
- requences of the primers for the second pair were: hygF2 5'-GACGGCAATTTCGATGATG-
- 713 3' and hygR2 5'- CCGGTCGGCATCTACTCTAT-3', and the amplified fragment was 205 bp.
- 714 Non-transgenic, null segregants were used as a control.

715 **RNA extraction and cDNA synthesis**

- 716 Total RNA from 7 DAP spikes was extracted using TRI Reagent and 1-bromo-3-
- chloropropane (BCP) (Sigma-Aldrich) according to the manufacturer's protocol. The purity
- and concentration of the isolated RNA were determined using a NanoDrop spectrophotometer
- 719 (NanoDrop ND-1000) and the integrity was checked by electrophoresis on 1.5% (w/v)
- agarose gels. To remove the residual DNA the RNA samples were treated with DNase I,
- 721 RNase-free (Thermo Fisher Scientific). Each time 1 µg of good quality RNA was used for
- 722 cDNA synthesis using the RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher
- 723 Scientific) following the manufacturer's instructions. The obtained cDNA was diluted 20
- times before use in RT-qPCR assays.
- 725 Quantitative RT-qPCR
- 726 RT-qPCR assays were performed for 6 target genes: *TaCKX1* (JN128583), *TaCKX2.1*
- 727 (JF293079)/2.2 (FJ648070), TaCKX11 (3) (JN128585), TaCKX5 (Lei et al. 2008), TaCKX9
- 728 (10) (JN128591). Primer sequences designed for each gene as well as for the reference gene
- are shown in **Table S1**. All real-time reactions were performed in a Rotor-Gene Q (Qiagen)
- thermal cycler using 1x HOT FIREPol EvaGreen qPCR Mix Plus (Solis BioDyne), 0.2 µM of
- each primer, and 4 µl of 20 times diluted cDNA in a total volume of 10 µl. Each reaction was
- carried out in 3 technical replicates at the following temperature profile: $95^{\circ}C 15$ min initial
- denaturation and polymerase activation $(95^{\circ}\text{C} 25 \text{ s}, 62^{\circ}\text{C} 25 \text{ s}, 72^{\circ}\text{C} 25 \text{ s}) \times 45 \text{ cycles},$
- 734 $72^{\circ}C 5$ min, with the melting curve at 72–99°C, 5 s per step. The expression of *TaCKX*
- 735 genes was calculated according to the two standard curves method using ADP-ribosylation
- 736 *factor* (*Ref 2*) as a normalizer.

737 Relative expression/silencing of *TaCKX1* was related to mean expression of the gene in non-

- silenced control plants set as 1.00. Relative expression of other *TaCKX* genes was related to
 each tested gene set as 1.00 in non-silenced plants.
- 740 Statistical analysis was performed using Statistica 13 (StatSoft) software. The normality of
- 741 data distribution was tested using the Shapiro-Wilk test. To determine whether the means of
- two sets of data of expression levels, phytohormone concentrations, and yield-related traits
- between non-silenced and silenced lines are significantly different from each other (for p
- value less than p<0.05) Student's t-test or the Mann-Whitney test was applied. Correlation
- 745 coefficients were determined using parametric correlation matrices (Pearson's test) or a
- nonparametric correlation (Spearman's test).

747 Quantification of ABA, auxins, cytokinins and GA₃

- 748 Chemicals used for quantification were: the standard of ABA, five standards of auxins: IAA,
- 749 indole-3-butyric acid (IBA), indole-3-propionic acid (IPA), 1-naphthaleneacetic acid (NAA),
- and 2-phenylacetic acid (PAA); twenty-seven standards of CKs: *tZ*, *trans*-zeatin riboside
- 751 (tZR), trans-zeatin-9-glucoside (tZ9G), trans-zeatin-7-glucoside (tZ7G), trans-zeatin-O-
- 752 glucoside (tZOG), trans-zeatin riboside-O-glucoside (tZROG), trans-zeatin-9-glucoside-O-
- 753 glucoside (*t*Z9GOG), *trans*-zeatin-9-glucoside riboside (tZ9GR), *c*Z, *cis*-zeatin-riboside
- 754 (cZR), cis-zeatin O-glucoside (cZOG), cis-zeatin 9-glucoside (cZ9G), cis-zeatin-O-glucoside-
- riboside (cZROG), dihydrozeatin (DZ), dihydrozeatin-riboside (DZR), dihydrozeatin-9-
- 756 glucoside (DZ9G), dihydrozeatin-7-glucoside (DZ7G), dihydrozeatin-*O*-glucoside (DZOG),
- 757 dihydrozeatin riboside-O-glucoside (DZROG), N^6 -isopentenyladenine (iP), N^6 -
- isopentenyladenosine (iPR), N^6 -isopentenyladenosine-7-glucoside (iP7G), para-topolin (pT),
- 759 meta-topolin (mT), ortho-topolin (oT), 6-benzylaminopurine (6-BAP), and standard of GA₃
- 760 were purchased from OlChemIm (Olomouc, Czech Republic). Methanol (MeOH), acetonitrile
- 761 (ACN), water (LC-MS purity), and formic acid (FA) were purchased from Merck KGaA
- 762 (Darmstadt, Germany).
- For the measurement of phytohormones, 200 mg of plant powders were placed into the 2 mL
- Eppendorf tubes, suspended in 1 mL of (v/v) 50% ACN and homogenized in a bead mill (50
- 765 Hz, 5 min; TissueLyser LT, Qiagen, Germany) using two 5 mm tungsten balls. Then, samples
- 766 were homogenized using the ultrasound processor VCX 130 (max. power 130 W, max.
- 767 frequency 20 kHz, 5 min) equipped with titanium probe (Sonics & Materials Inc., USA) and
- 768 mixed in laboratory shaker (90 rpm, dark, 5°C, 30 min; LC-350, Pol-Eko-Aparatura, Poland).
- 769 Samples were centrifuged (9000×g, 5 min; MPW-55 Med. Instruments, Poland) and collected

- in a glass tube. For quantification of ABA, AXs, CKs and GA_3 , $[^{2}H_{6}](+)$ -*cis,trans*-ABA (50
- 771 ng), $[{}^{2}H_{5}]$ IAA (15 ng), $[{}^{2}H_{6}]$ iP (50 ng), $[{}^{2}H_{5}]$ tZ (30 ng), $[{}^{2}H_{5}]$ -tZOG (30 ng), $[{}^{2}H_{3}]$ -DZR (30
- ng) and $[^{2}H_{2}]$ GA₃ (30 ng) were added to samples as internal standards.
- 773 Prepared extracts were purged using a Waters SPE Oasis HLB cartridge, previously activated
- and equilibrated using 1 mL of 100% MeOH, 1 mL water, and 1 mL of (v/v) 50% ACN
- (Simura et al., 2018). Then, extracts were loaded and collected to the Eppendorf tubes and
- eluted with 1 mL of 30% ACN (v/v). Samples were evaporated to dryness by centrifugal
- vacuum concentrator (Eppendorf Concentrator Plus, Germany), dissolved in 50 μ L of (v/v)
- 30% ACN and transferred into the insert vials. Detection of analyzed phytohormones was
- performed using an Agilent 1260 Infinity series HPLC system (Agilent Technologies, USA)
- including a Q-ToF LC/MS mass spectrometer with Dual AJS ESI source; 10 μ L of each
- sample was injected on the Waters XS elect C₁₈ column (250 mm \times 3.0 mm, 5 µm), heated up
- to 50 °C. Mobile phase A was 0.01% (v/v) FA in ACN and phase B 0.01% (v/v) FA in water;
- flow was 0.5 mL min⁻¹. Separation of above hormones was done in ESI-positive mode with
- the following gradient: 0-8 min flowing increased linearly from 5 to 30% A, 8-25 min 80% A,
- 785 25-28 min 100% A, 28-30 min 5% A.
- For the optimization of MS/MS conditions, the chemical standards of analysed
- phytohormones were directly injected to the MS in positive $([M + H]^+)$ ion scan modes, then
- areas of detected standard peaks were calculated. $[M + H]^+$ was chosen because of its
- significantly better signal-to-noise ratios compared to the negative ion scan modes.
- 790 Chlorophyll content was measured using a SPAD chlorophyll meter.
- 791
- 792

793 SUPPLEMENTAL DATA

- 794 Supplemental Table S1 Primer sequences designed for reference gene and each of 6 tested
- 795 *TaCKX* genes and amplicon length.
- Supplemental Table S2 Phenotypic traits and ratio indicator in silent T₁ and not silent, control
 plants.
- Supplemental Table S3 Phenotypic traits and ratio indicator in silent T₂ and not silent, control
 plants.
- 800 Supplemental Table S4 A. B. Correlation coefficients among expression of all tested *TaCKX*
- 801 genes and enzyme activity, and phenotypic traits in non-silent (A) and highly silent T₂ plants
- 802 (B). * non-parametric analysis; in bold significant at p<0.01.

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- 808

809 FIGURE LEGENDS

- Fig. 1 A, B. Relative expression level of silenced *TaCKX1* in segregating T_1 (A) and T_2 (B)
- 811 plants. The level of expression is related to the control set as 1.00.
- 812 Fig. 2. Comparison of means of relative CKX enzyme activity and selected gene expression
- levels in T_1 (bars) and T_2 (line) generation of silenced lines. * significant at p<0.05; ** -
- significant at p<0.01.
- Fig. 3. Comparison of phenotypic effect of silencing of *TaCKX1* in T₁ and T₂ generations
- 816 based on ratio indicators. * significant at p<0.05; ** significant at p<0.01.
- Fig. 4 A, B. Phytohormone content (ng/g biomass) measured in the group of control and silent
- 818 T₂ plants (A). Phytohormone ratio indicators (mean value in silent per mean value in non-
- silent, control plants) in silent T₂ plants (B). * significant at p<0.05. Trace amounts (≤ 1.00
- 820 ng/g biomass): tZR, tZ7G, cZ9G, cZOGR, DZ9G, iP, iP7G, BA. Not detected: cZR, DZ,
- 821 DZR, iPR, IBA, IPA, NAA, PAA.
- 822 Fig. 5. Graphic presentation of coordinated effect of *TaCKX1* silencing on expression of other
- 823 *TaCKX* genes, phytohormone levels as well as phenotype in 7 DAP spikes of T_2 plants based
- 824 on ratio indicators. * significantly increased comparing to T_1 ; ? expected changes.
- Fig. 6 A-H. Models of regulation of phytohormone levels and phenotypic traits by
- 826 coordinated expression of TaCKX genes based on correlation coefficients (cc) in non-silenced
- 827 and silenced wheat plants. (cc) correlation coefficient between expression and trait; bold -
- strong, significant correlations at $p \le 0.05$ (above 0.82); gray from 0.5 to 0.6
- 829 \uparrow increased; \downarrow decreased; cZOG cZ-0-glucoside (inactive); tZGs. mainly tZ-0-
- 830 glucoside + tZ-9-glucoside-0-glucoside.
- 831
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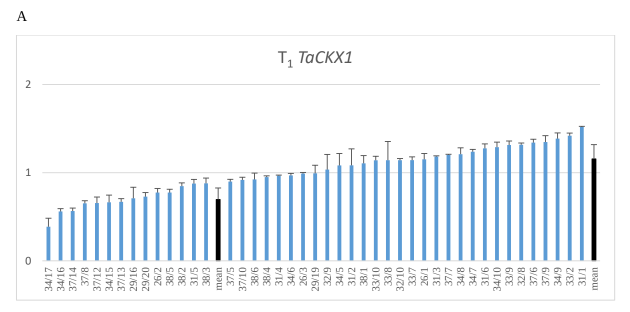
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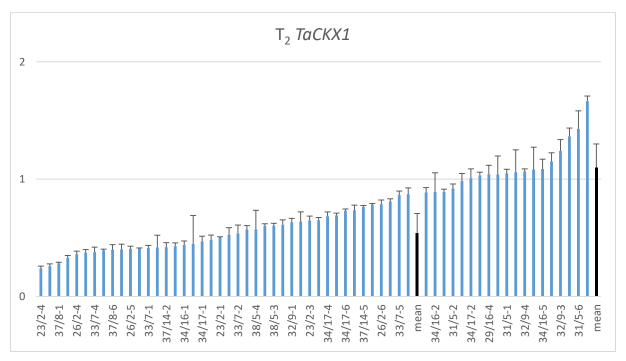


Fig. 1 A. B. Relative expression level of silenced *TaCKX1* in segregating T_1 (A) and T_2 (B) plants. The level of expression is related to the control set as 1.00.

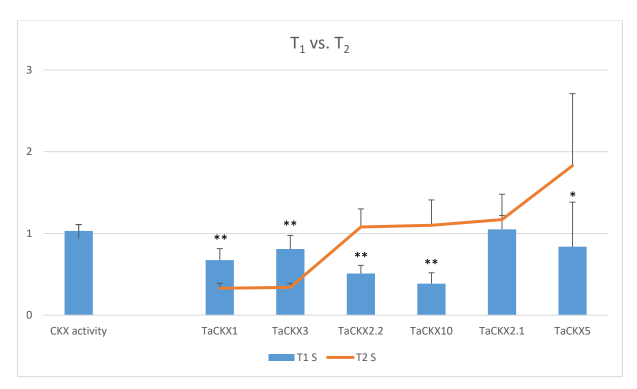


Fig. 2. Comparison of means of relative CKX enzyme activity and selected gene expression levels in T_1 (bars) and T_2 (line) generation of silenced lines. * - significant at p<0.05; ** - significant at p<0.01.

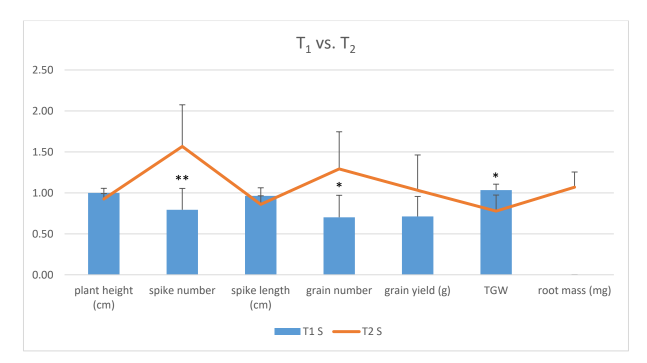
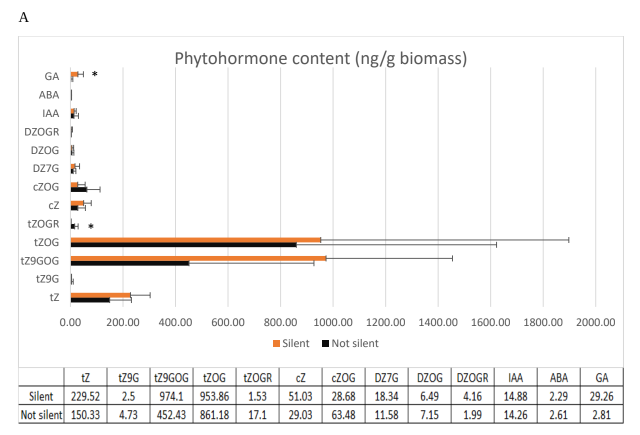


Fig. 3. Comparison of phenotypic effect of silencing of *TaCKX1* in T_1 and T_2 generations based on ratio indicators. * - significant at p<0.05; ** - significant at p<0.01.



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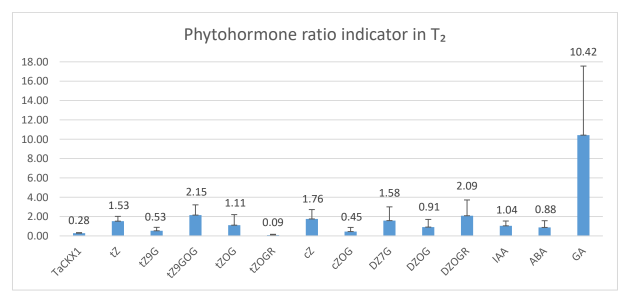
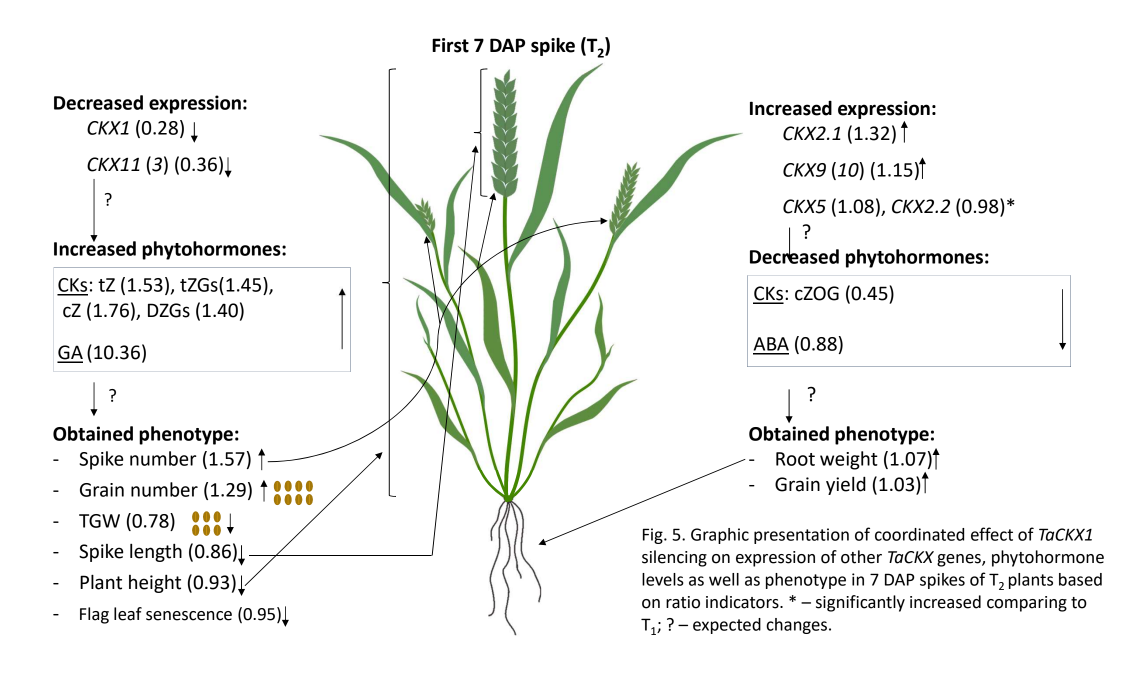


Fig. 4 A. B. Phytohormone content (ng/g biomass) measured in the group of control and silent T_2 plants (A). Phytohormone ratio indicators (mean value in silent per mean value in not silent, control plants) in silent T_2 plants (B). * - significant at p<0.05. Trace amounts (≤ 1.00 ng/g biomass): tZR, tZ7G, cZ9G, cZOGR, DZ9G, iP, iP7G, BA. Not detected: cZR, DZ, DZR, iPR, IBA,IPA, NAA, PAA.



	Non-silent	Observed phenotype		Silent T ₂	
	(cc) expression = phytohormone = expected phenotype	Not-silent Silent		(cc) expression = phytohormone = expected phenotype	
A	bioRxiv preprint doi: https://doi.org/10.1101/2020.01.07.897421; this (which was not certified by peer review) is the author/funder, who h ? - BA + IAA + GA	version posted January 8, 20 as granted bioRxiv a license t BY 4.0 International license. ← Plant l	We want	this preprint petuity. It is ? − tZ↑, tZGs↑	- smaller
В	? + cZOG – ABA	→ → ← Spike	length →	(+) $CKX2.2 \approx \rightarrow ?$ (+) $CKX5 \approx \rightarrow$ + CKX activity↓ \rightarrow – cZ↑	- shorter
С	? – tZ, cZ + GA	← TG	••••• •••••	 (-) CKX2.1↑ → + tZ↑, tZGs↑, cZ↑, iP↑ – GA (+) CKX11 (3)↓ → – tZGs↑, – iP 	- lower
D	(+) CKX1, CKX2.2, CKX5 - BA (-) CKX11 (3), CKX2.1 \rightarrow - tZGs	← Grain	yield →	(-) $CKX2.1\uparrow \rightarrow + tZGs\uparrow, cZ\uparrow$ (+) $CKX11(3)\downarrow \rightarrow + cZOG\downarrow$ $- GA\uparrow\uparrow$ (+) CKX activity≈	- higher
E	(+) CKX1, CKX2.2, CKX5 $+ iP\downarrow$ -BA (-) CKX2.1, CKX11 (3) $\rightarrow - iP$ lower	+ → → → → → → → → → → → → →	tumber →	(-) $CKX1 \downarrow \rightarrow - BA \downarrow$ (+) $CKX2.1\uparrow + CKX5 \approx \rightarrow + DZGs\uparrow$, iP↑ + GA↑↑	- higher
F	(+) <i>CKX1</i> , <i>CKX2</i> .2, <i>CKX5</i> + iP - BA - BA - iP	+ → → ← Grain n	umber →	(-) <i>CKX1</i> ↓ → − BA↓ (+) <i>CKX5</i> ≈ → + IAA≈	- higher
G	(+) <i>CKX9</i> (10) → ? (-) CKX activity	← Root v	veight →	(+) CKX11 (3) $\downarrow \rightarrow$ + cZOG \downarrow (-) CKX9 (10) $\uparrow \rightarrow$ - cZOG \downarrow (-) CKX2.2 $\approx \rightarrow$ - cZOG \downarrow (-) CKX2.1 $\uparrow \rightarrow$ + cZ \uparrow	higher
н	? + tZ, cZ – GA	← SPAD 1	st spike →	(-) <i>CKX2.1</i>↑ → + tZ ↑, tZGs, cZ ↑, DZGs↑ + GA ↑↑	- lower

Fig. 6 A-H. Models of regulation of phytohormone levels and phenotypic traits by coordinated expression of *TaCKX* genes based on correlation coefficients (cc) in non-silenced and silenced wheat plants. (cc) – correlation coefficient between expression and trait; bold – strong, significant correlations at $p \le 0.05$ (above 0.82); gray – from 0.5 to 0.6.

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