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Main Manuscript for

- Superoxide is promoted by sucrose and affects amplitude of circadian
- rhythms in the evening
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Abstract

 Plants must coordinate photosynthetic metabolism with the daily environment and adapt rhythmic physiology and development to match carbon availability. Circadian clocks drive biological rhythms which adjust to environmental cues. Products of photosynthetic metabolism, including sugars and reactive oxygen species (ROS), are closely associated with the plant circadian clock 33 and sugars have been shown to provide metabolic feedback to the circadian oscillator. Here, we
34 report a comprehensive sugar-regulated transcriptome of Arabidopsis and identify genes 34 report a comprehensive sugar-regulated transcriptome of Arabidopsis and identify genes
35 associated with redox and ROS processes as a prominent feature of the transcriptional re associated with redox and ROS processes as a prominent feature of the transcriptional response. 36 We show that sucrose increases levels of superoxide (O_2^-) which is required for transcriptional 37 and growth responses to sugar. We identify circadian rhythms of O_2 -regulated transcripts which 38 are phased around dusk and find that O_2^- is required for sucrose to promote expression of 39 *TIMING OF CAB1 (TOC1)* in the evening. Our data reveal a role for O_2^- as a metabolic signal affecting transcriptional control of the circadian oscillator in Arabidopsis.

Significance Statement

 Distinguishing the effects of light and sugars in photoautotrophic cells is challenging. The circadian system is a regulatory network that integrates light and metabolic signals and controls rhythmic physiology and growth. Our experimental approach has defined a light-independent, sugar-regulated transcriptome in Arabidopsis and revealed reactive oxygen species (ROS) as a prominent feature. ROS are by-products of photosynthetic metabolism and oscillate with circadian rhythms but have not previously been demonstrated as inputs to the plant circadian oscillator. Our data suggest a new role for superoxide as a rhythmic sugar signal which acts in the evening and affects circadian gene expression and growth.

Main Text

Introduction

 Plant metabolism is inextricably linked to daily photoperiodic cycles because of the requirement of light for photosynthesis. Anticipation and adaptation to changing light availability enables plants to optimise metabolism according to their immediate environment. Plant metabolism responds to environmental cues, such as light, temperature, biotic and abiotic stress by diverse mechanisms (1).

 Plant cells require signalling mechanisms to sense carbon and energy status and adjust metabolism. Snf1 RELATED KINASE 1 (SnRK1) and TARGET OF RAPAMYCIN 1 (TOR1) are

counteracting signalling hubs which are activated under low and replete carbon status,

 respectively (2, 3). Trehalose-6-phosphate (T6P) is an essential signalling sugar which indicates carbon status and acts through SnRK1 (4, 5).

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- Circadian clocks are an endogenous time-keeping mechanism which regulate rhythms of
- physiology and metabolism and control responses to environmental signals according to the time
- of day (6). The core circadian oscillator in Arabidopsis is a network of transcription factors
- comprised of Myb-like genes *CIRCADIAN CLOCK ASSOCIATED 1 (CCA1), LATE ELONGATED*
- *HYPOCOTYL (LHY)* and *REVIELLE (RVE)* expressed at dawn, *PSEUDO RESPONSE*
- *REGULATOR (PRR)* genes expressed through the day including *TIMING OF CAB 1 (TOC1)* at

 dusk, and the Evening Complex (EC) in the night. The phase and amplitude of gene expression 77 and protein levels are responsive to environmental cues and they, in turn, coordinate the 78 regulation of thousands of genes. regulation of thousands of genes.

 There is extensive transcriptional and post-transcriptional control of photosynthetic metabolism by the circadian clock and there is metabolic feedback on the circadian oscillator. Elevated SnRK1 activity under carbon limitation lengthens circadian period and sucrose shortens period by T6P- SnRK1 acting on the oscillator gene *PRR7* (7–9). Period also responds to glucose by a TOR- dependent mechanism (10). In continuous dark, circadian rhythms rapidly dampen, but can be sustained by addition of sugars. This effect of sugar requires GIGANTEA (GI), a clock protein which is stabilised by sucrose in the evening (11). Sugars can also reinitiate transcriptional rhythms in dark-adapted seedlings, setting phase according to the time of sugar application (8, 12), but the mechanism in unknown.

 Redox state and levels of reactive oxygen species (ROS), which are tightly linked to metabolism, are also associated with circadian rhythms in plants. There are circadian rhythms of hydrogen 92 peroxide (H₂O₂) and NADP(H)⁺ in Arabidopsis (13, 14). Circadian rhythms of peroxiredoxin oxidation have been detected across Kingdoms (15). These rhythms of redox state and associated ROS are generally considered as outputs of rhythmic metabolism controlled by the circadian clock (13), or even independent of the circadian oscillator (15). The defence hormone salicylic acid perturbs redox state and affects gating of immune response, dependent on the redox-sensitive transcription factor NON-EXPRESSOR OF PATHOGENESIS 1 (NPR1) (14). But there is presently no clear evidence of a role for redox signals as a mechanism of metabolic feedback to the circadian oscillator in plants.

 Distinguishing sugar and light signals can be challenging in photosynthetic cells since it is likely that sugar signalling will be activated in the light. Recent advances in our understanding of the impact of metabolic signalling to the plant circadian clock have relied on experiments in low light or darkness (7, 8, 10–12, 16). Here, we use an experimental approach based on the previous observation that sugar can activate expression of circadian clock genes in dark-adapted seedlings to define a light-independent, sugar-regulated transcriptome in Arabidopsis (8, 12). We compare the response of the transcriptome to sucrose in the dark and inhibition of photosynthesis in the light and identify redox and ROS processes as a prominent feature of transcriptional 109 responses to sugars. We demonstrate that superoxide $\overline{O_2}$ can act as a signal to alter gene 110 expression and growth in response to sucrose. This O_2 ⁻ signal acts to promote transcription of 110 expression and growth in response to sucrose. This O_2^- signal acts to promote transcription 111 circadian oscillator genes in the evening. These reveal that ROS can function as metabolic signals affecting circadian rhythms in Arabidopsis.

 Results To identify transcripts that are regulated by sugars in the presence and absence of light and photosynthesis, we designed an RNA-seq experiment based on the previous observation that sugars can reinitiate transcriptional circadian rhythms in dark-adapted Arabidopsis seedings (8, 12). Two-week old wild-type (Col-0) seedings were grown in the dark for 72 h to dampen circadian rhythms and establish a stabilised C starvation state. At subjective dawn, dark-adapted seedlings were transferred to media containing 10 mM mannitol (osmotic control) or sucrose and maintained in the dark or transferred to media containing 10 mM mannitol with or without 3-(3,4- dichlorophenyl)-1,1-dimethylurea (DCMU), an inhibitor of photosynthesis, and grown in the light. The four treatments provide conditions of no sugar/no light (Dark), sugar/no light (Suc), sugar/light (Light) and light/no sugar (DCMU) (Fig. 1A). We confirmed that seedling glucose content increased in the Suc and Light treatments but not in the Dark or DCMU treatments (Fig. 1B). To capture both early and late transcriptional responses within the timeframe of a typical photoperiod, shoot tissue was harvested at subjective dawn (0 h) and 0.5, 2 and 8 h after the treatments and prepared for RNA-Seq.

 We detected 5571 Suc-regulated genes that were differentially expressed between Dark and Suc treatments and 4628 DCMU-regulated genes differentially expressed between Light and DCMU (Fig. 1C, Dataset 1). The quantification of gene expression by RNA-seq was corroborated for 31 134 representative transcripts by qRT-PCR with a strong positive correlation (R^2 =0.91) (Fig. S1). The overlap of differentially expressed genes (DEGs) between time-points was relatively low (Fig. 1C), suggesting the sampling design captures a wide dynamic range of the transcriptional response. Comparison of our list of Suc-regulated genes to published microarray datasets (17, 18) indicated that we have captured a more extensive sugar-regulated transcriptome (Fig. S2A).

 To identify genes that are regulated by sugar, independent of light availability, we generated a list of genes that were upregulated by Suc in the dark and downregulated by DCMU in the light (sugar-activated; 927) or downregulated by Suc in the dark and upregulated by DCMU in the light (sugar-repressed; 1117) (Dataset 2; Fig. S3). The sugar-activated genes were enriched for Gene Ontology (GO) terms related to protein and cell wall synthesis (Fig. S3A). Sugar-repressed genes were enriched for GO terms related to light signalling, circadian rhythm and sugar metabolism (Fig. S3B, S3C). We compared our list of all 2042 sugar-regulated genes to published lists of genes regulated by SnRK1 and TOR, which are two major energy signalling hubs (2, 3). There was significant overlap with both datasets, but 1080 sugar-regulated genes were unique to this study (Fig. S3D), including 929 genes represented on ATH1 microarrays. These unique genes could represent responses either upstream or independent of SnRK1- and TOR-mediated signalling. Among the most significantly enriched GO terms in this list was Response to oxygen containing compound and Circadian rhythm (Fig. S3E).

 To define the temporal characteristics of the complete transcriptome dataset, we performed clustering analysis of expression of 18071 genes across all 53 samples using variational Bayesian Gaussian mixture models (Fig. 1D, Dataset 3). We opted for 14 clusters as a tradeoff between maximizing the explained variance and producing meaningful clusters (Fig. S4, Fig. 1D). Several clusters were associated with either sugar-repressed (clusters 1-4) or sugar-activated (clusters 11-14) genes (Fig. 1D). We searched for enriched GO terms within each cluster (Dataset 3) and summarised these using an enrichment map of the top 15 terms within each cluster (Fig. 1E, Dataset 4). Some highly enriched GO term networks were specific to one or two clusters such as inositol phosphate processes in cluster 13 or circadian rhythm and growth in clusters 8 and 13. Other enrichment GO term networks represent four or five clusters. The largest of these networks included terms associated with metabolism of sugars, nucleotides and phospholipids, chloroplast function and proteostasis. The second largest enrichment network included terms associated with reactive oxygen species (ROS) metabolism and signalling, metabolic stress and immune responses.

- Since GO terms associated with ROS appear to be a strong feature of the complete dataset, we hypothesised that ROS might be contributing to transcriptional responses to sugar. Indeed, Response to oxygen containing compound was the most significantly enriched GO term among all 2042 sugar-regulated genes and among Suc-regulated genes at 2 h (Fig. S2B). Within the former, 195 genes are associated with this GO term, including *ANNEXIN 2 (ANN2)* and six *WRKY* transcription factor genes (Fig. 2A, Dataset 5). We also identified 95 sugar-regulated genes previously reported as ROS-responsive (19), including *ASCORBATE PEROXIDASE 1 (APX1)* and *CATALASE 2 (CAT2)* (Fig. 2B, Dataset 5).
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 To test whether treatment of Arabidopsis seedlings with sucrose affects production of ROS in 179 dark-adapted seedlings, we used histochemical stains for hydrogen peroxide (H_2O_2) and 180 superoxide (O_2^-) (Fig. 2C,D). Treatment of dark-adapted seedlings with sucrose led to a decrease 181 in staining for H_2O_2 within 30 min. By contrast, sucrose treatment of dark-adapted seedlings
182 increased stain for O_2^- within 2 h, compared to mannitol controls. The elevated NBT stain was 182 increased stain for O_2 ⁻ within 2 h, compared to mannitol controls. The elevated NBT stain was observed throughout the shoot, including hypocotyl, cotyledons and leaves. To corroborate this 184 observation, we used a L-012 luminescence assay, which does not discriminate between H_2O_2
185 and O_2 , but provides better temporal resolution of ROS production than histochemical stains. 185 and O_2 , but provides better temporal resolution of ROS production than histochemical stains. 186 Consistent with the NBT stains for O_2 , we detected elevated L-012 luminescence within 2 h in 186 Consistent with the NBT stains for O_2^- , we detected elevated L-012 luminescence within 2 h in 187 sucrose-treated seedlings compared to mannitol-treated controls (Fig. 2E). Presumably, this 188 assay underestimates the difference in O_2 ⁻ production since the signal in sucrose-treated 189 seedlings will be the sum of the reduced \overline{H}_2O_2 and the increased $O_2^{\prime\prime}$ (Fig. 2C). The ROS- response detected in both the histochemical and luminescent assays is concomitant with the timing of the transcriptional response associated with ROS-related genes that we detected after 2 h (Fig. 2A, 2B, S2B, Dataset 1).

194 The accumulation of O_2^- in sucrose-treated seedlings might be a by-product of increased energy metabolism or could be contributing as a signal to affect transcriptional changes. We looked for 196 chemicals that could inhibit the sucrose-induced production of $O_2^{\text{-}}$. Diphenyleneiodonium (DPI) is 197 an inhibitor of NADPH oxidases, which generate O_2 ⁻ at the plasma membrane. Methyl viologen 198 (MV) interferes with electron transport from PS I and elevates O_2^- . 3-amino-1,2,4-triazole (3-AT) 198 (MV) interferes with electron transport from PS I and elevates O_2^- . 3-amino-1,2,4-triazole (3-AT)
199 is a catalase inhibitor which promotes H_2O_2 accumulation. We tested the effect of these chemicals on induction of a circadian-regulated luciferase reporter for *COLD, CIRCADIAN RHYTHM REGULATED 2 (CCR2)*. DPI strongly inhibited the increase of luciferase luminescence in sucrose-treated, dark-adapted *CCR2p:LUC* seedlings, whereas MV and 3-AT did not (Fig. 3A). Similarly, DPI, but not MV or 3-AT, also inhibited sucrose-induced L-012 luminescence (Fig. 3B) 204 and histochemical staining for O₂⁻but did not affect sucrose-induced changes in staining for H₂O₂ (Fig. 3C, D).

 We used the transcriptional response of *CCR2p:LUC* to generate a dose-response curve of 208 inhibition by DPI. This response was inhibited by 30% at 1 μ M DPI and by >70% at concentrations above 5 µM (Fig. 3E). Similar dose-dependent effects were also observed for two other NADPH oxidsase inhibitors, VAS2870 (20) and apocynin (21), but not for the xanthine dehydrogenase inhibitor, allopurinol (22) (Fig S5). We confirmed that DPI also inhibited sucrose- induction of *CCR2* and *WRKY60* transcripts by qRT-PCR (Fig. 3F) as well as *WRKY11p:ß- GLUCURONIDASE (GUS)* and *WRKY30p:GUS* reporters (Fig. S6). Thus, DPI effectively inhibits transcriptional regulation of multiple sugar-regulated genes.

 DPI could be inhibiting transcriptional responses to sugar in our assay by affecting uptake of sucrose, altered sugar metabolism, or inhibition of sugar sensing or signalling. We measured soluble sugars glucose, fructose and sucrose in sucrose-treated dark-adapted seedlings in the presence of DMSO or DPI. We did not detect a difference from controls for any sugar within 8 h of sucrose treatment (Fig. S7), suggesting that inhibition of sugar uptake or sucrose catabolism cannot account for the dramatic inhibition of the transcriptional response by DPI.

 Since DPI can inhibit transcriptional responses to sugar, we sought to establish whether DPI also affects other sugar-regulated processes in Arabidopsis. Seed germination in both dormant and non-dormant seeds is inhibited by exogenous sugar, acting through abscisic acid-dependent pathways (23). Similarly to sucrose, DPI also inhibits germination (24) (Fig. S8). If DPI inhibits germination by the same pathway as sucrose, we expected that their effects would be non- additive. However, the effect of DPI on inhibition of germination was detected both with and without sucrose in dormant and non-dormant seeds (Fig. S8). This suggests that DPI does not affect the regulatory pathways through which sucrose inhibits seed germination.

 Sugars promote growth. To test the effect of DPI on growth promotion by sucrose, we measured effects on hypocotyl elongation and root growth in dark-grown seedlings. This growth assay enables quantification of effects of sugar on cell elongation in the hypocotyl and cell division in the root in the absence of light signals. Seedlings growing on media containing DPI had slightly reduced hypocotyl length and root length in control media, and DPI strongly attenuated the 237 positive effects of sucrose on both hypocotyl and root length (Fig 3G). These data suggest that DPI inhibits the signalling or metabolism of sucrose to promote cell elongation and cell division.

 NADPH oxidases are encoded by a family of ten *RESPIRATORY BURST OXIDASE HOMOLOG (RBOH)* genes in Arabidopsis. We tested whether *rboh* mutants had altered ROS production in dark-adapted seedlings using L-012 luminescence assays. Both the *rbohb* and *rbohc* mutants had similar response to sucrose as wild type, but *rboha* mutants and *rbohd rbohf* double mutants had reduced L-012 luminescence (Fig. S9A), similar to wild type treated with DPI, VAS2890 or apocynin (Fig. S5B). We also tested whether *rboh* mutants had altered growth responses to sucrose (Fig. S9B). The *rbohd rbohf* double mutant had reduced root and hypocotyl length on control media compared to wild type but growth was still responsive to sucrose in the mutant. Stimulation of hypocotyl growth by sucrose was reduced in the *rboha* mutant compared to wild type, but stimulation of root growth was unaffected. Thus, although we detected small growth effects in the mutants, none of those tested were able to phenocopy the effect of DPI. Similarly, the transcriptional response of *CCR2* or *WRKY60* to sucrose in dark adapted seedlings was not 252 reduced in *rboh* mutants (Fig. S9C). These suggest that there is residual O_2 ⁻ accumulation in these mutants sufficient to elicit a response and that there is genetic redundancy in the molecular targets of DPI contributing to these sugar responses.

 Sugars affect period of circadian rhythms (8) and the circadian clock contributes to rhythms of ROS homeostasis (13). We tested the effect of DPI, MV and 3-AT on circadian rhythms in media with or without sucrose. We measured circadian rhythms of *TOC1p:LUC* in continuous low light 259 (10 μ mol m⁻² s⁻¹) because the effect of exogenous sucrose on circadian rhythms is more pronounced in these conditions (8). Circadian period was significantly shorter in seedlings grown on sucrose compared to mannitol for all ROS modifiers, similar to the DMSO control (Fig. 4A, 4B). This suggests that these chemicals did not affect the adjustment of period by exogenous sucrose.

 Sugars also affect amplitude of circadian rhythms (11). Luciferase signal is dramatically elevated in *TOC1p:LUC* seedlings transferred to media containing sucrose compared to mannitol (Fig 4A, 4C). This transcriptional response does not require GI (Fig. S10), a clock protein which is post- transcriptionally regulated by sucrose (11). The effect of sucrose in *TOC1p:LUC* seedlings was strongly attenuated in the presence of DPI, elevated in the presence of MV and unaffected by 3- 270 AT (Fig. 4C), which is consistent with the effects of these compounds on O_2^- levels. The effects of 271 DPI and MV were particularly pronounced during the night and were not observed in *CCA1p:LUC*
272 or *PRR7p:LUC* seedlings (Fig. 4C), suggesting O₂⁻ acts on specific components of the oscillator. 272 or PRR7p:LUC seedlings (Fig. 4C), suggesting \tilde{O}_2^- acts on specific components of the oscillator.

 Since the effects of DPI and MV differed between the morning-phased CCA1p:LUC and PRR7p:LUC and evening-phased TOC1p:LUC, we wondered whether this might reflect a global 276 pattern of O_2 on transcriptional rhythms. We used a set of previously reported O_2 and H_2O_2 - responsive transcripts (19) to determine their phases in continuous light from a published RNA-278 seq dataset (25). The distribution of phases of transcripts up- and down-regulated by O_2^- or H₂O₂ 278 seq dataset (25). The distribution of phases of transcripts up- and down-regulated by O_2^- or H_2O_2
279 deviated significantly from expectations (Fig. 4D, Dataset 5). The phase of transcripts 280 upregulated by H_2O_2 were enriched several hours after subjective dawn and downregulated transcripts were enriched before subjective dawn. This is consistent with the reported role of 282 CCA1 in driving rhythms of H_2O_2 which peak in the early morning (13). By contrast, the phase of 283 transcripts upregulated by O_2^- , which included *TOC1*, *GI*, *PRR5* and *LUX*, were enriched around 283 transcripts upregulated by O₂⁻, which included *TOC1, GI, PRR5* and *LUX*, were enriched around subjective dusk. About 20% of these genes are direct *TOC1* targets (26) (Dataset 5). Transcripts 285 down-regulated by O₂⁻, including LHY and RVE8, were enriched around subjective dawn. This 286 suggests that H₂O₂ and O₂⁻ production or signalling are antiphased and is consistent with a role 287 of $\overline{O_2}$ contributing to promoting oscillations of circadian transcripts in the evening.

292 **Discussion**

293 294 We have identified ROS-regulated genes as a prominent feature in the response of the 295 Arabidopsis transcriptome to sugars in both dark and light (Fig.1). The transcriptional response to 296 sucrose in dark-adapted seedlings coincides with an increase in ROS levels, including O_2^- (Fig. 297 2). Both the accumulation of O_2 and transcriptional response to sucrose were strongly attenuated 298 in seedlings treated with DPI, a chemical inhibitor of flavoenzymes including NADPH oxidases 299 (Fig. 3). DPI also inhibited the promotion of hypocotyl elongation and root growth by sucrose, 300 demonstrating a broader impact of the ROS signal in sugar responses. Finally, we found that DPI 301 inhibited the effect of sucrose on the evening expressed *TOC1* and identified a highly significant 302 anti-phasing of rhythmic transcripts that are up- and down-regulated by O_2^- to dusk and dawn, 302 anti-phasing of rhythmic transcripts that are up- and down-regulated by O_2^- to dusk and dawn,
303 respectively (Fig. 4). This is different to the redox effects of salicylic acid on both morning and 304 evening genes (14). Thus, we propose that $O₂$ functions as a metabolic signal associated with 304 evening genes (14). Thus, we propose that O_2^- functions as a metabolic signal associated variant 305 sugar levels which acts positively on the circadian oscillator in the evening. An association 306 between cellular sugar status and redox state has been long recognised in the context of 307 metabolism and oxidative stress (27), but our data provide evidence of a role for O_2^- as a 308 dynamic sugar signal affecting daily rhythms of gene expression. This effect of sugar on the 309 oscillator appears to be distinct from the T6P/SnRK1-mediated effect on period *via* transcriptional 310 regulation of *PRR7* (7) (Fig. 4) and the post-transcriptional control of GI (11) (Fig. S9) revealing 311 an additional layer of metabolic control of circadian rhythms in plants. 312

313 DPI is a potent inhibitor of NADPH oxidases which generate extracellular O_2^- at the plasma 313 DPI is a potent inhibitor of NADPH oxidases which generate extracellular O_2^- at the plasma
314 membrane activated by intracellular signals (28). We observed reduced sucrose-activated ROS 315 production and modest growth phenotypes in *rboha* and *rbohd rbohf* mutants, but the 316 transcriptional response to sucrose was similar to wild type (Fig. S8). Notwithstanding that the 317 five *rboh* mutants examined here represent over 90% of total *RBOH* gene expression (Dataset 1), 318 the subtle phenotypes in the *rboh* mutants compared to DPI-treated seedlings probably reflects 319 functional redundancy within this gene family. This will be challenging to verify, since higher order 320 mutants would be expected to be lethal. It is possible that effects of DPI on $O_2^{\frac{1}{2}}$ -mediated 320 mutants would be expected to be lethal. It is possible that effects of DPI on O_2^- -mediated 321 responses to sugar can be attributed to inhibition of other flavoenzymes. For example, in 322 photosynthetic organisms DPI inhibits O_2^- production from xanthine dehydrogenases, glutathione 322 photosynthetic organisms DPI inhibits O_2^- production from xanthine dehydrogenases, glutathione
323 reductases and mitochondrial NAD(P)H dehydrogenases (29–31). However, the similar effects of 324 VAS2890 and apocynin, but not allopurinol, on sugar responses support the role of NADPH 325 oxidases (Fig. S5). 326

327 MV interferes with electron transport from PSI, as well as in mitochondria (32), and leads to 328 accumulation of O_2^- , so the opposite effects on transcriptional responses might be expected 328 accumulation of O₂⁻, so the opposite effects on transcriptional responses might be expected
329 compared to DPI. MV was unable to induce a transcriptional response in *CCR2p:LUC* seedlings 330 without sucrose (Fig. 3A), which suggests that $O₂^-$ alone does not activate circadian gene 331 expression or that the site of O_2^- accumulation in MV-treated seedlings is not sufficient to act as 331 expression or that the site of O_2^- accumulation in MV-treated seedlings is not sufficient to act a
332 the signal. However, MV elevated the response to sucrose in *TOC1p:LUC* seedlings (Fig. 4C) 333 suggesting that O_2^- and sucrose might act synergistically.

334

335 O_2^- is generated in mitochondria, chloroplasts, peroxisomes and the apoplast (28). O_2^- is typically 336 scavenged quickly by superoxide dismutases. Elevation of O_2^- could be due to increased 337 production or reduced scavenging. The increase in $O₂^-$ triggered by sucrose in dark-adapted 337 production or reduced scavenging. The increase in O_2^- triggered by sucrose in dark-adapte
338 seedlings by histochemical stain and L-012 assay was relatively low and slow compared to 339 elicitor-induced respiratory burst (33) but faster than a ROS effect reported for cell-wall damage 340 (34). It might be that sucrose generates O_2^- in specific cell-types or subcellular locations and the 341 signal might be diluted in bulk tissues or our detection methods might have insufficient sensitivity. 342 This might explain why we couldn't detect L-012 luminescence in *rbohd rbohf* double mutants (Fig 343 S8A). Thus, it will be useful to map the cellular and subcellular location of the O_2^- signal using the

 expanding toolset of available redox probes (35–37). This will also provide clearer identity of candidate proteins producing the signal.

 Reversible oxidation of redox-sensitive proteins by ROS can alter their activity. In Arabidopsis, 348 redox-sensitive proteins that are oxidised by H_2O_2 have been identified in most cellular
349 compartments (38). These include plasma membrane receptors (39), glycolytic enzyme

compartments (38). These include plasma membrane receptors (39), glycolytic enzymes (38, 40)

 which can localise in the nucleus and associate with DNA (41, 42) and transcription factors (43). Thus, localised changes in redox state could affect signalling pathways and gene expression by

352 various mechanisms. Changes in localised O_2^- concentration could modify protein function 353 indirectly after dismutation to H_2O_2 or directly by affecting Fe-S proteins (28).

355 It is experimentally difficult to separate the effects of H_2O_2 , O_2^- or other ROS on protein oxidation. 355 It is experimentally difficult to separate the effects of H_2O_2 , O_2^- or other ROS on protein oxida
356 Differences in target specificity for ROS might depend on their redox dynamics or subcellular 357 location. H₂O₂ is regarded as the most likely ROS signal because it is relatively stable compared 358 to the more reactive O₂⁻ (28). However, our phase analyses of H₂O₂ and O₂⁻ regulated transcripts 358 to the more reactive O_2^- (28). However, our phase analyses of H_2O_2 and O_2^- regulated transcripts indicates clear temporal separation of their effects (Fig. 4). This might reflect differences in spatial organisation of oxidative metabolism at different times of day. The mechanism by which sugar- activated O₂⁻ production affects gene regulation will depend on its cellular location. 361
362

 By examining the effects of sugar on the Arabidopsis transcriptome independently of light, we 364 have uncovered a role for redox status, exemplified by accumulation of O_2^- , that promotes 364 have uncovered a role for redox status, exemplified by accumulation of O_2^- , that promotes
365 responses to sugar including growth and circadian rhythms. In contrast to the previously reported 366 association of circadian rhythms of H₂O₂, which are phased in the morning (13), the O₂⁻-activated 366 association of circadian rhythms of H_2O_2 , which are phased in the morning (13), the O_2^- -activated 367 transcriptome peaks in the evening and includes core genes within the circadian oscillator. Sugar 368 promotes O_2 ⁻ which alters gene expression by either an extracellular or intracellular redox signal which could transmit to the nucleus via signalling or protein localisation. We propose that this metabolic signal functions to coordinate rhythmic physiology and growth in response to environmental conditions that affect photosynthetic metabolism.

Materials and Methods

 Details of plant materials and growth conditions, RNA-Seq and clustering, qRT-PCR, histochemical stains, luminescence assays and sugar quantification are described in SI Appendix. Primers are listed in Dataset 6.

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Figures legends

 Figure 1. A light-independent sugar-regulated transcriptome of Arabidopsis. (A) Two week old seedlings were grown in the dark for 72 h, then transferred to 10 mM mannitol (Dark) or sucrose 503 (Suc) in the dark, or into the light with 10 mM mannitol (Light) or 20 μ M DCMU and 10 mM mannitol (DCMU). Shoot tissue was collected at 0, 0.5, 2 and 8 h for RNA-Seq. (B) Leaf glucose content in seedlings grown as in (A) (means ± SD, *N* = 3; * *P* < 0.05 from Dark; Bonferroni- corrected *t*-test). (C) Venn diagrams of differentially expressed genes at each time-point in samples collected in the dark (left) or light (right). (D) Expression trajectories of 14 clusters of co- expressed genes identified by variational Bayesian Gaussian mixture model. Pink and blue lines indicate genes identified as up/down or down/up regulated by sucrose/DCMU, respectively. The number of genes within each cluster are in parentheses. (E) Gene Ontology enrichment maps of the top 15 terms in each cluster in (D). Node colours correspond to the cluster(s) represented in (D). Node sizes are proportional to the number of genes. Selected nodes are labelled with significantly enriched, representative GO terms for each network. See Dataset 4 for the fully annotated networks.

 Figure 2. Sucrose promotes superoxide production and ROS-regulated transcripts in dark- adapted seedlings. Transcript levels of representative ROS-associated genes identified as sugar- regulated from RNA-seq that are (A) from the GO class 'responsive to oxygen-containing compound' or (B) identified from a previous study (19) (means ± SD*, N* = 3). (C) Histochemical stains for hydrogen peroxide (DAB) and superoxide (NBT) in 10 d old, dark-adapted Col-0 seedlings treated with 30 mM mannitol or sucrose. (D) DAB and NBT stain intensity in seedlings grown as in (C) (means ± SD, *N* = 6; *P* < 0.05 from mannitol; Bonferroni-corrected *t*-test). (E) L-

 012 luminescence in dark-adapted Col-0 treated with 30 mM mannitol or sucrose (means ± SEM*,* $N = 6$).

Figure 3. Modifiers of superoxide inhibit responses to sucrose. (A) Luciferase luminescence in
526 dark-adapted *CCR2p:LUC* seedlings treated with 30 mM mannitol or sucrose in the presence of dark-adapted *CCR2p*:*LUC* seedlings treated with 30 mM mannitol or sucrose in the presence of DMSO, 10 µM DPI, 2 µM MV or 200 µM 3-AT (means ± SEM, *N* = 6). (B) L-012 luminescence in dark-adapted Col-0 treated as in (A) (means \pm SEM, N = 6). (C) Histochemical NBT stain for O₂⁻ 529 and DAB stains for H_2O_2 in dark-adapted Col-0 seedlings treated with 30 mM mannitol or sucrose 530 in the presence of 0.1% DMSO or 10 μ M DPI. (D) Stain intensity in Col-0 seedlings 4 h (NBT) or in the presence of 0.1% DMSO or 10 μ M DPI. (D) Stain intensity in Col-0 seedlings 4 h (NBT) or 0.5 h (DAB) after treatment as in (A) (*N* = 6; * *P* < 0.05; *t*-test). (E) Inhibition of response of luciferase luminescence to 30 mM sucrose in dark-adapted *CCR2p:LUC* seedlings in the presence of 0 (0.1% DMSO), 1, 5 or 25 µM DPI. (means ± SEM, *N* = 3; * *P* < 0.05 from DMSO; Bonferroni-corrected *t*-test). (F) Transcript level of *CCR2* and *WRKY60,* relative to *UBQ10* in dark-adapted Col-0 seedlings 8 h after treatment with 30 mM mannitol, sucrose or sucrose with 536 10 μ M DPI (means \pm SD, $N = 4$; μ ϵ 0.05 from mannitol; Bonferroni-corrected *t*-test. (G) Hypocotyl length and root length of 5 d old dark-grown Col-0 seedlings grown on ½ MS with or without 30 mM mannitol or sucrose, 0.1% DMSO or 1 µM DPI (means ± SD*, N* = 10; * *P* < 0.05 from ½ MS; Bonferroni-corrected *t*-test).

 Figure 4. Modifiers of superoxide affect modulation of circadian rhythms by sucrose. (A) Normalised luciferase luminescence in *TOC1p:LUC* seedlings in continuous low light with 30 mM 542 mannitol (blue) or sucrose (red) in the presence of 0.1% DMSO or 10 μ M DPI, 2 μ M MV or 200 µM 3-AT (means ± SD, *N* = 4). (B) Circadian period estimates of luciferase luminescence in *TOC1p:LUC* seedlings in (A) (means ± SD, *N* = 4; * *P* < 0.05 from mannitol; Bonferroni-corrected *t*-test). (C) Luciferase luminescence in *TOC1p:LUC, PRR7p:LUC* and *CCA1p:LUC* seedlings for 546 24 h in light/dark treated as in (A) (means \pm SD, N = 4). (D) Phase of rhythmic O₂⁻- and H₂O₂- responsive transcripts in continuous light. Values are enrichment (observed/expected) of up- and 548 down-regulated genes in each 4-h phase window ($P < 0.01$; χ^2).

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37 **Supplementary Information Text**

38 **Materials and Methods**

Plant materials and growth conditions. Col-0 was used as wild-type *Arabidopsis thaliana. CCR2p:LUC, CCA1p:LUC, PRR7p:LUC* and *TOC1p:LUC* transgenic lines have been described previously (1). Mutants *rboha, rbohb, rbohc/root hair defective2-1* and *rbohd rbohf* and *WRKY11p:GUS* and *WRKY30p:GUS* transgenic lines were obtained from Arabidopsis Biological Resource Centre (ABRC). Mutant *tps1-12* (2) was backcrossed twice to Col-0. 44

45 Seeds were surface sterilised with 30% (v/v) bleach, 0.02% (v/v) Triton X-100, washed three
46 times with sterile deionised water and sown on 1/2 strength Murashige & Skoog (1/2 MS), pH 5. 46 times with sterile deionised water and sown on $\frac{1}{2}$ strength Murashige & Skoog ($\frac{1}{2}$ MS), pH 5.7 or 47 modified Hoagland media, pH 5.7 (3) solidified with 0.8% (w/v) agar Type M (Sigma). After 2 d in 47 modified Hoagland media, pH 5.7 (3) solidified with 0.8% (w/v) agar Type M (Sigma). After 2 d in
48 the dark at 4°C, seedlings were grown at 20°C in 12 h light/12 dark cycles (LD) under 100-140 48 the dark at 4°C, seedlings were grown at 20°C in 12 h light/12 dark cycles (LD) under 100-140
49 umol m⁻² s⁻¹ light. Concentrations of DPI. MV and 3-AT were based on a previous study (4) μ mol m⁻² s⁻¹ light. Concentrations of DPI, MV and 3-AT were based on a previous study (4)

50

51 For dark growth assays, seeds were germinated on $\frac{1}{2}$ MS in LD for 48 h. Within 1 h of dawn
52 before photomorphogenesis, germinated seeds were transferred to $\frac{1}{2}$ MS with 1% (w/v) agar 52 before photomorphogenesis, germinated seeds were transferred to $\frac{1}{2}$ MS with 1% (w/v) agar 53 containing treatments, wrapped in foil and grown vertically for 3 d. Plates were photographed 53 containing treatments, wrapped in foil and grown vertically for 3 d. Plates were photographed and 54 root and hypocotyl lengths were quantified with ImageJ (NIH). root and hypocotyl lengths were quantified with ImageJ (NIH). 55

56 **RNA-seq.** Col-0 seeds were sown on nylon membrane on modified Hoagland's solution and 57 grown at 45° angle. Two week old seedlings were wrapped in aluminium foil before dawn and 58 grown in the dark for 72 h. Under dim green light, dark-adapted seedlings were transferred to 58 grown in the dark for 72 h. Under dim green light, dark-adapted seedlings were transferred to
59 Hoagland's media containing 10 mM mannitol or 10 mM sucrose and maintained in the dark o 59 Hoagland's media containing 10 mM mannitol or 10 mM sucrose and maintained in the dark or 10 mM mannitol with or without 20 μM DCMU and returned to the light. Shoots of 40 seedlings were 61 collected at 0, 0.5, 2 and 8 h after treatments, snap-frozen in liquid nitrogen and stored at -80 $^{\circ}$ C 62 until processing. The RNA-seq samples were taken from two independent experiments; the first 63 produced three biological replicates for all conditions, and the second, three further replicates for 63 produced three biological replicates for all conditions, and the second, three further replicates for the dark-grown 0, 2 and 8 h conditions. RNA was extracted with RNeasy Plant Mini Kit including 65 on-column DNase treatment (Qiagen). RNA quantity and purity were confirmed using a 66 Nanodrop spectrophotometer (ThermoScientific), and samples were run on an Agilent 2100 Bioanalyzer, with RNA 6000 Nano kit, to confirm RNA integrity (all samples displayed RINs of $>$ 68 $-$ 7). mRNA sequencing libraries were prepared from 1 μ g total RNA using the NEBNext RNA Ultra 69 Directional Library preparation kit for Illumina (New England BioLabs Inc.), in conjunction with the 69 Directional Library preparation kit for Illumina (New England BioLabs Inc.), in conjunction with the 70 NEBNext Poly(A) mRNA Magnetic Isolation Module and NEBNext multiplex oligos for Illumina NEBNext Poly(A) mRNA Magnetic Isolation Module and NEBNext multiplex oligos for Illumina 71 (dual 8 bp indexing primers set 1), according to the manufacturer's instructions. Libraries were
72 pooled at equimolar ratios, and the pool was sent for 2×150 base paired end sequencing on a 72 pooled at equimolar ratios, and the pool was sent for 2 x 150 base paired end sequencing on a
73 HiSeg 3000 at the University of Leeds Next Generation Sequencing Facility. Each sample was HiSeg 3000 at the University of Leeds Next Generation Sequencing Facility. Each sample was 74 sequenced twice on two separate lanes, except replicate 3 of the light 2 h condition, which failed 75 and was resequenced on one lane only, and replicate 1 of the 0 h condition in experiment 2, 75 and was resequenced on one lane only, and replicate 1 of the 0 h condition in experiment 2, which also failed and was not resequenced. Raw reads have been uploaded to the European 77 Nucleotide Archive, ENA accession PRJEB40453 [these will be made public on acceptance].

78

79 RNA-seq samples were quantified with Salmon v0.8.2 (5) using options -I ISR, --seqBias, --
80 gcBias, --useVBOpt and --numBootstraps 30 and providing both lanes of sequencing for ead

80 gcBias, --useVBOpt and --numBootstraps 30 and providing both lanes of sequencing for each
81 sample as input. The reference was Araport11 files Araport11 genes.201606.cdna.fasta.gz an 81 sample as input. The reference was Araport11 files Araport11_genes.201606.cdna.fasta.gz and 82 Araport11 GFF3 genes transposons.201606.gtf.gz. downloaded from 82 Araport11_GFF3_genes_transposons.201606.gtf.gz, downloaded from

83 https://www.arabidopsis.org/download/index-
84 auto.isp?dir=%2Fdownload_files%2FGenes%

84 auto.jsp?dir=%2Fdownload_files%2FGenes%2FAraport11_genome_release on 26 April 2017
85 (included in Drvad repository II). A map of transcript names to gene names to use with Salmon 85 (included in Dryad repository []). A map of transcript names to gene names to use with Salmon
86 option -g was created with the following Unix one liner:
87 cut -f9 Araport11 GFF3 genes transposons.201606.gtf | sort | u option -g was created with the following Unix one liner:

- 87 cut -f9 Araport11_GFF3_genes_transposons.201606.gtf | sort | uniq | 88 perl -ne 'print "S1\t\$2\n" if /transcript id "(.+)"; gene id "(.+)"
- 88 perl -ne 'print " $\frac{1}{51}$ + $\frac{1}{52}$ if /transcript_id "(.+)"; gene_id "(.+)";/'
89 > Araport11 GEE3 gene transposons 201606 salmon geneMap tsy
- 90 Salmon output was converted to sleuth-compatible format with wasabi
91 (https://github.com/COMBINE-lab/wasabi.commit f31c73e). These files
- 91 [\(https://github.com/COMBINE-lab/wasabi,](https://github.com/COMBINE-lab/wasabi) commit f31c73e). These files will be included in a
92 Drvad repository (https:datadryad.org) on acceptance but can be accessed during peer revie
- Dryad repository (https:datadryad.org) on acceptance but can be accessed during peer review
- 93 here https://drive.google.com/drive/folders/18zc1PCFvZaRTnxTce3lVdhwPFZ11inCm/.
- 94 95

96 Differential expression was analysed with Sleuth v0.29.0 (6) with multiple testing correction by
97 stageR v0.1.0, commit 59af4d7 (7), against the Araport11 gene annotation (8) imported from 97 stageR v0.1.0, commit 59af4d7 (7), against the Araport11 gene annotation (8) imported from
98 Ensembl Genomes release 36 (9) with biomaRt (10). Models were run with a log2 transforma 98 Ensembl Genomes release 36 (9) with biomaRt (10). Models were run with a log2 transformation
99 function on the counts (log2(x+0.5)). A Sleuth model was built for each pairwise comparison 99 function on the counts (log2(x+0.5)). A Sleuth model was built for each pairwise comparison (Dark vs Sucrose 0.5 h, Dark vs Sucrose 2 h, Dark vs Sucrose 8 h, Light vs DCMU 0.5 h, Light vs 101 DCMU 2 h, Light vs DCMU 8 h) with differentially expressed genes detected with a Wald test for 102 each comparison. A full model was run on all samples including control 0 h samples with each comparison. A full model was run on all samples including control 0 h samples with 103 differentially expressed genes detected with a likelihood ratio test. Screening p-values for stageR 104 were taken from the full model's likelihood ratio test and confirmation p-values from the pairwise
105 models' Wald tests, stageR results targeted a 10% overall false discovery rate using the Holm models' Wald tests. stageR results targeted a 10% overall false discovery rate using the Holm 106 method for family-wise error rate correction. R code to run Sleuth and stageR analyses is 107 provided in our Dryad repository (run sleuth R, run stageR R). Comparisons between ge 107 provided in our Dryad repository (run_sleuth.R, run_stageR.R). Comparisons between gene lists
108 vere made using a Venn diagram tool http://bioinformatics.psb.ugent.be/webtools/Venn/. Gene were made using a Venn diagram tool [http://bioinformatics.psb.ugent.be/webtools/Venn/.](http://bioinformatics.psb.ugent.be/webtools/Venn/) Gene 109 ontology (GO) enrichment of these lists used PANTHER Classification System (11) accessed 110 through The Arabidopsis Information Resource (TAIR). through The Arabidopsis Information Resource (TAIR).

111

q**RT-PCR.** cDNA was prepared from 0.5 µg RNA in 10 µl reactions using Tetro cDNA synthesis 113 kit (Bioline). 0.5 ng/µl of cDNA was used in each PCR reaction with 0.2 µM primers in the 113 kit (Bioline). 0.5 ng/μl of cDNA was used in each PCR reaction with 0.2 μM primers in the 114 SensiFAST SYBR no-ROX kit (Bioline) on a CFX96 Touch Real-time PCR detection syste SensiFAST SYBR no-ROX kit (Bioline) on a CFX96 Touch Real-time PCR detection system (Bio-115 Rad). PCR reaction efficiencies were determined for each primer pair using LinRegPCR (12) and 116 transcript levels were determined for target and reference genes using (mean PCR efficiency)^{-Ct}. 116 transcript levels were determined for target and reference genes using (mean PCR efficiency)^{-ct}.
117 Primer sequences are listed in Dataset 5. Primer sequences are listed in Dataset 5.

118

127

119 **Transcriptome Clustering.** Genes were clustered based on Sleuth scaled_reads_per_base
120 abundance values for each sample, using scikit-learn's BayesianGaussianMixture (13) 120 abundance values for each sample, using scikit-learn's BayesianGaussianMixture (13)
121 bittos://scikit-learn.org/stable/modules/generated/sklearn.mixture.BayesianGaussianMix 121 [https://scikit-learn.org/stable/modules/generated/sklearn.mixture.BayesianGaussianMixture.html\)](https://scikit-learn.org/stable/modules/generated/sklearn.mixture.BayesianGaussianMixture.html)
122 with maximum 1000 iterations. Numbers of clusters from 2 to 20 were tested, with the 14 cluster with maximum 1000 iterations. Numbers of clusters from 2 to 20 were tested, with the 14 cluster 123 output chosen for further analysis. Gene Ontology Enrichment analysis for each cluster was 123 output chosen for further analysis. Gene Ontology Enrichment analysis for each cluster was 124 performed with R's clusterProfiler (14) 124 performed with R's clusterProfiler (14)
125 https://bioconductor.org/packages/rele

125 [https://bioconductor.org/packages/release/bioc/vignettes/clusterProfiler/inst/doc/clusterProfiler.ht](https://bioconductor.org/packages/release/bioc/vignettes/clusterProfiler/inst/doc/clusterProfiler.html) 126 [ml\)](https://bioconductor.org/packages/release/bioc/vignettes/clusterProfiler/inst/doc/clusterProfiler.html). R code for clustering is provided in the Dryad repository (cluster analysis.R).

Histochemical stains. Seeds were sown on 1/2 MS and grown in LD and 11 d old seedlings were 129 wrapped in aluminium foil at dusk. After 72 h, at subjective dusk under dim green light, seedlings 129 wrapped in aluminium foil at dusk. After 72 h, at subjective dusk under dim green light, seedlings 130 were transferred into 0.5 ml liquid 1/2 MS containing 0.1% (v/v) DMSO or chemical treatments in 130 were transferred into 0.5 ml liquid $\frac{1}{2}$ MS containing 0.1% (v/v) DMSO or chemical treatments in 131 48-well plates. At the following subjective dawn in dim green light. 0.5 ml of 60 mM mannitol or 131 48-well plates. At the following subjective dawn in dim green light, 0.5 ml of 60 mM mannitol or 132 sucrose was added (30 mM final sugar concentration). For H_2O_2 stains, 1 mg/ml (w/v) 3'3-
133 diaminobenzidine tetrahydrochloride hydrate was dissolved in 50 mM Tris acetate (pH 5.0) diaminobenzidine tetrahydrochloride hydrate was dissolved in 50 mM Tris acetate (pH 5.0). For 134 O₂-stains. 2 mg/ml (w/y) nitroblue tetrazolium was dissolved in 10 mM potassium phosphate O² – 134 stains, 2 mg/ml (w/v) nitroblue tetrazolium was dissolved in 10 mM potassium phosphate 135 buffer (pH 7.8), 10 mM NaN₃. Seedlings were vacuum infiltrated for 1 min in freshly prepared
136 staining solutions and incubated in the dark for 24 h. Samples were cleared by boiling for 5 m staining solutions and incubated in the dark for 24 h. Samples were cleared by boiling for 5 min in 137 1:1:4 lactic acid:glycerol:ethanol then transferred to 1:4 glycerol:ethanol. GUS-stains of 138 transgenic lines was performed overnight as previously (15) . Stained seedlings were mounted 139 under coverslips on microscope slides and imaged immediately with a SMZ800 stereomicrosco 139 under coverslips on microscope slides and imaged immediately with a SMZ800 stereomicroscope
140 (Nikon) or a V370 Photo flatbed scanner (Epson). DAB and NBT stain intensity were quantified in (Nikon) or a V370 Photo flatbed scanner (Epson). DAB and NBT stain intensity were quantified in 141 whole shoots by dividing integrated density by area of individual seedlings and subtracting 142 background signal in ImageJ (NIH). background signal in ImageJ (NIH). 143

144 **L-012 luminescence assay.** Clusters of 7 d old seedlings grown on ½ MS or 6 mm leaf discs from 4 week old plants grown in LD were transferred to 96-well luminescence plates (Greiner)

146 containing 250 µl liquid ½ MS before dusk (ZT12), wrapped in aluminium foil and placed in the 147 dark for 72 h. At subjective dawn under dim green light, media was replaced with 100 µl 100 µl or 147 dark for 72 h. At subjective dawn under dim green light, media was replaced with 100 μl 100 μM
148 L-012, 20 μg/ml horseradish peroxidase containing 0.01% DMSO, 10 μM DPL 2 μM MV, 0.2 mM L-012, 20 µg/ml horseradish peroxidase containing 0.01% DMSO, 10 µM DPI, 2 µM MV, 0.2 mM 149 3-AT, 20 μM VAS2870, 500 μM apocynin or 500 μM allopurinol. After 1 h of chemical pre-
150 treatment 100 μl of 60 mM sucrose or mannitol was added to each well (final sugar concer treatment 100 µl of 60 mM sucrose or mannitol was added to each well (final sugar concentration 151 30 mM). Luminescence was measured in the dark at 90 s intervals in a Lumistar Omega plater 152 reader (BMG) using a 4 mm orbital well scan.

153

154 **Luciferase luminescence assays.** For sugar-response assays, *CCR2p:LUC* seeds were sown
155 on ½ MS and grown in LD. Pairs of 10 d old seedlings were transferred into 96-well luminescenc 155 on ½ MS and grown in LD. Pairs of 10 d old seedlings were transferred into 96-well luminescence
156 olates (Greiner) containing 200 ul ½ MS with agar at dusk, wrapped in foil and grown in the dark. 156 plates (Greiner) containing 200 µl ½ MS with agar at dusk, wrapped in foil and grown in the dark.
157 1 mM D-luciferin, K-salt (Promega) was applied twice under dim green light. After 84 h in the dark 157 1 mM D-luciferin, K-salt (Promega) was applied twice under dim green light. After 84 h in the dark 158 (subjective dawn). 20 ul of 0.5% (y/y) DMSO, 50 uM DPI, 10 uM MV or 1 mM 3-AT was applied to 158 (subjective dawn), 20 µl of 0.5% (v/v) DMSO, 50 µM DPI, 10 µM MV or 1 mM 3-AT was applied to 159 seedlings under dim green light. 1 h before addition of 30 µl of 30 mM mannitol or sucrose. For 159 seedlings under dim green light, 1 h before addition of 30 μ l of 30 mM mannitol or sucrose. For 160 to the dose response curves, seedlings were transferred under dim green light to 1/2 MS media the dose response curves, seedlings were transferred under dim green light to $\frac{1}{2}$ MS media 161 containing DMSO, DPI, VAS2870, apocynin or allopurinol 12 h before application of sugar at 162 subjective dawn. Luminescence was measured in the dark at 1 h intervals in a Lumistar Omega
163 plate reader (BMG) using a 4 mm orbital well scan. plate reader (BMG) using a 4 mm orbital well scan.

164

165 To measure circadian rhythms, clusters of 5 seeds were sown on $\frac{1}{2}$ MS and grown in LD.
166 Clusters of 7 d old seedlings were transferred at dawn to $\frac{1}{2}$ MS containing 30 mM mannito Clusters of 7 d old seedlings were transferred at dawn to $\frac{1}{2}$ MS containing 30 mM mannitol or Sucrose with 0.1% (v/v) DMSO, 10 μ M DPI, 2 μ M MV or 0.2 mM 3-AT. 1 mM D-luciferin, K-salt 168 (Promega) was applied to seedlings twice prior to imaging. Luciferase was imaged in 10 min 168 (Promega) was applied to seedlings twice prior to imaging. Luciferase was imaged in 10 min integrations following 120 s of dark at 1 hr intervals with an HRPCS5 intensified CCD camera 170 (Photek) fitted with LB3 red (640 nm) and blue (470 nm) LED arrays providing light at 60 µmol m⁻² $\,$ 171 s⁻¹ for 1 LD followed by continuous low light at 10 µmol m⁻² s⁻¹. Luminescence counts were 171 s⁻¹ for 1 LD followed by continuous low light at 10 μ mol m⁻² s⁻¹. Luminescence counts were 172 extracted from ROIs using Image32 software (Photek) and circadian rhythms were analysed by
173 Fast Fourier Transform Non-linear Least Squares using Biodare2 (16). Fast Fourier Transform Non-linear Least Squares using Biodare2 (16).

174

Sugar quantification. Seedlings were grown as for the RNA-Seq experiment or pairs of seeds 176 were sown on 1/2 MS and grown in LD. Seven d old seedlings were wrapped in foil at dusk and 176 were sown on $\frac{1}{2}$ MS and grown in LD. Seven d old seedlings were wrapped in foil at dusk and 177 grown in the dark. After 72 h, seedlings were transferred under dim green light into 96 well plat grown in the dark. After 72 h, seedlings were transferred under dim green light into 96 well plates 178 containing $\frac{1}{2}$ MS with 0.1% DMSO or 10 μ M DPI. At subjective dawn, seedlings were treated with 179 30 μ L 30 mM mannitol or sucrose. 30 seedlings were harvested per biological replicate, frozen in 179 30 μ L 30 mM mannitol or sucrose. 30 seedlings were harvested per biological replicate, frozen in 180 liquid N and stored at -80 $^{\circ}$ C until processing. Soluble sugars were extracted in 80% (v/v) ethanol liquid N and stored at -80 $^{\circ}$ C until processing. Soluble sugars were extracted in 80% (v/v) ethanol 181 measured using a Sucrose/Glucose/Fructose calorimetric assay kit (Megazyme) scaled down for 182
182 96-well plates. 96-well plates.

186

187 **Fig. S1.** Quality control of RNA-seq transcript data. (A) and (B) comparison of quantification of 8
188 representative marker genes determined by RNA-Seq (A) and qRT-PCR relative to geometric 188 representative marker genes determined by RNA-Seq (A) and qRT-PCR relative to geometric 189 mean of *PP2AA3 and IPP2* (B) (means ± SD, *N* = 3). (C) and (D) comparison of quantification of 189 mean of PP2AA3 and IPP2 (B) (means \pm SD, $N = 3$). (C) and (D) comparison of quantification of 190 31 transcripts by qRT-PCR (PCR efficiency^{-Ct}) and RNA-seq (scaled reads per base). Plots are the same data coloure 191 the same data coloured by transcript (C) or treatment (D). Values are individual biological replicates. replicates.

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- 194

- 198 Comparison of genes identified as sugar-regulated in the dark in this study with two previous
199 studies (17, 18). (B) Gene Ontology enrichment of 2772 differentially-expressed genes after 2
- 199 studies (17, 18). (B) Gene Ontology enrichment of 2772 differentially-expressed genes after 2 h
200 treatment with mannitol or sucrose in the dark showing GO categories with a z-score > 2. treatment with mannitol or sucrose in the dark showing GO categories with a *z*-score > 2.

204

205 **Fig. S3.** Light-independent sugar-regulated genes in Arabidopsis. (A) Gene Ontology enrichment 206 of 927 genes that are up-regulated by sucrose in the dark and down-regulated by DCMU in the 206 of 927 genes that are up-regulated by sucrose in the dark and down-regulated by DCMU in the 207 light. (B) Gene Ontology enrichment of 1117 genes that are down-regulated by sucrose in the 207 light. (B) Gene Ontology enrichment of 1117 genes that are down-regulated by sucrose in the 208 dark and up-requiated by DCMU in the light. Fold-enrichment and z-score are plotted on the 208 dark and up-regulated by DCMU in the light. Fold-enrichment and *z-*score are plotted on the same scale. (C) RNA-seq transcript level of light-signalling genes identified as down-regulated by 210 sucrose and up-regulated by DCMU. (D) Comparison of 2042 genes identified as sugar-regulated 211 in (A) and (B) to genes reported as regulated by SnRK1 (19) and TOR (20). (E) Gene Ontology 211 in (A) and (B) to genes reported as regulated by SnRK1 (19) and TOR (20). (E) Gene Ontology
212 enrichment of 1080 sugar-regulated genes not previously identified as SnRK1- or TOR-regulate enrichment of 1080 sugar-regulated genes not previously identified as SnRK1- or TOR-regulated 213 showing GO categories with a *z*-score > 2. Fold-enrichment and *z-*score are plotted on the same scale.

Fig. S4. Optimisation of gene clustering. Elbow plot of percentage of total variance within clusters for clustering runs with k=2 to k=20. Grey is cluster with largest variance, usually representing unclustered genes for clustering runs with k=2 to k=20. Grey is cluster with largest variance, usually representing

unclustered genes.

222

225 **Fig S5.** Effects of NADPH oxidase inhibitors. (A) Inhibition of response of luciferase
226 luminescence to 30 mM sucrose in dark-adapted *CCR2p:LUC* seedlings by DPI, VA 226 luminescence to 30 mM sucrose in dark-adapted *CCR2p:LUC* seedlings by DPI, VAS2870,
227 apocynin or allopurinol in the presence of four concentrations of each chemical inhibitor or D 227 apocynin or allopurinol in the presence of four concentrations of each chemical inhibitor or DMSO
228 (means \pm SEM, $N = 6$; $\pm P < 0.05$ from DMSO; Bonferroni-corrected *t*-test). (B) Luciferase 228 (means ± SEM, *N* = 6; * *P* < 0.05 from DMSO; Bonferroni-corrected *t*-test). (B) Luciferase 229 luminescence in dark-adapted *CCR2p*:*LUC* seedlings treated with 30 mM mannitol or sucrose in 230 the presence of 0.1% DMSO, 10 μ M DPI, 20 μ M VAS2870, 500 μ M apocynin or 500 μ M
231 allopurinol (means ± SEM, N = 6). (C) L-012 luminescence in dark-adapted Col-0 treated 231 allopurinol (means \pm SEM, $N = 6$). (C) L-012 luminescence in dark-adapted Col-0 treated with 30
232 mM mannitol or sucrose in the presence of DMSO, 10 μ M DPI, 20 μ M VAS2870 or 500 μ M 232 mM mannitol or sucrose in the presence of DMSO, 10 μ M DPI, 20 μ M VAS2870 or 500 μ M
233 apocynin or 500 μ M allopurinol (means ± SEM, N = 12). (D) Representative images and (E) 233 apocynin or 500 μ M allopurinol (means \pm SEM, $N = 12$). (D) Representative images and (E) 234 guantification of NBT stains in dark-adapted Col-0 seedlings 4 h after treatment with 30 mM 234 quantification of NBT stains in dark-adapted Col-0 seedlings 4 h after treatment with 30 mM
235 mannitol or sucrose in presence of 0.1% DMSO, 10 uM DPI, 30 uM VAS2870, 500 uM Apoc 235 mannitol or sucrose in presence of 0.1% DMSO, 10 μ M DPI, 30 μ M VAS2870, 500 μ M Apocynin 236 or 500 μ M allopurinol (means ± SD, N = 3; * P < 0.05 from DMSO+Sucrose ; Bonferroni-corrected 236 or 500 µM allopurinol (means ± SD, *N* = 3; * P < 0.05 from DMSO+Sucrose ; Bonferroni-corrected *t*-test). 238

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Fig. S6. Sugar and DPI affect *WRKY* promoter activity. (A) GUS stains of dark-adapted 10 d old *WRKY11p-GUS* and *WRKY30p-GUS* seedlings treated with 30 mM mannitol or sucrose, pre-246 treated for 30 min with DMSO or 10 μ M DPI. (B) RNA-seq transcript levels of *WRKY11* and 247 *WRKY30* (means \pm SD, *N* = 3). *WRKY30* (means ± SD*, N* = 3).

251 **Fig. S7.** Soluble sugar content in DPI-treated seedlings. Glucose, sucrose and fructose content in dark-adapted Col-0 seedlings treated with 30 mM sucrose in the presence of 0.1% DMSO or 10 μ M DPI. Values are mean dark-adapted Col-0 seedlings treated with 30 mM sucrose in the presence of 0.1% DMSO or 10 µM DPI. Values are means ± SD, *N* = 4. No significant difference was identified between DMSO or DPI treated seedlings by *t*-test with Bonferroni correction, *P* < 0.05.

Fig. S8. Additive effects of DPI and sucrose on seed germination. (A) Percentage of germinated (A) non-dormant Col-0 seeds following 2 d chilling at $\frac{4}{9}$ C or (B) dormant seeds without chilling 260 sown on $\frac{1}{2}$ MS with or without 30 mM mannitol or sucrose and 0.1% DMSO or DPI. Values are 260 sown on 1/2 MS with or without 30 mM mannitol or sucrose and 0.1% DMSO or DPI. Values are
261 mean ± SD of four independent seed populations. mean \pm SD of four independent seed populations.

Fig. S9. NADPH oxidases contribute redundantly to sugar responses. (A) L-012 luminescence in dark-adapted Col-0 (with or without 10 µM DPI), *rboha, rbohb, rbohc* and *rbohd rbohf* seedlings 268 after treatment with 30 mM mannitol or sucrose (means \pm SEM, N = 6). (B) Hypocotyl length and 269 root length of 5 d old dark-grown Col-0, *rboha, rboha, rbohb, rbohc, rbohd rbohf* and *tps1-12* root length of 5 d old dark-grown Col-0, *rboha, rboha, rbohb, rbohc, rbohd rbohf* and *tps1-12* seedlings grown on ½ MS with 30 mM mannitol or sucrose (means ± SD*, N* = 10; * *P* < 0.05 from 271 mannitol, *t*-test). (C) Transcript level of *CCR2* and *WRKY60*, relative to *UBQ10* in dark-adapted
272 Col-0 and *rboh* mutant seedlings (control) or 12 h after treatment with 30 mM mannitol or sucros 272 Col-0 and *rboh* mutant seedlings (control) or 12 h after treatment with 30 mM mannitol or sucrose
273 (means ± SD, N = 3; * P < 0.05 from Col-0; Bonferroni-corrected *t*-test). (means ± SD, *N* = 3; * *P* < 0.05 from Col-0; Bonferroni-corrected *t*-test).

278 **Fig. S10.** Effects of ROS chemicals on circadian rhythms. Luciferase luminescence in Col-0
279 TOC1p:LUC and gi-2 TOC1p:LUC seedlings in continuous light with or without 90 mM sucros

- *TOC1p:LUC* and *gi-2 TOC1p:LUC* seedlings in continuous light with or without 90 mM sucrose $(means \pm SEM, N = 4).$
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- **Dataset 1 (separate file).** Differentially expressed genes between Dark and Suc or Light and DCMU.
- **Dataset 2 (separate file).** Lists of sugar-activated and sugar-repressed genes.
- **Dataset 3 (separate file).** Gene lists and GO enrichment of 14 clusters.
- **Dataset 4 (separate file).** Complete GO enrichment map of top 15 terms from 14 gene clusters.
- **Dataset 5 (separate file).** Gene lists and phase analysis of ROS-regulated genes.
- **Dataset 6 (separate file).** Primer sequences.

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