1 **Title:**

2 Cell–cell adhesion in plant grafting is facilitated by β-1,4-glucanases

3

Authors: Michitaka Notaguchi^{1,2,3*}, Ken-ichi Kurotani¹, Yoshikatsu Sato^{3,4}, Ryo
Tabata², Yaichi Kawakatsu², Koji Okayasu², Yu Sawai², Ryo Okada², Masashi Asahina⁵,

6 Yasunori Ichihashi^{6,7}, Ken Shirasu^{6,8}, Takamasa Suzuki⁹, Masaki Niwa² and Tetsuya

- 7 Higashiyama^{3,4,8}
- 8

9 Author affiliations:

- ¹Bioscience and Biotechnology Center, Nagoya University, Furo-cho, Chikusa-ku,
- 11 Nagoya 464-8601, Japan.
- 12 ²Graduate School of Bioagricultural Sciences, Nagoya University, Furo-cho,
- 13 Chikusa-ku, Nagoya 464-8601, Japan.
- ¹⁴ ³Institute of Transformative Bio-Molecules, Nagoya University, Furo-cho, Chikusa-ku,
- 15 Nagoya 464-8601, Japan.

⁴Graduate School of Science, Nagoya University, Furo-cho, Chikusa-ku, Nagoya
464-8601, Japan.

⁵Department of Biosciences, Teikyo University, Utsunomiya, Tochigi 320-8551, Japan.

19 ⁶Center for Sustainable Resource Science, RIKEN, Tsurumi, Yokohama, Kanagawa

- 20 230-0045, Japan.
- ⁷RIKEN BioResource Research Center, Tsukuba, Ibaraki 305-0074, Japan.
- 22 ⁸Graduate School of Science, The University of Tokyo, Hongo, Bunkyo-ku, Tokyo
- 23 113-0033, Japan.

⁹College of Bioscience and Biotechnology, Chubu University, Matsumoto-cho, Kasugai

- 25 487-8501, Japan.
- 26
- 27 ^{*}Author for correspondence: Michitaka Notaguchi
- 28 Tel: +81 52 789 5714
- 29 E-mail: notaguchi.michitaka@b.mbox.nagoya-u.ac.jp
- 30

Plant grafting is conducted for vegetative propagation in plants, whereby a piece of 1 2 living tissue is attached to another tissue through establishment of cell-cell adhesion. Plant grafting has a long history in agriculture and has been applied to improve 3 4 crop traits for thousands of years¹. Plant grafting has mostly relied on the natural ability of a plant for wound healing. However, the compatibility of cell-cell adhesion 5 typically limits graft combinations to closely related species²⁻⁴, and the mechanism 6 7 by which cell-cell adhesion of injured tissues is established is largely unknown. Here, 8 we show that a subclade of β -1,4-glucanases secreted into the extracellular region 9 facilitates cell-cell adhesion near the graft interface. *Nicotiana* shows a propensity 10 for cell-cell adhesion with a diverse range of angiosperms, including vegetables, fruit trees, and monocots, in which cell wall reconstruction was promoted in a 11 similar manner to conventional intrafamily grafting⁵⁻⁷. Using transcriptomic 12 approaches, we identified a specific clade of β -1,4-glucanases that is upregulated 13 14 during grafting in successful graft combinations but not in incompatible grafts and 15 precedes graft adhesion in inter- and intrafamily grafts. Grafting was facilitated 16 with an overexpressor of the β -1,4-glucanase and, using *Nicotiana* stem as an 17 interscion, we produced tomato fruits on rootstocks from other plant families. Our 18 results demonstrate that the mechanism of cell-cell adhesion is partly conserved in 19 plants and is a potential target to enhance plant grafting techniques.

20 Plant grafting is the procedure of connecting two or more pieces of living plant 21 tissues to grow as a single plant, for which the healing of the wound site is accomplished through the adhesion of proliferated cells^{2–4}. Use of grafting is necessary for propagation 22 23 of many fruit trees worldwide, such as apples, pears, grapes, and citrus, for vegetable 24 cultivation in Asian and European countries to obtain the benefits of certain rootstocks, such as disease resistance and tolerance of unfavorable soil conditions, and to control the 25 quantity and quality of fruit^{1,8}. Recently, grafting has been used in scientific studies to 26 27 explore the mechanisms of systemic signaling in plants where long-distance transport of 28 phytohormones, RNAs, and proteins in the vascular system has an important molecular basis^{9–11}. Although grafting is a useful technique, for practical use the scion–stock 29 30 combination (the shoot and root parts of a graft, respectively) is limited to closely related

1 plant species. In general, grafting is successful between members of the same species, 2 genus, and family, but not between members of different families because of graft incompatibility^{2–4,9}. However, several interfamily graft combinations have been 3 reported^{12–17}, including a combination we have studied previously involving a *Nicotiana* 4 benthamiana scion (Nb, Solanaceae) and an Arabidopsis thaliana stock (At, 5 Brassicaceae)¹⁸, in which the *Nb* scion grew slowly but distinctly (Supplementary Movie 6 7 1, Extended data Fig. 1a, b). Moreover, in nature, plants that parasitize species from a different plant family have evolved a haustorium, a specialized organ that invades host 8 plant tissues and absorbs nutrients following tissue adhesion¹⁹. Therefore, plants 9 10 potentially have the ability to achieve cell-cell adhesion between members of different 11 families.

12 *Nicotiana* potential for interfamily grafts

13 We expanded the range of graft combinations to include other angiosperms and observed 14 that *Nicotiana* shows strong potential for cell-cell adhesion with phylogenetically distant 15 plant species and interfamily grafts are alive for more than 1 month (Fig. 1, 16 Supplementary Tables 1, 2). In the case where Chrysanthemum morifolium (Cm, 17 Asteraceae) was used as the stock and we conducted Cm/Cm homografts (scion/stock 18 notation) and interfamily grafts with *Glycine max* (*Gm*, soybean, Fabaceae) and *Nb*, the 19 Cm/Cm homografts established and the Cm scions produced flowers, whereas the Gm/Cm20 interfamily grafts did not establish and the Gm scion died (Fig. 1a, b). By contrast, in 21 *Nb/Cm* interfamily grafts, the *Nb* scion continued to grow four weeks after grafting (Fig. 22 1c). The Nb scion grew for more than three months until setting seeds. The viability of Nicotiana interfamily grafting was confirmed in combinations using Nb as the stock 23 (Extended data Fig. 1c). In transverse sections of the graft junctions for these 24 25 combinations, a necrotic layer formed at the graft boundary in unsuccessful Gm/Cm 26 interfamily grafts but developed only weakly in successful Cm/Cm homografts and 27 *Nb/Cm* interfamily grafts two weeks after grafting (Fig. 1d–f). Necrotic layer formation is an indicator of incompatibility in cell-cell adhesion in grafting¹⁵⁻¹⁷. These results 28 29 indicated that Nb/Cm grafts showed cell-cell adhesion despite the interfamily 30 combination. Transmission electron microscopy (TEM) revealed folded cell wall

1 remnants caused by graft injury at the graft interface of Gm/Cm unsuccessful interfamily grafts (Fig. 1g). In contrast, a thin cell wall formed in some areas of the graft interface in 2 3 *Nb/At* interfamily grafts (Fig. 1h). Serial sections indicated a decrease in cell wall 4 thickness at the graft interface (Fig. 1j-k, Extended data Fig. 2). These results are consistent with previous observations of cellular morphology at graft interfaces of 5 compatible interfamily grafts¹⁶. Taken together, these findings indicate that certain plant 6 7 species, such as Nb and other species studied, can accomplish cell-cell adhesion in 8 interfamily combinations. We then examined how widespread this capability may be 9 among angiosperms. We conducted grafting experiments using plants of seven Nicotiana 10 species (Nb was predominantly used) and an interfamily partner from 84 species in 42 families, chosen from among 416 angiosperm families²⁰. Ability for cell-cell adhesion 11 12 was evaluated based on scion viability 4 weeks after grafting, because in incompatible 13 combinations scion viability is lost soon after grafting and the loss of viability is visible after transferring grafts to a low-humidity environment. We observed that Nicotiana 14 15 species showed compatibility in interfamily grafting with 73 species from 38 families, 16 including two species of magnoliids, five species of monocots, and 65 species of eudicots, 17 such as important vegetable, flower, and fruit tree crops, with Nicotiana plants used as 18 either the scion or stock (Fig. 11, Extended data Fig. 3, Supplementary Tables 1, 2). These 19 observations indicated that the cell-cell adhesion capability of Nicotiana plants could be 20 extended for grafting with a diverse range of angiosperms.

21 *Nicotiana* promotes cell wall reconstruction during interfamily grafting

22 To examine the underlying mechanism of Nicotiana interfamily grafting, we performed 23 transcriptome analysis on graft junction samples from Nb/At interfamily grafts 2 h after 24 grafting and 1, 3, 5, 7, 10, 14, and 28 days after grafting (DAG) with the following 25 controls: intact Nb stem, and graft junctions of Nb/Nb homografts at 1, 3, 5, and 7 DAG 26 (Fig. 2). The transcriptome was distinctly changed 2 h after grafting compared with that 27 of intact Nb and the transcriptome changed gradually over time after grafting (Fig. 2a, b). 28 On the basis of clustering data, genes previously reported to be associated with grafting^{21,22} (Supplementary Table 3) were upregulated in response to Nb/At interfamily 29 grafting, including genes associated with auxin action, wound repair, and cambium, 30

1 provascular and vascular development (Fig. 2b). The expression level was comparable or 2 relatively higher than that observed in Nb/Nb homografts (Fig. 2c), which may indicate 3 that Nicotiana interfamily grafting requires greater contribution of these genes to achieve 4 graft establishment of more weakly compatible combinations. These molecular responses 5 at the graft junction were consistent with morphological changes in the Nicotiana interfamily grafts, in which cell proliferation and xylem bridge formation were observed 6 7 in the grafted region but the xylem bundle was obviously thin (Extended data Fig. 4a-e). 8 Dye tracer experiments using toluidine blue, an apoplastic tracer, and carboxyfluorescein, 9 a symplasmic tracer, provided evidence for establishment of both apoplastic and 10 symplasmic transport at 3 DAG or later (Extended data Fig. 4f-h). Moreover, transport of 11 mRNAs¹⁸ and GFP proteins across the graft junction was also detected (Extended data Fig. 4i, j), although the amount detected was weaker than that for homografts. Hence, the 12 13 viability of the Nb scions was preserved by parenchymatous tissue formation at the graft 14 interface.

15 To elucidate the molecular events in the early stage of cell-cell adhesion, we 16 extracted early-upregulated genes in the Nb scion of Nb/At interfamily grafts and 17 identified 189 genes (Fig. 2d, Supplementary Table 3). In a gene ontology (GO) enrichment analysis for these genes, the top-ranked GO terms were 'Extracellular region', 18 19 'Cell wall' and 'Apoplast' (Fig. 2e, Methods), which indicated that cell wall modification 20 was undertaken in Nicotiana interfamily grafting. Genes encoding cell wall 21 modification/reconstruction enzymes, including β -1,4-glucanase, β -1,3-glucanase, 22 xyloglucan hydrolase, and expansin, were promoted at 1 to 28 DAG (Fig. 2f). Laser 23 microdissection samples of Nb/At interfamily graft junctions confirmed the enhanced 24 expression level of a number of these genes in the cells proliferated from the cambial or 25 pith region of the graft boundary (Fig. 2g, h). Expression of genes associated with cell wall dynamics was also implicated in previous transcriptomic studies of conventional 26 intrafamily grafting⁵⁻⁷ and wounding response²³, which implies that *Nicotiana* activates a 27 mechanism for cell wall reconstruction in either intra- or interfamily grafting. 28

29 *Nicotiana* interfamily grafting requires a secreted type of β-1,4-glucanase

1 We next investigated the characteristics of Nicotiana grafting by comparing the 2 transcriptome with that of interfamily grafting using soybean (Gm), which was 3 incompatible in interfamily grafting combinations. We screened genes that were 4 upregulated in the Nb scions of Nb/At interfamily grafts but not in the Gm scions of 5 Gm/At interfamily grafts. For comparisons, we selected each homologous gene in Gm that showed the highest homology using tblastx (see Methods). Using the gene 6 7 information obtained, we extracted genes that were upregulated in Nb scions but not in 8 Gm scions. Of 189 genes upregulated in Nb scions (Fig. 2d), upregulation of 79 genes 9 was not observed in *Gm* scions (Fig. 3a, Methods). We assumed that these genes may 10 explain the difference in graft compatibility of Nb and Gm. Among the 79 genes, genes 11 associated with 'Extracellular region' and 'Cell wall' were highly conserved (Fig. 3b) (in 12 comparison with Fig. 2e, the number of genes associated with 'Extracellular region' and 13 'Cell wall' was nine out of 14, whereas the number of genes associated with the other GO 14 terms was 16 out of 50). This result again suggested that cell wall reconstruction is a 15 critical event for success of interfamily grafting of Nicotiana. This population included a 16 gene encoding β -1,4-glucanase of the glycosyl hydrolase 9B (GH9B) family, designated 17 *NbGH9B3* based on similarity to *At* genes. Expression of *NbGH9B3* was significantly 18 upregulated in the Nb/At interfamily grafts in contrast to corresponding genes in the At 19 stocks of Nb/At and the Gm scions of Gm/At interfamily grafts (Fig. 3c). Given that 20 β -1,4-glucanases of the GH9B family show cellulolytic activities and play roles in cellulose digestion, relaxation of cell wall, and cell wall construction during plant growth 21 processes, such as root elongation^{24,25}, we hypothesized that NbGH9B3 facilitates cell-22 23 cell adhesion of opposing cells at the graft boundary and further analyzed NbGH9B3 24 function in grafting.

We applied virus-induced gene silencing (VIGS) to examine the function of *NbGH9B3* in *Nb/At* interfamily grafting (Fig. 3d–f). We prepared non-infected and vector control samples for comparison. VIGS targeting of *NbGH9B3* caused failure of *Nb/At* interfamily grafting 2 weeks after grafting; the *Nb* scion was easily detached from the *At* stocks and the *Nb* tissues formed a necrotic layer on the graft surface (Fig. 3d). The expression level of *NbGH9B3* was consistent with the percentage success of grafting (Fig.

3e, f). At the graft interface of Nb scions in which NbGH9B3 was down-regulated by 1 2 VIGS, folded cell walls were frequently observed in contrast to the non-infected control 3 (Fig. 3g-j). We generated a knock-out line of NbGH9B3 (NbGH9B3-KO) using a 4 clustered, regularly interspaced, short palindromic repeats (CRISPR)/CRISPR-associated 5 protein 9 nuclease (CRISPR/Cas9) editing method (see Methods) and conducted grafting 6 experiments. The percentage success of grafting wild-type Nb scions onto At stocks was 7 91%, whereas that of NbGH9B3-KO interfamily grafting was 60% (Fig. 3k). These data 8 suggested that the β -1,4-glucanase encoded by *NbGH9B3* functions in cell wall digestion 9 at the graft interface and facilitates graft establishment in *Nicotiana* interfamily grafting.

10 **GH9B3** plays a crucial role in graft establishment

11 We hypothesized that *Nicotiana* interfamily grafting with a diverse range of plants is 12 achieved through a common mechanism of cell-cell adhesion during graft formation. To 13 test this hypothesis, we examined whether β -1,4-glucanase also functions in conventional 14 intrafamily grafting for other genera (Fig. 4a, b). We prepared homograft samples of 15 soybean (Gm), morning glory (In), maize (Zm), and Arabidopsis (At). Among GH9 16 family genes, one gene from each plant species was distinctly upregulated at 1 to 7 DAG 17 in all homografts except maize homografts, which failed to graft successfully because monocot species lack cambial activity in the stem²⁶; the genes all belonged to the *GH9B3* 18 clade (Fig. 4a, b, Extended data Fig. 5). Moreover, the GH9B3 genes were temporally 19 20 upregulated at 1 DAG but expression was not increased subsequently in *Gm* and *In* scions 21 grafted onto At stocks (Fig. 4b). These data suggested that upregulation of GH9B3 genes 22 during graft adhesion was conserved among these plants and, in the case of Nicotiana, 23 this mechanism can be switched on even in interfamily grafting. In Zm grafts, an 24 orthologous gene was not upregulated in both homografts and interfamily grafts, which 25 may imply that during the evolution of maize the mechanism to promote expression of 26 GH9B3 clade genes in response to stem injury was lost.

To examine the role of *GH9B3* genes in grafting in other plant genera, we performed seedling micrografting in *Arabidopsis* using wild-type and two T-DNA insertion mutant lines for *AtCEL3*, a *GH9B3* clade gene that was upregulated in *At* homografts (Fig. 4b). Although a significant difference in percentage success was not

observed among wild-type and mutant homografts, shoot growth after grafting was 1 2 significantly decreased in grafts of both mutant lines compared with that of the wild type 3 (Fig. 4c), which indicated that GH9B3 is required for straightforward establishment of 4 the graft connection in At and that dependency on GH9B3 is higher in Nicotiana 5 interfamily grafting (Fig. 3f, k). We next examined the effect of GH9B3 overexpression on grafting. We generated transgenic lines of Arabidopsis that overexpressed NbGH9B3 6 under the control of a wound-induced RAP2.6 promoter (NbGH9B3-OX)²⁷. The 7 percentage success of micrografting using the NbGH9B3-OX line was significantly 8 9 higher than that of wild-type grafting (Fig. 4d). Thus, it was demonstrated that GH9B3 10 functions in graft formation in plants other than Nicotiana.

11 The aforementioned results indicated that Nicotiana plants activate a 12 mechanism for graft adhesion even in interfamily grafting, which is generally activated 13 only in the case of intrafamily healable grafting. The ubiquity of GH9B3, an enzyme 14 secreted into the extracellular region, in plants enables the success of Nicotiana 15 interfamily grafting with a diverse range of angiosperms. To exploit this capability, we 16 examined whether Nicotiana could act as an intermediate in the grafting of different plant 17 families. We chose tomato as the scion because the fruit exhibit several favorable traits, such as umami flavor, nutrient richness, and lycopene production, and are cultivated 18 worldwide²⁸. We grafted the tomato scion onto At or Cm stocks using a Nicotiana 19 interscion, where the junction between the tomato scion and the Nicotiana interscion 20 21 represented an intrafamily graft. The tomato scions were successfully stabilized and 22 ultimately produced fruit 3–4 months after grafting (Fig. 4e–g, Extended data Fig. 6a). 23 We also achieved other interfamily grafts in which the scion, interscion, and stock all 24 belonged to different plant families (Supplementary Table 4, Extended data Fig. 6b). One 25 of the stock plants we used was Cm, a member of the Asteraceae, one of largest family in angiosperm, which is economically important family for oils and leaf vegetables such as 26 sunflower seeds and lettuce, and at the same time, is recognized as invasive weeds in 27 various circumstances including non-arable region²⁹. Our results therefore demonstrated 28 that grafting might increase the utilization of beneficial root systems found in natural 29 30 resources with minimal destruction of ecosystems (Fig. 4h).

1 Grafting is reliant on plants' ability for tissue adhesion and healing of wounds, 2 which is fundamental for the hardiness and vigor of plants in nature. Grafting is achieved 3 through sequential cellular processes, including wound response, cell regeneration, cell 4 proliferation, cell-cell adhesion, and cell differentiation into specific tissues, and is an important topic in plant science⁹⁻¹². In regard to cell-cell adhesion, the cell wall 5 polysaccharide matrix is heterogeneous and varies considerably in composition among 6 7 plant species, therefore in interspecific grafting differences in cell wall composition may account for incompatibility. Nicotiana shows cell-cell adhesion compatibility with 8 9 diverse plant species through the function of a conserved clade of extracellular-localized 10 β -1,4-glucanases, the GH9B3 family, which probably target cellulose, a core structural component of the cell wall in plants^{24,30}, together with other components associated with 11 cell wall dynamics (Fig. 2). Thus, we identified a typical biological function of a specific 12 13 clade of glycosyl hydrolase large gene family (hundreds number of genes are included in 14 the family, Fig 4a) through a study on grafting. Cellular processes involved in grafting 15 require characterization and the outcomes of such studies may enhance grafting 16 techniques in plant science research and agriculture worldwide.

17

18 Methods

19 Plant materials

20 Nicotiana benthamiana seeds were surface sterilized with 5% (w/v) bleach for 5 min, 21 washed three times with sterile water, incubated at 4°C for 3 days, and sown on half-22 strength Murashige and Skoog (1/2 MS) medium supplemented with 0.5% (m/v) sucrose 23 and 1% agar. The pH was adjusted to pH 5.8 with 1 M KOH. Seedlings were grown at 22°C for At and 27°C for Nb under continuous illumination of 100 μ mol m⁻² s⁻¹. For 24 25 generation of the Nb NbGH9B3-KO lines, three artificially synthesized DNA fragments (5'-TGTCAAGTTCAATTTCCCAA-3', 5'-CAATGCTTTCTTGGAACACA-3'. 26 and 5'-CCGATTACTTCCTCAAGTGT-3') were cloned as sgRNA into the pTTK352 vector 27 for CRISPR/Cas9 editing³¹. Binary vectors were introduced into Agrobacterium 28 tumefaciens strain EHA105 by electroporation and transformed into Nb plants by the leaf 29 disk transformation method³². For generation of the At NbGH9B3-OX lines, a 1687 bp 30

1 promoter sequence of RAP2.6 (At1g43160) and a 3078 bp cDNA sequence of 2 Niben101Scf01184g16001 amplified separately by PCR from the RAP2.6 plasmid vector 3 and Nb cDNA library, respectively, were cloned into pENTR/D-TOPO (Thermo Fisher Scientific, Waltham, MA, USA) using InFusion[®] (Takara Bio, Ohtsu, Japan) as an entry 4 clone. The entry clone was transferred into the pGWB1 vector³³ using the LR reaction. 5 The sequences of the primers used for PCR amplification were as follows: 6 7 gw RAP2.6pro F2, 5'-CACCTCTAGATGGGATGGTGTACTACGGATG-3' and 8 gw RAP2.6pro R2, 9 5'-GATCGGGGGAAATTCGGTACCCCTCTAGGTTTGAAATTGCGGTGGTAG-3' for 10 **RAP2.6** RAP2.6 01184g16001 F, the promoter; 11 5'-TTCAAACCTAGAGGGATGGCGTTTAGAGTGAAAG-3' and

pgwb_01184g16001_R, 5'-GATCGGGGGAAATTCGCTAACGTTTGGAACTATCAA-3'
for the *Niben101Scf01184g16001* CDS. Binary vectors were introduced into *Agrobacterium tumefaciens* strain GV3101 by electroporation. Plant transformation was
performed using the floral dip method³⁴. The *At cel3* T-DNA insertion lines
SALK_032323 and CS_803355 were obtained from the Arabidopsis Biological Resource
Center (ABRC; Ohio State University, Columbus, OH, USA). The *At* ecotype Columbia
(Col) was used as the wild type.

19

20 Grafting

21 For the grafting of Nb as the scion, 1- to 2-month-old Nb inflorescence stems were used. 22 For the other graft combinations, plants of sufficient size to perform grafting were used. 23 For the majority of herbaceous species, 2-week-old to several-month-old plants were 24 used and, for tree species, several-year-old plants were used. Wedge grafting was 25 performed on the epicotyls, stems, petioles, or peduncles. For stock preparation, stems (or 26 other organs) were cut with a 2-3 cm slit at the top. For scion preparation, the stem was 27 cut and trimmed into a V-shape. The scion was inserted into the slit of the stock and 28 wrapped tightly with parafilm. A plastic bar was set along the stock and the scion for 29 support. The entire scion was covered with a plastic bag, which had been sprayed inside 30 with water beforehand. Grafted plants were grown for 7 days in an incubator at 27°C

under continuous light (~30 μ mol m⁻² s⁻¹), or in a greenhouse at 22–35°C under natural 1 light (500–1500 μ mol m⁻² s⁻¹ during the day). After this period, the plastic bags were 2 partly opened by cutting the bags and making holes for acclimation. The next day, the 3 4 plastic bags were removed and the grafted plants were grown in an incubator at 22–25°C under continuous light (~100 μ mol m⁻² s⁻¹), in a plant growth room at 22–30°C under 5 continuous light (~80 μ mol m⁻² s⁻¹), or in a greenhouse. Grafting was determined to be 6 7 successful if the scion was alive at 4 weeks post-grafting for the combinations listed in 8 Supplemental Tables 1-3. For the test of GH9B loss- and gain-of-function effect on a 9 percentage of grafting success (Fig. 3f, k, Fig. 4d) was evaluated based on scion viability 10 2 weeks after grafting. When three plants were grafted using an interscion, grafting 11 manipulations were performed either all at once or in two steps. In the latter case, two 12 graft combinations were performed first; in one case, the future stock was grafted with a 13 Nb scion and, in the other case, the future scion was grafted onto a Nb stock. After 14 establishment of each graft, a second grafting was performed using the Nb parts of each 15 graft. To compare watering and grafting (Extended data Fig. 1b), Nb primary stems of 7 16 cm length were cut and the cut edge was trimmed into a V-shape. Expanded leaves (more 17 than 1 cm width) were removed so that water absorption was directed to stem growth dependent on the cutting sites. Half of the trimmed Nb stems were watered only and the 18 19 other half were grafted as scions onto At stocks. The stem length was measured once per 20 week after these treatments (n = 24 per treatment). All other plant materials for stem 21 grafting used in this study are listed in Supplemental Tables 1-3.

22 Micrografting of At was performed using a microscaled device constructed for micrografting following a protocol described previously³⁵. Seeds were sown on the 23 devices and the devices were put on the Hybond-N+ nylon membrane (GE Healthcare, 24 25 Chicago, IL, USA) which was placed on 1/2 MS medium supplemented with 0.5% 26 sucrose and 1% agar. Four-day-old seedlings underwent micrografting on the device. 27 After grafting, the device containing grafted seedlings were transferred to fresh 1/2 MS medium supplemented with 0.5% sucrose and 2% agar and grown at 22°C (for the test of 28 29 NbGH9B3-OX line) or 27°C (for the test of At cel3 mutant lines) for 6 days. After this 30 period, the grafted seedlings were taken out from the device and transferred to fresh 1/2

MS medium supplemented with 0.5% sucrose and 1% agar and grown at 22°C. The
phenotype was examined at 10 DAG.

3

4 Microscopy

To capture brightfield images of hand-cut sections of grafted regions, a stereomicroscope
(SZ61, Olympus, Tokyo, Japan) equipped with a digital camera (DP21, Olympus) or an
on-axis zoom microscope (Axio Zoom.V16, Zeiss, Göttingen, Germany) equipped with a
digital camera (AxioCam MRc, Zeiss) was used.

9 To observe xylem tissues, hand-cut transverse sections of the grafted stem 10 region were stained with 0.5% toluidine blue or 1% phloroglucinol. Phloroglucinol 11 staining (Wiesner reaction) was performed using 18 μ L of 1% phloroglucinol in 70% 12 ethanol followed by addition of 100 μ L of 5 N hydrogen chloride to the section samples. 13 Brightfield images were captured using a stereomicroscope or a fluorescence imaging 14 microscope.

To determine apoplasmic transport, the stems of *At* stocks were cut, and the cut edge was soaked in 0.5% toluidine blue solution for 4 h to overnight. Hand-cut transverse sections of the grafted regions were observed using a stereomicroscope (SZ61, Olympus) or a fluorescence imaging microscope (BX53, Olympus) equipped with a digital camera (DP73, Olympus) for high-magnification images. The water absorption sites were stained blue.

To determine symplasmic transport, cut leaves from At stocks were treated with 21 0.01% 5(6)-carboxyfluorescein diacetate (CF; stock solution 50 mg ml⁻¹ in acetone), 22 23 together with 0.1% propidium iodide (PI) to distinguish symplasmic transport (indicated 24 by CF fluorescence) from apoplastic transport (indicated by PI fluorescence) for 4 h to 25 overnight. Transverse sections of the grafted regions and the apical regions of Nb scions 26 were hand-cut and observed. To examine GFP protein transport, Nb scions were grafted onto transgenic 35S::EGFP At stocks ³⁶. Hand-cut transverse sections of the grafted 27 regions were made and the fluorescence images were captured using a fluorescence 28 29 imaging microscope (BX53, Olympus) or a confocal laser scanning microscopy 30 (LSM780-DUO-NLO, Zeiss). To quantify the GFP fluorescence signal in the grafted

region, lambda mode scanning was performed by collecting emissions in the 490–658 nm range with excitation at 488 nm and extracted against the GFP reference spectrum. The tile scan mode was also used to capture a wide view of the entire graft section. The z-sectioning images were processed using ZEN 2010 software to create maximum-intensity projection images.

To observe resin-embedded sections, samples were fixed with 2% 6 7 paraformaldehyde and 2% glutaraldehyde in 0.05 M cacodylate buffer (pH 7.4) at 4°C 8 overnight. After fixation, the samples were washed three times with 0.05 M cacodylate 9 buffer for 30 min each, and then postfixed with 2% osmium tetroxide in 0.05 M 10 cacodylate buffer at 4°C for 3 h. The samples were dehydrated in a graded ethanol series 11 (50%, 70%, 90%, and 100%). The dehydration schedule was as follows: 50% and 70% 12 for 30 min each at 4°C, 90% for 30 min at room temperature, and four changes of 100% 13 for 30 min each at room temperature. Dehydration of the samples was continued in 100% 14 ethanol at room temperature overnight. The samples were infiltrated with propylene 15 oxide (PO) two times for 30 min each and transferred to a 70:30 mixture of PO and resin 16 (Quetol-651, Nisshin EM Co., Tokyo, Japan) for 1 h. Then, the caps of the tubes were 17 opened and PO was volatilized overnight. The samples were transferred to fresh 100% 18 resin and polymerized at 60°C for 48 h. For light microscopy, the polymerized samples 19 were sectioned (8 µm thickness) with a microtome and mounted on glass slides. For light 20 microscopic observation, sections of 1.5 µm thickness were stained with 0.5% toluidine 21 blue (pH 7.0), mounted on the glass slides with Mount-Quick (Daido Sangyo Co., Tokyo, 22 Japan), and observed using a digital microscope (DMBA310, Shimadzu RIKA Co., 23 Tokyo, Japan). For transmission electron microscopic analysis, the polymerized samples 24 were ultra-thin-sectioned at 80-120 nm with a diamond knife using an ultramicrotome 25 (ULTRACUT UCT, Leica, Tokyo, Japan). The sections were mounted on copper grids 26 and stained with 2% uranyl acetate at room temperature for 15 min, then washed with 27 distilled water followed by secondary staining with lead stain solution (Sigma-Aldrich 28 Co., Tokyo, Japan) at room temperature for 3 min. The grids were observed using a 29 transmission electron microscope (JEM-1400Plus, JEOL Ltd, Tokyo, Japan) at an

acceleration voltage of 80 kV. Digital images were captured with a CCD camera
 (VELETA, Olympus Soft Imaging Solutions GmbH, Münster, Germany).

3

4 Transcriptome analysis

5 The grafted or intact plants were harvested at the respective time points. Approximately 6 10-15 mm of graft junction or stem tissue at a similar location was sampled. Each 7 biological replicate comprised the pooled tissues from 10 grafts or 10 intact plants. Total 8 RNA was extracted from the samples using the RNeasy Mini Kit (Oiagen, Hilden, 9 Germany) following the manufacturer's protocol. The cDNA libraries were prepared with 10 an Illumina TruSeq Stranded Total RNA kit with Ribo-Zero Plant or the BrAD-Seq method^{37,38} and sequenced for 86 bp single end with an Illumina NextSeq 500 platform 11 12 (Illumina, San Diego, CA, USA). Data preprocessing was performed as follows. Raw 13 sequence quality assessed with FastOC v0.11.4 was 14 (http://www.bioinformatics.babraham.ac.uk/projects/fastgc/). Adapters were removed 15 and data trimmed for quality using Trimmomatic v0.36 with the settings 16 TruSeq3-PE-2.fa: 2:40:15, SLIDINGWINDOW: 4:15, LEADING: 20, TRAILING: 20, and MINLEN: 30 (http://www.usadellab.org/cms/). FastQC quality control was repeated 17 18 to ensure no technical artifacts were introduced. Trimmed reads were mapped on the 19 genome assembly using HISAT2 version 2.1.0 with the settings -q -x "\$index" --dta 20 -dta-cufflink (http://daehwankimlab.github.io/hisat2/). The generated SAM files were 21 merged using converted to BAM format and SAMtools version 1.4.1 22 (http://samtools.sourceforge.net). Gene expression levels (fragments per kilobase of 23 transcript per million fragments mapped; FPKM) were estimated using Cufflinks version 24 2.1.1 with the -G option (http://cole-trapnell-lab.github.io/cufflinks/). The expression 25 fluctuation profiles were generated using Cuffdiff version 2.1.1. The reference sequences 26 version used for mapping and annotation follows: and were as Nb. 27 https://btiscience.org/our-research/research-facilities/research-resources/nicotiana-bentha 28 miana/, Nicotiana benthamiana draft genome sequence v1.0.1; At, 29 https://www.arabidopsis.org, TAIR10 release; genome Gm, 30 https://phytozome.jgi.doe.gov/pz/, Phytozome v7.0 (Gmax 109); In,

1 <u>http://viewer.shigen.info/asagao/</u>, Asagao_1.2; and

Zm,

2 <u>https://plants.ensembl.org/Zea_mays/</u>, Zea_mays.AGPv4.

3 Extraction of upregulated genes (Fig. 2d) was performed using the Cuffdiff 4 results of Nb/At and Gm/At graft samples, with three biological replicates for each, 5 according to the following criteria: (i) for evaluation using the ratio between the two 6 samples, genes whose expression level (FPKM value) is >0 in the 0 DAG samples, (ii) 7 the ratio of the value at 1 DAG to that at 0 DAG is 2 or more, (iii) the value at 3 DAG is 8 higher than that at 1 DAG, (iv) the values at 5 and 7 DAG are higher than that at 0 DAG, 9 and (v) the value at 1 DAG is higher than 10. Based on the Nb transcript sequence of the 10 extracted 189 genes, homology analysis of the amino acid sequence was performed with 11 tblastx using the At transcript as a reference, and the At gene ID closest to each gene was 12 obtained. А GO enrichment analysis was performed with DAVID 13 (https://david.ncifcrf.gov) using the obtained At gene IDs and a Venn diagram was 14 created. For each of the 189 genes, homology analysis on Gm was performed using 15 tblastx to obtain the orthologous gene of Gm. Data classification for Fig. 3a was 16 performed according to the following criteria: (i) the FPKM values at 1 and 3 DAG were lower than 15, and (ii) the ratio at 3 DAG to 1 DAG was 1.5 or less. For construction of 17 18 phylogenetic tree for plant glycosyl hydrolase gene family (Fig. 4a), GH9B3 clade genes 19 were isolated from Nb, Gm, In, Zm, and At as well as the other GH genes from At on 20 database (http://www.phytozome.net) or the TAIR the Phytozome database 21 (https://www.arabidopsis.org) and were used to test the phylogeny. For the At genes, we 22 handled 88 genes which harbor GH numbers on the annotations. The tree shown in Fig. 23 4a was reconstructed with a part of the entire phylogenetic tree using the 24 neighbor-joining method. Upper panel shows a tree topology for the GH9B3 clade genes 25 of Nb, Gm, In, Zm, and At. Lower panel shows a tree for all At GH genes where only the 26 primary branches for each GH groups were drawn. The number of genes included in each 27 clade is shown in triangles. The GH9B3 clade we called includes AtGH9B3 and 28 AtGH9B4. Data extraction for Fig. 4b was performed using the Cuffdiff results of Nb/At 29 samples as described above and Gm/Gm, Gm/At, In/In, In/At, Zm/Zm, Zm/At, and At/At 30 graft samples in biological replicates for each tissue at each time point.

1 For laser microdissection (LMD) samples, stem segments of the graft junction 2 of Nb/At interfamily grafts (~15 mm) were frozen in liquid nitrogen. Frozen samples 3 were embedded with Super Cryoembedding Medium (Section-Lab, Hiroshima, Japan) in 4 a dry ice/hexane cooling bath, and then cryosectioned into 15-µm-thick transverse 5 sections using a cryostat (CM1860, Leica) in accordance with the method of Kawamoto 39 . The sections that adhered to films were desiccated in a -20° C cryostat chamber for 6 7 30-60 min. Sections of three tissue regions from heterografts (vascular tissue adjacent to 8 the graft union, pith tissue adjacent to the graft union, and Nb scion tissue) were 9 microdissected using a LMD6500 laser microdissection system (Leica) and separately 10 collected into RNA extraction buffer composed of buffer RLT (Qiagen) and 0.01% 11 β-mercaptoethanol. Total RNA was extracted using a QIAshredder (Qiagen) and the 12 RNeasy Plant Mini kit (Qiagen) in accordance with the manufacturer's instructions. The 13 cDNA libraries were constructed using an Ovation RNA-Seq System V2 (NuGEN 14 Technologies, Redwood City, CA, USA) in accordance with the manufacturer's 15 instructions. RNA sequencing (RNA-Seq) analysis was performed as described above. 16 RNA-Seq data are available from the DNA Data Bank of Japan (DDBJ; 17 http://www.ddbj.nig.ac.jp/).

18

19 VIGS experiments

For VIGS of *Niben101Scf01184g16001* (*NbGH9B3*), a 294 bp portion of the region spanning from the 5' UTR to the first exon of *NbGH9B3* was amplified by PCR using primers harboring a partial sequence of *NbGH9B3* and CMV-A1 vector: GA_F_NbGH9B3,

24 5'-GTCACCCGAGCCTGAGGCCTGAAAAAGACACTTGATCGAAAAGC-3'; and

25 GA_R_NbGH9B3,

5'-GGGGAGGTTTACGTACACGCGTGTCCTTCAAAGAACAAAATGG-3'. For the
no-silencing vector control, a 292 bp portion of the *GFP* gene was amplified by PCR
using the following primers: GA_F_GFP,
5'-GTCACCCGAGCCTGAGGCCTGACTCGTGACCACCCTGACCTAC-3'; and
GA R GFP,

1 5'-GGGGAGGTTTACGTACACGCGGCTTGTCGGCCATGATATAGA-3'. The

2 amplified fragments were cloned between the StuI and MluI sites of the CMV-A1 vector using the Gibson assembly method⁴⁰. Plasmids containing the full-length cDNA of the 3 4 viral RNA were transcribed in vitro and leaves of 3-week-old Nb plants were dusted with 5 carborundum and rub-inoculated with the transcripts. Successful infection of the virus in 6 the upper leaves of Nb plants without deletion of the inserted sequences was confirmed 7 RT-PCR of the viral RNA using the primers CMV RNA2 2327 F, bv 8 5'-ATTCAGATCGTCGTCAGTGC-3', CMV RNA2 2814 R, and 9 5'-AGCAATACTGCCAACTCAGC-3'. Primary inoculated leaves were used for 10 secondary inoculation of Nb plants. Primary inoculated leaves were ground in 100 mM 11 phosphate buffer (pH 7.0) and 10 µL of the homogenate was placed dropwise on the 12 three expanded leaves of a new 3-week-old Nb plant and rub-inoculated. One week after 13 inoculation (corresponding to age 4 weeks), the infected Nb stem was grafted onto the 14 bolting stem of 5-week-old At plants. Two weeks after grafting, the percentage success of 15 grafting was scored based on scion survival.

16 Quantitative reverse-transcription PCR was performed using cDNA templates prepared from total RNA from the stem of intact Nb or grafted plants 3 days after grafting. 17 The PCR conditions were 50°C for 2 min, 95°C for 10 min, and 40 cycles of 95°C for 15 18 19 s followed by 60°C for 1 min; The primers used were qPCR Cellulase5 F2, 20 5'-ATTGGGAGCCAATGATGTACC-3', and qPCR Cellulase5 R2, 21 5'-TGTCATTTCCAACAACGCTTC-3'. NbACT1 (Niben101Scf09133g02006.1) was 22 used as the internal standard and amplified with the primers NbACT-F, 23 5'-GGCCAATCGAGAAAAGATGAC-3', and NbACT-R. 24 5'-AACTGTGTGGCTGACACCATC-3'. All experiments were performed with three 25 independent biological replicates and three technical replicates.

26

27 References

Mudge, K., Janick, J., Scofield, S. & Goldschmidt, E. E. A history of grafting.
 Hortic. Rev. 35, 437–493. (2009).

30 2. Hartmann, H. T. & Kester, D. E. Plant propagation: Principles and practices. 3rd

- 1 *ed.*, 314–427 (Prentice-Hall, 1975).
- Andrews, P. K. & Marquez, C. S. Graft incompatibility. *Hortic. Rev.* 15, 183–231
 (1993).
- Melnyk, C. W. Plant grafting: insights into tissue regeneration. *Regeneration* 4, 3–14.
 (2017).
- Cookson, S. J. et al. Graft union formation in grapevine induces transcriptional
 changes related to cell wall modification, wounding, hormone signalling, and
 secondary metabolism. *J. Exp. Bot.* 64, 2997–3008 (2013).
- 9 6. Ren, Y. et al. Involvement of metabolic, physiological and hormonal responses in the
 graft-compatible process of cucumber/pumpkin combinations was revealed through
 the integrative analysis of mRNA and miRNA expression. *Plant Physiol. Biochem.*12 129, 368–380 (2018).
- 13 7. Xie, L., Dong, C. & Shang, Q. Gene co-expression network analysis reveals
 14 pathways associated with graft healing by asymmetric profiling in tomato. *BMC*15 *Plant Biol.* 19, 373 (2019).
- Lee, J. M. & Oda, M. Grafting of herbaceous vegetable and ornamental crops.
 Hortic. Rev. 28, 61–124 (2003).
- Goldschmidt, E. E. Plant grafting: new mechanisms, evolutionary implications.
 Front. Plant Sci. 5, 727 (2014).
- 20 10. Wang, J., Jiang, L. & Wu, R. Plant grafting: how genetic exchange promotes
 21 vascular reconnection. *New Phytol.* 214, 56–65 (2017).
- 11. Tsutsui, H. & Notaguchi, M. The use of grafting to study systemic signaling in
 plants. *Plant Cell Physiol.* 58, 1291–1301 (2017).
- Gaut, B. S., Miller, A. J. & Seymour, D. K. Living with two genomes: grafting and
 its implications for plant genome-to-genome interactions, phenotypic variation, and
 evolution. *Annu. Rev. Genet.* 53, 195–215 (2019).
- Simon, S. V. Transplantationsversuche zwischen Solanum melongena und Iresine
 Lindeni. Jb. wiss. Bot. 72, 137–160 (1930).
- 29 14. Nickell, L. G. Heteroplastic grafts. *Science* **108**, 389 (1948).
- 30 15. Moore, R. & Walker, D. B. Studies of vegetative compatibility-incompatibility in

higher plants. II. A structural study of an incompatible heterograft between *Sedum telephoides* (*Crassulaceae*) and *Solanum pennellii* (*Solanaceae*). *Am. J. Bot.* 68,
 831–842 (1981).

- Kollmann, R. & Glockmann, C. Studies on graft unions. I. Plasmodesmata between
 cells of plants belonging to different unrelated taxa. *Protoplasma* 124, 224–235
 (1985).
- Flaishman, M. A., Loginovsky, K., Golobowich, S. & Lev-Yadun, S. *Arabidopsis thaliana* as a model system for graft union development in homografts and
 heterografts. J. Plant Growth Regul. 27, 231–239 (2008).
- 10 18. Notaguchi, M., Higashiyama, T. & Suzuki, T. Identification of mRNAs that move
 11 over long distances using an RNA-Seq analysis of Arabidopsis/*Nicotiana*12 *benthamiana* heterografts. *Plant Cell Physiol.* 56, 311–321 (2015).
- 13 19. Westwood, J. H., Yoder, J. I., Timko, M. P. & dePamphilis, C. W. The evolution of
 parasitism in plants. *Trends Plant Sci.* 15, 227–235 (2010).
- APG IV the angiosperm phylogeny group. An update of the Angiosperm Phylogeny
 Group classification for the orders and families of flowering plants: APG IV. *Bot. J. Linn. Soc.* 181, 1–20 (2016).
- 18 21. Matsuoka, K. et al. Differential cellular control by cotyledon-derived
 phytohormones involved in graft reunion of Arabidopsis hypocotyls. *Plant Cell Physiol.* 57, 2620–2631 (2016).
- 21 22. Melnyk, C. W. et al. Transcriptome dynamics at *Arabidopsis* graft junctions reveal
 22 an intertissue recognition mechanism that activates vascular regeneration. *Proc. Natl.* 23 *Acad. Sci. U. S. A.* 115, E2447–E2456 (2018).
- 24 23. Cheong, Y. H. et al. Transcriptional profiling reveals novel interactions between
 25 wounding, pathogen, abiotic stress, and hormonal responses in Arabidopsis. *Plant*26 *Physiol.* 129, 661–77 (2002).
- 27 24. Cosgrove, D. J. Growth of the plant cell wall. *Nat. Rev. Mol. Cell Biol.* 6, 850–61
 28 (2005).
- 29 25. Lewis, D. R. et al. A kinetic analysis of the auxin transcriptome reveals cell wall
 30 remodeling proteins that modulate lateral root development in Arabidopsis. *Plant*

1 *Cell* **25**, 3329–46 (2013).

- 2 26. Roodt, D., Li, Z., Van de Peer, Y. & Mizrachi, E. Loss of wood formation genes in
 3 monocot genomes. *Genome Biol. Evol.* 11, 1986–1996 (2019).
- 4 27. Matsuoka, K. et al. RAP2.6L and jasmonic acid–responsive genes are expressed
 5 upon Arabidopsis hypocotyl grafting but are not needed for cell proliferation related
 6 to healing. *Plant Mol. Biol.* 96, 531–542 (2018).
- 7 28. Bergougnoux, V. The history of tomato: from domestication to biopharming.
 8 *Biotechnol. Adv.* 32, 170–89 (2014).
- 9 29. Mandel, J. R. et al. A fully resolved backbone phylogeny reveals numerous
 10 dispersals and explosive diversifications throughout the history of *Asteraceae*. *Proc.*11 *Natl. Acad. Sci. U. S. A.* **116**, 14083–14088 (2019).
- 30. Urbanowicz, B. R. et al. Endo-1,4-b-glucanases (cellulases) of glycosyl hydrolase
 family 9. *Plant Physiol.* 144, 1693–1696 (2007).
- 14 31. Tsutsui, H. & Higashiyama, T. pKAMA-ITACHI vectors for highly efficient
 15 CRISPR/Cas9-mediated gene knockout in *Arabidopsis thaliana*. *Plant Cell Physiol*.
 16 58, 46–56 (2017).
- 32. Gallois, P. & Marinho, P. Leaf disk transformation using *Agrobacterium tumefaciens*-expression of heterologous genes in tobacco. *Methods Mo. Biol.* 49, 39–48 (1995).
- 33. Nakagawa, T. et al. Development of series of gateway binary vectors, pGWBs,
 for realizing efficient construction of fusion genes for plant transformation. *J. biosci. bioeng.* 104, 34–41 (2007).
- 34. Clough, S. J. & Bent, A. F. Floral dip: a simplified method for
 Agrobacterium-mediated transformation of *Arabidopsis thaliana*. *Plant J*. 16, 735–
 43 (1998).
- 35. Tsutsui, H. et al. Micrografting device for testing environmental conditions for
 grafting and systemic signaling in Arabidopsis. Preprint at
 https://doi.org/10.1101/2019.12.20.885525 (2019).
- 36. Notaguchi, M., Daimon, Y., Abe, M. & Araki, T. Adaptation of a seedling
 micro-grafting technique to the study of long-distance signaling in flowering of

1 *Arabidopsis thaliana. J. Plant Res.* **122**, 201–214 (2009).

- 37. Townsley, B. T., Covington, M. F., Ichihashi, Y., Zumstein, K. & Sinha, N. R.
 Brad-Seq: Breath Adapter Directional Sequencing: A Streamlined, Ultra-Simple and
 Fast Library Preparation Protocol for Strand Specific Mrna Library Construction." *Front. Plant Sci.* 6, 366 (2015).
- 6 38. Ichihashi, Y., Fukushima, A., Shibata, A. & Shirasu. K, High impact gene discovery:
 7 Simple strand-specific mRNA library construction and differential regulatory
 8 analysis based on gene co-expression network. *Methods Mol. Biol.* 1830, 163–189
 9 (2018).
- 39. Kawamoto, T. Use of a new adhesive film for the preparation of multi-purpose
 fresh-frozen sections from hard tissues, whole-animals, insects and plants. *Arch. Histol. Cytol.* 66, 123–143 (2003).
- 40. Otagaki, S., Kawai, M., Masuta, C. & Kanazawa, A. Size and positional effects of
 promoter RNA segments on virus-induced RNA-directed DNA methylation and
 transcriptional gene silencing. *Epigenetics* 6, 681–91 (2011).
- 16

17 Acknowledgements

We thank D. Kurihara, T. Araki, K. Shiratake, S. Otagaki, S. Ishiguro, and Japan 18 19 Tobacco Inc., Japan for plant materials and A. Iwase for the RAP2.6 plasmid vector. We 20 thank T. Shinagawa, H. Fukada, A Ishiwata, R. Masuda, Y. Hakamada, M. Hattori, M. 21 Matsumoto, I. Yoshikawa, A. Yagi, A. Shibata, and A. Furuta for technical assistance, 22 and T. Akagi and Y. Hattori for discussions. This work was supported by grants from the 23 Japan Society for the Promotion of Science Grants-in-Aid for Scientific Research 24 (18KT0040, 18H03950 and 19H05361 to M.N. and 2516H06280 and 19H05364 to Y.S.), 25 the Cannon Foundation (R17-0070), the Project of the NARO Bio-oriented Technology 26 Research Advancement Institution (Research Program on Development of Innovative 27 Technology 28001A and 28001AB) to M.N., and the Japan Science and Technology 28 Agency (ERATO JPMJER1004 to T.H. and START15657559 and PRESTO15665754 to 29 M.N.).

30

1 Author Contributions

2 M.N., K.K., Y.S., and M. Niwa conceived of the research and designed experiments.

3 M.N., and K.O. performed grafting experiments. M.N., and Y. Sawai analyzed tissue

- 4 sections. M.N. collected microscopic data with Y.S.'s support. R.T. performed VIGS
- 5 experiments. Y.K. performed micrografting experiments. M.A. collected LMD samples.
- 6 K.K., R.O., and Y.I. generated RNA-Seq libraries. T.S. performed sequencing. K.K.
- 7 analyzed transcriptome data. M.N., M. Niwa, K.S., and T.H. supervised the experiments.
- 8 M.N., K.K. and M. Niwa wrote the paper.
- 9

10 **Competing interests:** Nagoya University has filed for patents regarding the following 11 topics: "Interfamily grafting technique using *Nicotiana*," inventor M.N. (patent 12 publication nos. WO 2016/06018 and JP 2014-212889); "Grafting facilitation technique 13 using cellulase," inventors M.N., K.K., R.T. and Y.K. (patent application nos. JP 14 2019-052727 and JP 2020-042379). We declare no financial conflicts of interest in 15 relation to this work.

1 Figure legends

2

3 Figure 1 *Nicotiana* established cell–cell adhesion in interfamily grafting.

4 **a**–**c**, Four weeks after grafting of the *Cm*, *Gm*, and *Nb* scion on the *Cm* stock. Scale bars, 5 10 cm. d-f, Transverse sections at graft junctions of (a-c). Dashed rectangles indicate 6 the position of insets. In the Gm/Cm interfamily graft, a necrotic layer is formed at the 7 graft interface (e), but is less developed in the Cm/Cm homograft (d) and Nb/Cm8 interfamily graft (f). Scale bars, 1 mm. g, Transmission electron micrograph (TEM) 9 near the Gm/Cm graft junction showing folding of the cell walls. Scale bar, 5 µm. h, 10 Stacked cell walls is not observed near the junction of the *Nb/At* interfamily graft. Scale 11 bar, 5 µm. i–k, TEMs for serial sections of a cell–cell boundary at the graft interface of 12 a Nb/At interfamily graft 2 weeks after grafting. Arrows indicate the thickness of the 13 cell wall between the cells. Scale bars, 1 µm. I, Phylogenetic tree showing angiosperm 14 families with which *Nicotiana* species (an arrowhead) form compatible interfamily 15 grafts (arrows). Families including major crops are indicated in red.

16

Figure 2 Transcriptomic analysis revealed conventional graft-associated gene expression in *Nicotiana* interfamily grafting.

19 **a**, Principal component analysis of the transcriptome of the *Nb* intact stem and the scion 20 of *Nb/At* interfamily grafts at five time points (three biological replicates for each time 21 points) distinguishes intact stems, wound response a short time after grafting, and the 22 graft wound healing process. PC, principal component. b, Hierarchical clustering with 23 Euclidean distance and Ward's minimum variance method over ratio of RNA-seq data 24 from five time points after Nb/At grafting against intact plants resolves nine gene 25 clusters. Genes associated with grafting reported in previous studies are marked. c, 26 Expression levels of genes associated with auxin action, wound repair, and cambium, 27 provascular xylem, and phloem development in Nb/At interfamily grafts and Nb/Nb 28 homografts. Supplementary Information Table 3 provides details. d, Extraction of 29 early-upregulated genes associated with heterograft formation. Bold line indicates 30 average of 189 genes. e, GO enrichment analysis of 189 genes shows enrichment of 31 'Extracellular region', 'Cell wall', 'Apoplast', and 'Plasmodesmata'. Genes in the four 32 categories overlap. Each numerical value represents the P-value of the GO analysis. f, 33 Expression profile of representative genes among the 189 early-upregulated genes after

grafting. g, Laser microdissection of *Nb/At* heterograft tissue was performed for the
 RNA-seq analysis. In, Vas, and Pith represent the inner central area of *Nb* scion tissue,
 and the cambial and pith area of the graft boundary for RNA extraction samples,
 respectively. h, LMD-RNA-seq of genes presented in (f) shows significant expression
 in Vas.

6

7 Figure 3 Cell wall modification involved in *Nicotiana* interfamily grafting.

8 **a**, Search for genes involved in *Nb* interfamily grafting. Of 189 upregulated genes in *Nb* 9 (Fig. 2d), 110 genes were upregulated but 79 genes were not in incompatible Gm/At 10 interfamily grafts. Expression patterns of the 110 and 79 genes in Nb/At and Gm/At are 11 shown. b, GO enrichment analysis of 79 genes shows the genes overlapping in two 12 categories, 'Extracellular region' and 'Cell wall', were mostly extracted after 13 classification in (a) and are marked in red. c, Expression profile of β -1,4-glucanase 14 (NbGH9B3) in represented samples. d, Nb/At grafts two weeks after grafting in VIGS 15 experiments (upper panels). Nb scions infected with CMV virus harboring a partial 16 sequence of NbGH9B3 to trigger gene silencing (NbGH9B3-VIGS), with no virus infection (NI) and vector control (VC) were grafted. Lower panels show transverse 17 18 sections of each graft junction. Inset indicates the intercept of At tissues separated. e, 19 Suppression of *NbGH9B3* expression by VIGS was verified by qRT-PCR. Expression 20 levels were normalized against *NbACT1* and adjusted to be relative to the NI sample. **f**, 21 Effect of the *NbGH9B3*-VIGS on graft establishment. Differences between the sample 22 groups were tested with Fisher's exact tests with α set at P < 0.05 (*) or P < 0.01 (**), n = 30 grafts for each sample fraction. g-j, Transverse sections of grafted stem sample as 23 24 represented. g, h, Optical microscopic images. Arrowheads indicate the boundary of Nb 25 and At. Scale bars, 100 µm. i, j, TEM images. Yellow and red 'P' indicate the plastid of At and Nb, respectively. * indicates a gap formed between Nb and At cells. Scale bars, 5 26 27 um. k, Effect of CRISPR knock-out (KO) of NbGH9B3 on graft establishment. The 28 effect of KO was evaluated using Fisher's exact test (P < 0.05). Graft establishment was 29 confirmed for 41 of 45 wild-type grafts and 28 of 47 KO grafts.

30

31 Figure 4 *Glycosyl hydrolase 9B3* is essential for graft wound healing in plants.

32 **a**, Phylogeny of plant glycosyl hydrolase gene family including the *GH9B3* clade.

33 Upper panel shows a tree for the *GH9B3* clade genes and lower panel shows a tree for

1 all GH clades. The number of At genes included in each clade is shown in triangles (see 2 Methods). b. GH9B3 clade genes located the same clade in as 3 Niben101Scf01184g16001 show a common expression pattern; expression of genes 4 up-regulated at an early stage after grafting is maintained when the graft is established, 5 or continues to rise subsequently, and is not maintained if the graft is not established. c, Increase in shoot fresh weight after grafting in two lines of mutants for AtCEL3, a 6 7 GH9B3 clade gene in At, and the wild type. Experiments were performed on 14–19 8 seedlings for each sample fraction. Student's *t*-tests were conducted (*P < 0.05). d. An 9 At overexpression line of NbGH9B3 using a RAP2.6 wound-inducible promoter 10 (NbGH9B3-OX) increased percentage success of grafting compared with wild-type 11 grafting. In the experiment, graft trials were performed on 64 NbGH9B3-OX and 102 12 wild-type seedlings. Viability of the scion was determined two weeks after grafting and 13 the effect of overexpression was evaluated by Fishers' exact test (P < 0.05). e-g Grafts 14 of tomato scion onto At (e; 3 weeks after grafting, f; 4 months after grafting) or Cm (g; 3 15 months after grafting) using a Nb interscion. Arrowheads indicate grafted points. Scale 16 bars, 1 cm. h, Proposed method to perform interfamily grafting mediated by a Nicotiana interscion. 17



Figure 1 Nicotiana established cell-cell adhesion in interfamily grafting.

a–**c**, Four weeks after grafting of the *Cm*, *Gm*, and *Nb* scion on the *Cm* stock. Scale bars, 10 cm. **d**–**f**, Transverse sections at graft junctions of (**a**–**c**). Dashed rectangles indicate the position of insets. In the *Gm/Cm* interfamily graft, a necrotic layer is formed at the graft interface (**e**), but is less developed in the *Cm/Cm* homograft (**d**) and *Nb/Cm* interfamily graft (**f**). Scale bars, 1 mm. **g**, Transmission electron micrograph (TEM) near the *Gm/Cm* graft junction showing folding of the cell walls. Scale bar, 5 μ m. **h**, Stacked cell walls is not observed near the junction of the *Nb/At* interfamily graft. Scale bar, 5 μ m. **i**–**k**, TEMs for serial sections of a cell–cell boundary at the graft interface of a *Nb/At* interfamily graft 2 weeks after grafting. Arrows indicate the thickness of the cell wall between the cells. Scale bars, 1 μ m. **l**, Phylogenetic tree showing angiosperm families with which *Nicotiana* species (an arrowhead) form compatible interfamily grafts (arrows). Families including major crops are indicated in red.



Figure 2 Transcriptomic analysis revealed conventional graft-associated gene expression in *Nicotiana* interfamily grafting.

a, Principal component analysis of the transcriptome of the *Nb* intact stem and the scion of *Nb/At* interfamily grafts at five time points (three biological replicates for each time points) distinguishes intact stems, wound response a short time after grafting, and the graft wound healing process. PC, principal component. **b**, Hierarchical clustering with Euclidean distance and Ward's minimum variance method over ratio of RNA-seq data from five time points after *Nb/At* grafting against intact plants resolves nine gene clusters. Genes associated with grafting reported in previous studies are marked. **c**, Expression levels of genes associated with auxin action, wound repair, and cambium, provascular xylem, and phloem development in *Nb/At* interfamily grafts and *Nb/Nb* homografts. Supplementary Information Table 3 provides details. **d**, Extraction of early-upregulated genes associated with heterograft formation. Bold line indicates average of 189 genes. **e**, GO enrichment analysis of 189 genes shows enrichment of 'Extracellular region', 'Cell wall', 'Apoplast', and 'Plasmodesmata'. Genes in the four categories overlap. Each numerical value represents the *P*-value of the GO analysis. **f**, Expression profile of representative genes among the 189 early-upregulated genes after grafting. **g**, Laser microdissection of *Nb/At* heterograft tissue was performed for the RNA-seq analysis. In, Vas, and Pith represent the inner central area of *Nb* scion tissue, and the cambial and pith area of the graft boundary for RNA extraction samples, respectively. **h**, LMD-RNA-seq of genes presented in (**f**) shows significant expression in Vas.



Figure 3 Cell wall modification involved in Nicotiana interfamily grafting.

a, Search for genes involved in Nb interfamily grafting. Of 189 upregulated genes in Nb (Fig. 2d), 110 genes were upregulated but 79 genes were not in incompatible Gm/At interfamily grafts. Expression patterns of the 110 and 79 genes in Nb/At and Gm/At are shown. **b**, GO enrichment analysis of 79 genes shows the genes overlapping in two categories, 'Extracellular region' and 'Cell wall', were mostly extracted after classification in (a) and are marked in red. c, Expression profile of β -1,4-glucanase (NbGH9B3) in represented samples. **d**, Nb/At grafts two weeks after grafting in VIGS experiments (upper panels). Nb scions infected with CMV virus harboring a partial sequence of NbGH9B3 to trigger gene silencing (NbGH9B3-VIGS), with no virus infection (NI) and vector control (VC) were grafted. Lower panels show transverse sections of each graft junction. Inset indicates the intercept of At tissues separated. e, Suppression of NbGH9B3 expression by VIGS was verified by qRT-PCR. Expression levels were normalized against NbACT1 and adjusted to be relative to the NI sample. f, Effect of the NbGH9B3-VIGS on graft establishment. Differences between the sample groups were tested with Fisher's exact tests with α set at P < 0.05 (*) or P < 0.01 (**), n = 30 grafts for each sample fraction. g-j, Transverse sections of grafted stem sample as represented. g, h, Optical microscopic images. Arrowheads indicate the boundary of Nb and At. Scale bars, 100 µm. i, j, TEM images. Yellow and red 'P' indicate the plastid of At and Nb, respectively. * indicates a gap formed between Nb and At cells. Scale bars, 5 µm. k, Effect of CRISPR knock-out (KO) of *NbGH9B3* on graft establishment. The effect of KO was evaluated using Fisher's exact test ($P \le 0.05$). Graft establishment was confirmed for 41 of 45 wild-type grafts and 28 of 47 KO grafts.



Figure 4 Glycosyl hydrolase 9B3 is essential for graft wound healing in plants.

a, Phylogeny of plant glycosyl hydrolase gene family including the *GH9B3* clade. Upper panel shows a tree for the *GH9B3* clade genes and lower panel shows a tree for all *GH* clades. The number of *At* genes included in each clade is shown in triangles (see Methods). **b**, *GH9B3* clade genes located in the same clade as *Niben101Scf01184g16001* show a common expression pattern; expression of genes up-regulated at an early stage after grafting is maintained when the graft is established, or continues to rise subsequently, and is not maintained if the graft is not established. **c**, Increase in shoot fresh weight after grafting in two lines of mutants for *AtCEL3*, a *GH9B3* clade gene in *At*, and the wild type. Experiments were performed on 14–19 seedlings for each sample fraction. Student's *t*-tests were conducted (*P < 0.05). **d**, An *At* overexpression line of *NbGH9B3* using a RAP2.6 wound-inducible promoter (*NbGH9B3*-OX) increased percentage success of grafting compared with wild-type grafting. In the experiment, graft trials were performed on 64 *NbGH9B3*-OX and 102 wild-type seedlings. Viability of the scion was determined two weeks after grafting and the effect of overexpression was evaluated by Fishers' exact test (P < 0.05). **e**-**g** Grafts of tomato scion onto *At* (**e**; 3 weeks after grafting) or *Cm* (**g**; 3 months after grafting) using a *Nb* interscion. Arrowheads indicate grafted points. Scale bars, 1 cm. **h**, Proposed method to perform interfamily grafting mediated by a *Nicotiana* interscion.