Protection of Human Colon Cells from Shiga Toxin by Plant-based Recombinant Secretory IgA

Katsuhiro Nakanishi1, Shotaro Morikane1, Shiori Ichikawa1, Kohta Kurohane1, Yasuo Niwa2, Yoshihiro Akimoto3, Sachie Matsubara4, Hayato Kawakami3, Hirokazu Kobayashi2 & Yasuyuki Imai2

Shiga toxin is a major virulence factor of food-poisoning caused by Escherichia coli such as O157:H7. Secretory immunoglobulin (Ig) A (SIgA) is supposed to prevent infection of the mucosal surface and is a candidate agent for oral immunotherapy. We previously established a recombinant monoclonal antibody (mAb) consisting of variable regions from a mouse IgG mAb specific for the binding subunit of Shiga toxin 1 (Stx1) and the Fc region of mouse IgA. Here we produced a secretory form of the recombinant IgA (S-hyIgA) with transgenic Arabidopsis thaliana plant. All the S-hyIgA cDNAs (heavy, light, J chain and secretory component) were expressed under the control of a bidirectional promoter of a chlorophyll a/b-binding protein of A. thaliana without using a viral promoter. The plant-based S-hyIgA exhibited antigen binding, and was modified with plant-specific N-linked sugar chains. The Ig heavy chain and secretory components were observed in an intracellular protein body-like structure of the transgenic leaves on immuno-electron microscopy. An extract of the transgenic leaves neutralized the cytotoxicity of Stx1 toward butyrate-treated Caco-2 cells, a human colon carcinoma cell line. These results will contribute to the development of edible therapeutic antibodies such as those for the treatment of mucosal infection.

The intestinal mucosal surface is always exposed to many types of antigens derived from food, exogenous microbes and enteric bacteria. The invasion and potential harmful effects of these antigens on the body are blocked by mucosal immune systems including immunoglobulin A (IgA)1. IgA is the major antibody isotype on the mucosal surface, and it forms secretory IgA (SIgA). Two IgA monomers are linked through a joining (J) chain (termed dimeric IgA), followed by association with a secretory component (SC) that yields SIgA. SIgA is produced by two types of cells, IgA-secreting plasma cells and mucosal epithelial cells2. Plasma cells produce dimeric IgA and secrete it into the lamina propria. After binding to the polymeric immunoglobulin receptors (plgR) on the epithelial cell membrane, the secreted dimeric IgA is transcytosed from the basolateral to the apical side. The extracellular domains of plgR are cleaved by a protease to form SC. The dimeric IgA and SC complex is released into the intestinal lumen as SIgA. SIgA can function in a protease-rich environment such as the interior of the intestine because it is resistant to proteases3,4. SIgA is also important for protection of children from infection on oral feeding of breast milk containing SIgA5. In a porcine system, orally fed SIgA, but not IgG, was shown to protect piglets from infection of enterotoxigenic Escherichia coli (ETEC)6. Thus, SIgA specific for antigens on pathogens will be useful for treating mucosal infections based on the mechanism of oral passive immunity.

It is difficult to produce monoclonal SIgA using single mammalian cells because two types of cells are involved in nature. Although there have been several reports, the production of SIgA in a single mammalian cell is still challenging7,8. The first report of SIgA production using a plant expression system was earlier than that as to a mammalian cell system, and such reports on plant systems have recently been increasing7–12. A plant expression system has advantages as to oral passive immunity due to the ability of oral administration without or with minimal formulation when edible plant hosts are used13. In addition, such a system is more cost effective and exhibits

1Laboratory of Microbiology and Immunology, School of Pharmaceutical Sciences, University of Shizuoka, Shizuoka City, Shizuoka 422-8526, Japan. 2Laboratory of Plant Molecular Improvement, Graduate Division of Nutritional and Environmental Sciences, University of Shizuoka, Shizuoka City, Shizuoka 422-8526, Japan. 3Department of Anatomy, Kyorin University School of Medicine, Mitaka, Tokyo 181-8612, Japan. 4Laboratory for Electron Microscopy, Kyorin University School of Medicine, Mitaka, Tokyo 181-8612, Japan. Correspondence and requests for materials should be addressed to Y.I. (email: imai@u-shizuoka-ken.ac.jp)
higher scalability than mammalian ones for the production of therapeutic proteins\textsuperscript{14,15}. Antibodies produced in a plant system are called plantibodies for short\textsuperscript{14}. Thus, plant expression systems will be valuable for the production of SlgA and other agents aiming at oral passive immunity.

We previously established recombinant IgA antibodies aiming at oral passive immunization against Shiga toxin 1 (Stx1)\textsuperscript{16}. Stx1 is responsible for food-poisoning caused by enterohemorrhagic \textit{Escherichia coli} and \textit{Shigella dysenteriae}\textsuperscript{17}. Stx1 is an AB\textsubscript{5} toxin consisting of a cytotoxic A subunit and a pentamer of cell-binding B subunits (StxB)\textsuperscript{18}. StxB1 binds to the target cells via a cell surface glycolipid, globotriaosylceramide (Gb\textsubscript{3}). Gb\textsubscript{3} is expressed on kidney, microvascular, leukocytic and neuronal cells, as well as many types of tumour cells\textsuperscript{19-21}. Butyrate treatment of some cancer cell lines has been reported to induce expression of Gb\textsubscript{3}\textsuperscript{17}. Because the cytotoxicity of Stx1 starts with the binding of StxB to cell surface Gb\textsubscript{3}, inhibition of the binding of StxB to Gb\textsubscript{3} will be effective for preventing Stx1 toxicity. We established a recombinant antibody, termed hybrid-IgG/IgA (hyIgA for short), which has an amino acid sequence derived from a variable to the hinge region of the heavy chain as well as the light chain of mouse monoclonal IgG specific for Stx1B and the Fc region of the mouse IgA heavy chain. The dimeric hybrid-IgG/IgA expressed in Chinese hamster ovary cells together with mouse J chain efficiently neutralized Stx1 toxicity \textit{in vitro}. The neutralizing activity of the dimeric one was stronger than that of the recombinant monomeric IgA and the original IgG mAb\textsuperscript{22,23}. We then established transgenic \textit{Arabidopsis thaliana} expressing the \textit{H} and \textit{L} chain genes is controlled by a chlorophyll \textit{a/b}-binding protein (CAB) promoter derived from \textit{A. thaliana}, whereas the \textit{J} chain gene is controlled by the cauliflower mosaic virus 35S promoter. The CAB promoter is known to induce the co-expression of two proteins bi-directionally, and the expression level was reported to be high in leaf tissue\textsuperscript{25}. In this study, we developed hybrid-IgG/IgA transgenic plants that express a secretory form of hybrid-IgG/IgA (S-hyIgA) only using the CAB promoter and terminators through a one-step transformation of four genes in a construct. We examined the protein assembly of the secretory form of IgA in leaf tissues and its neutralizing activity against Stx1 \textit{in vitro}.

**Results**

**Establishment of secretory hybrid-IgG/IgA transgenic \textit{Arabidopsis thaliana}.** A binary vector was constructed for the expression of a plant-based monoclonal secretory hybrid-IgG/IgA (S-hyIgA) specific for Stx1. S-hyIgA consists of immunoglobulin heavy and light chains, a joining (J) chain, as well as a secretory component (SC), which is a product of mucosal epithelial cells. The S-hyIgA expression vector is formed from the pBCH1 binary vector\textsuperscript{26} harbouring the hybrid-IgG/IgA heavy (H) and light (L) chain expression cassette, and the J chain and SC expression cassette (Fig. 1a). The \textit{H} chain, \textit{L} chain, \textit{J} chain and SC genes were expressed under the control of a bidirectional CAB promoter and terminator derived from \textit{A. thaliana} (\textit{P}_{CAB}, \textit{T}_{CAB1} and \textit{T}_{CAB2}). The S-hyIgA expression binary vector was introduced into \textit{A. thaliana} through \textit{Agrobacterium}-mediated transformation. After selection with hygromycin B, we obtained 12 transgenic lines. Leaf extracts of the transgenic lines were screened for the IgA protein expression level by means of sandwich ELISA, in which anti-\textit{κ} was used for capture and anti-\textit{α} for detection. IgA levels in each leaf extract were compared by the absorbance readings relative to that of the
Figure 2. **Expression of S-hyIgA proteins in A. thaliana.** (a,b) Detection of assembled hybrid-IgG/IgA by sandwich ELISA. The hybrid-IgG/IgA proteins in each sample were captured with immobilized anti-κ antibodies and detected with anti-α (a) or anti-plgR (b) antibodies. Crude leaf extracts and 50%-saturated ammonium sulphate (AmS)-concentrated ones were compared. (c) Detection of SC protein by sandwich ELISA. The SC protein in the crude leaf extracts was captured with goat anti-plgR and detected with rabbit anti-plgR antibodies. The symbols represent crude extracts from secretory Tg (open circles) or WT plants (open triangles), or represent AmS-concentrated ones from secretory Tg (open diamonds) or WT plants.
(open squares). Data are expressed as means of triplicate determinations. All error bars (SD) are shorter than the size of symbols. The results are representative of four experiments. (d,e) Immunoblot analysis of S-hyIgA expressed in transgenic A. thaliana leaves. Leaf proteins precipitated with 50%-saturated ammonium sulphate were separated by SDS-PAGE under reducing (d; 12% gel, 15μg protein/lane) or non-reducing (e; 4–20% gradient gel, 5μg protein/lane) conditions, and then blotted onto a PVDF membrane. Each polypeptide chain of secretory IgA was detected with anti-α, anti-κ, anti-ι and anti-plgR antibodies. Secretory Tg (quadruple transgenic for H, L, J and SC), dimer Tg (triple transgenic for H, L and J), and WT (wild-type) A. thaliana, and mouse saliva (secretory IgA positive control) were analysed.

Expression of S-hyIgA proteins in transgenic A. thaliana plant leaves. Total soluble proteins (TSP) were extracted from secretory Tg leaves and then hybrid-IgG/IgA protein in the extract was quantitated by ELISA (Fig. 2a). The hybrid-IgG/IgA was captured with an immobilized anti-κ antibody, followed by detection with an anti-α antibody. The signals representing antibodies with both H and L chains increased with increasing TSP in the crude extract from secretory Tg leaves (open circles). In contrast, no signal was detected for wild type leaves (open triangles). The hybrid-IgG/IgA concentration was calculated by comparison with IgA myeloma TEPC 15 as a standard. The production volume of assembled hybrid-IgG/IgA reached 8.0μg/g leaf tissue (0.07% of TSP). Because SC consists of the extracellular domains of plgR, we used anti-plgR antibodies to detect SC. The S-hyIgA in the secretory Tg leaf extract was detectable with anti-plgR antibodies (Fig. 2b). The S-hyIgA-specific signals also increased with increasing TSP in the secretory Tg sample (open circles). No such signal was detected for the wild type leaves (open triangles). To measure the total SC concentration, sandwich ELISA was performed using goat anti-plgR as a capture antibody and rabbit anti-plgR as a detection antibody. The signals representing SC increased with increasing TSP from secretory Tg leaves (Fig. 2c). Compared with a standard curve generated with recombinant plgR, the total SC concentration was calculated. The production volume of the total SC reached 57.7μg/g leaf tissue (0.31% of TSP).

Proteins were concentrated from the leaf extracts by ammonium sulphate (AmS) precipitation. When concentrated leaf samples were examined (open diamonds), approximately two-fold enrichment was observed for both IgA (Fig. 2a) and SlgA (Fig. 2b) by comparing with the crude extract (open circles). No signal was detected in the wild type leaf concentrate (open squares).

The presence of each component of S-hyIgA in the concentrate was investigated by SDS-PAGE and immunoblotting using specific antibodies. In this experiment, we also performed comparison with the dimer hybrid-IgG/IgA transgenic A. thaliana (dimer Tg) expressing H, L and J chains but no SC, which had been established previously25. Under reducing conditions, α, κ and J chains were detected in the secretory Tg as well as in the dimer Tg, whereas SC was only detected in the secretory Tg (Fig. 2d, arrowheads). The bands representing the α chain and SC were observed around 50 kDa and 70 kDa, respectively, for the transgenic plants. In contrast, secretory IgA in mouse saliva exhibited values of 60 kDa and 90 kDa, respectively, for the α chain and SC. Some additional bands were observed by the anti-ι chain (around 30 kDa) and anti-κ chain (under 20 kDa) in the secretory Tg and dimer Tg leaves, but we did not carry out further identification of these bands.

Under non-reducing conditions, bands corresponding to the assembled S-hyIgA were detected with antibodies specific for each immunoglobulin chain in the secretory Tg leaves (Fig. 2e, arrows). These bands corresponded to smaller molecular masses than that of secretory IgA from mouse saliva. Some other bands were seen for the secretory Tg sample. These bands may represent the H chain dimer with J chain and an SC, free SC and their fragments. The secretory Tg but not the dimer Tg sample gave bands detectable with anti-plgR. These results indicate that S-hyIgA is expressed and assembled in the secretory Tg plant leaves.

Subcellular localization of hybrid-IgG/IgA in leaf tissue of transgenic A. thaliana. To determine the intracellular localization of S-hyIgA in plant cells, we prepared ultrathin sections of leaf tissue and analysed them by means of immuno-electron microscopy using antibody-conjugated colloidal gold particles (Fig. 3). A protein body-like structure was observed in the transgenic plant cells (arrows). Co-localization of the colloidal gold-labelling signals representing the α chain (18 nm-particle, open arrowhead) as well as SC (12 nm-particle, closed arrowhead) was observed within the protein body-like structure. The diameter of the protein body-like structures approximately ranged from 1 to 6μm. Such a structure was not seen in the wild type plant cells. In the isotype controls, colloidal gold-labelling signals were scarcely observed in the protein body-like structure. Glycosylation of plantibodies. Protein bodies are classified into endoplasmic reticulum (ER)-derived protein body-I (PB-I) and protein storage vacuole protein body-II (PSV or PB-II)28. Glycoproteins accumulated in the PSV are believed to have undergone sugar chain processing in the Golgi apparatus in plant cells29. Thus, it is...
useful to determine whether S-hylgA has undergone plant-specific modification of sugar chains for classifying S-hylgA accumulated in the protein body-like structure. Two types of modifications associated with the core of N-glycans were examined. One was Fuc α(1,3) GlcNAc and the other xylose β(1,2) Man. Plantibodies were captured with immobilized anti-κ antibodies and then detected with appropriate detection antibodies on ELISA. Because of the relatively high background for plant samples due to non-specific adsorption on ELISA wells, the signals from wells with BSA blocking only were subtracted from those from wells coated with anti-κ antibodies. Successful capture of S-hylgA was demonstrated by detection with anti-α antibodies. Samples from secretory Tg as well as mouse IgA myeloma TEPC 15 proteins produced α chain-specific signals, whereas samples from wild type plants did not. Samples from secretory Tg leaves exhibited the binding of antibodies specific for α1,3-linked Fuc as well as one for β1,2-linked Xyl (Fig. 4). TEPC 15 did not show plant-specific modifications, consistent with

**Figure 3.** Immuno-electron microscopic analysis of leaf tissue of transgenic A. thaliana. Transverse sections of A. thaliana leaf tissue were stained with anti-α chain (18-nm gold label, open arrowhead) or anti-pIgR (12-nm gold label, closed arrowhead) antibodies. Arrows indicate intracellular protein body-like structures containing S-hylgA. V, vacuole; Chl, chloroplast; CW, cell wall. Bars represent 1 μm (low magnification) or 500 nm (high magnification).
it being a mouse protein. In wild type plants, signals of plant-type glycosylation remained, but the signal levels were lower than those for transgenic plants. We concluded that the S-hyIgA was equipped with plant-type sugar chains, because the signals for plant-specific glycosylation were shown to be dependent on anti-κ capture.

The S-hyIgA binds to Stx1B. We then investigated whether the S-hyIgA plantibodies bound to Stx1B by means of ELISA. Anti-α as well as anti-κ second antibodies revealed dose dependent-binding of the plantibodies from the secretory Tg as well as dimer Tg leaves (Fig. 5a and b). Anti-pIgR antibodies only detected the binding of the secretory plantibodies from the secretory Tg leaves, while no signal was seen for the dimer Tg leaves (Fig. 5c). Stx1B-specific antibody binding was not observed using wild type leaf samples or TEPC 15. These results demonstrated that Stx1B-specific S-hyIgA is produced by the secretory Tg but not by dimer Tg leaves.

Neutralization of Stx1-mediated cytotoxicity by plantibody. Toxin neutralization by plantibodies was then examined. We first used a human colonic epithelial cell line, Caco-2, to assess Stx1-induced cytotoxicity. We think that direct cytotoxicity of Stx1 toward colonic epithelial cells may initiate food poisoning. Caco-2 cells are known to upregulate cell surface expression of Gb3, the target glycolipid for Stx1, on treatment with butyrate22. Under our current experimental conditions, Caco-2 cells became susceptible to Stx1 after butyrate treatment for 4 days. Thus, the viability of butyrate-treated Caco-2 cells decreased after 48 h exposure to Stx1 in a dose-dependent manner, whereas that of untreated Caco-2 cells did not (Fig. 6).

Butyrate-treated Caco-2 cells were used as a target to determine Stx1 neutralization. The reduced viability caused by exposure to 100 pg/mL Stx1 was reversed in the presence of increasing concentrations of the plantibody sample from secretory Tg or dimer Tg leaves (Fig. 7a). The reduced viability was not reversed in the presence of a wild type leaf sample or TEPC 15.

Because Vero cells are highly sensitive to Stx1 toxicity, we used them as another target to determine Stx1 neutralization. The reduced viability caused by exposure to 10 pg/mL Stx1 was reversed in the presence of increasing concentrations of a sample from secretory Tg or dimer Tg leaves, but not from wild type plants (Fig. 7b).

Because Stx1 is known to induce apoptosis of sensitive cells9,10, Stx1-induced apoptosis of butyrate-treated Caco-2 cells was assessed by means of DNA fragmentation assay. The genomic DNA of Caco-2 cells became fragmented into length of multiples of 180 bp after 48 h exposure to 100 pg/mL Stx1 (Fig. 7c, lane 2). High molecular weight DNA did not enter the agarose gel, and was not visible for the Stx1-untreated cells (lane 1). When Stx1 was pre-incubated with plantibodies, the DNA fragmentation of Caco-2 cells was inhibited (lanes 5 to 8). One μg/mL of plantibodies from secretory Tg (lane 6) or dimer Tg (lane 8) leaves completely prevented the DNA fragmentation induced by 100 pg/mL Stx1. Samples from wild-type leaves and TEPC 15 did not prevent DNA fragmentation (lanes 3 and 4).

In conclusion, a functional secretory IgA can be produced in transgenic A. thaliana using a promoter of plant origin. The in vitro neutralization capacities were similar between samples from secretory Tg and dimer Tg plants on the basis of the IgA concentrations.

Discussion
Plant expression systems are useful tools for the production of oral immunotherapeutic agents including SlgA because of easy administration, i.e., eating edible host plants13. We previously established a hybrid-IgG/IgA consisting of mouse monoclonal IgG against Stx1B and mouse IgA constant regions14. To produce SlgA that can neutralize Stx1 toxicity, we established transgenic plants expressing S-hyIgA by means of the Agrobacterium-method.
Viral promoters such as the cauliflower mosaic virus 35S promoter are often used in plant expression system. However, it will be useful to test whether an alternative expression system works without using virus-derived DNA elements with respect to public acceptance. We chose the Arabidopsis-derived CAB promoter to express S-hyIgA (Fig. 1a). The CAB promoter is known to allow a high level of expression as well as bi-directional transcription. Thus, this promoter is suitable for the co-expression of two proteins in leaf tissue. Regarding the protein expression levels, the SC protein was expressed at a high level (57.7 μg/g leaf tissue). This expression level is comparable to that in other reports describing stable expression of therapeutic proteins in plants. On the other hand, the expression level of assembled hybrid-IgG/IgA that has both heavy and light chains was 8.0 μg/g leaf tissue, which was lower than in other reports for stable expression of IgA in plants. Proteolytic degradation of heavy chains is considered as one of the reasons why the hybrid-IgG/IgA was expressed at a low level in contrast to the high level of SC expression. On western blot analysis, a large amount of degraded heavy chains was observed around 20–30 kDa (Fig. 2d). Some reports described that the hinge region of a human IgG plantibody has a weak spot as to proteolysis in plant cells. The hinge region of the hybrid-IgG/IgA is derived from that of mouse IgG, and the amino acid sequence around the weak spot is similar to that reported, that is KDVKKIVP in our construct and KDVKKVEP in other reports. Replacement of the hinge region may improve hybrid-IgG/IgA expression in plant cells.

Figure 5. Binding of S-hyIgA to immobilized Stx1B. Leaf extract samples (50%-saturated ammonium sulphate precipitation fraction) from secretory Tg (open circles), dimer Tg (open squares), or wild-type (open triangles) plants, or purified mouse IgA myeloma protein TEPC 15 (open diamonds) were allowed to bind to immobilized Stx1B. The bound antibodies were detected with anti-α (a), anti-κ (b), or anti-pIgR (c) antibodies. Data are expressed as means ± SD of triplicate determinations. Error bars underneath the symbols are not visible. For comparison, the results for secretory Tg and dimer Tg leaves, and TEPC 15 are plotted against IgA concentration, while those for secretory Tg and wild-type leaves against TSP, on the abscissa. The results are representative of three experiments.

Figure 6. Butyrate treatment increased the sensitivity of Caco-2 cells to Stx1. Caco-2 cells were cultured with 2 mM sodium butyrate-containing medium or normal medium for 4 days. The cells were then exposed to various concentrations of Stx1 (0, 10, 30, 100 or 300 pg/mL) for 48 h. The viable cells were quantitated by means of WST-8 cell viability assaying. Data for butyrate-treated cells (open circles) and untreated cells (open triangles) are shown. Viability was calculated as the relative value to that without toxin exposure. Data are expressed as means ± SD of triplicate determinations. Error bars underneath the symbols are not visible.
Intracellular localization of S-hylIgA is a key factor as to protein yield and stability. Translated S-hylIgA was accumulated in a protein body-like structure, which was observed only in transgenic plants (Fig. 3). A protein body-like structure was also reported in other transgenic plant leaves. Plant-specific glycosylation on S-hylIgA (Fig. 4) suggested that S-hylIgA was accumulated in a PSV-like vacuole. Storage proteins in PSV are known to be transported from the Golgi apparatus, in which plant-specific glycosylation occurs. PSV localization of plant-expressed IgA was also reported in rice. PSV functions to supply amino acids for plant seed development through degradation of storage proteins. Thus, a high rate of IgA degradation may be due to the localization in this structure. Modification of intracellular localization of S-hylIgA may increase its yield.

The EHEC infection is difficult to treat because a large amount of Stx is released from bacteria upon treatment with conventional antibiotics. Oral passive immunotherapy is a candidate treatment for EHEC infection as an alternative one to antibiotics. Despite that life-threatening symptoms are dependent on the entry of Stx into the blood circulation, it is still controversial as to how Stx initially enters the blood.

It is natural to consider that Stx enters due to a breach in the intestinal epithelial barrier. However, it is not clear whether Stx-induced damage to intestinal epithelial cells is responsible for the breach. One hypothesis is as follows: local inflammation due to the EHEC infection induces transmigration of neutrophils through the intestinal epithelium. The neutrophil transmigration triggers defects in the intestinal barrier that allow passive transfer of Stx into the bloodstream. A monolayer of human colon carcinoma T84 cells was used to demonstrate neutrophil transmigration in the presence of inflammatory cytokines. T84 cells lack Gb3, and are resistant to the direct cytotoxicity of Stx. Despite this, the translocation of Stx was reported to be enhanced with neutrophil transmigration using a polarized T84 monolayer system. In a mouse system, we demonstrated that the binding sites for Stx1B are restricted to the distal part of the colon.

With this background, we used human colon epithelial Caco-2 cells to measure Stx1 cytotoxicity in this study. The cytotoxicity was enhanced after butyrate treatment (Fig. 6), which is consistent with an increase in Gb3 on Caco-2 cells. The leaf samples from S-hylIgA transgenic plants inhibited Stx1 cytotoxicity not only against Vero cells but also against butyrate-treated Caco-2 cells (Fig. 7). Stx1-induced apoptosis of Caco-2 cells was also inhibited by the leaf samples from S-hylIgA transgenic plants. These results indicated that Stx1 is cytotoxic to human colon epithelial cells under certain circumstances. Stx1-sensitive human colon cells were protected by a plantibody specific for Stx1B. We already observed that the current construct is applicable to the expression of S-hylIgA in lettuce leaves (manuscript in preparation). Continuous efforts to increase production volume will provide a key for success toward practical edible IgA therapeutics.

We have previously established A. thaliana producing dimeric hyIgA where IgA heavy and light chains are produced under a CAB promoter while J chain is expressed under a 35S promoter. In contrast, the current S-hylIgA is expressed only using two CAB promoters in the absence of 35S promoter. Despite this difference, the production volume of S-hylIgA was 8.0 μg/g fresh weight and that of dimeric hyIgA was 11 μg/g. Although...
and SC proteins, a Jc-SC expression cassette was constructed as follows: mouse J chain cDNA was amplified by PCR using J chain-specific primers named JCF15 and JCR-15, and subcloned J chain cDNA was digested with Xho I and ligated into the pGEM5zf vector (Invitrogen, Carlsbad, CA, USA). The subcloned J chain cDNA was digested with Xhol and ligated into the pGEM5zf vector (Promega, Madison, WI, USA) harbouring chlorophyll a/b-binding protein terminator (TcAB)25, resulting in the production of pGEM5zf/Jc-TcAB. To subclone mouse SC cDNA, PCR was performed using pcDNA3.1/mouse SC45 as a template, and SC-specific SCF-Xho and SCR-Xba as primers. Following XbaI/Xhol digestion and generation of blunt ends, SC cDNA was inserted into EcoRV-digested pGEM5zf harboured TcAB (pGEM5zf/TcAB), resulting in the production of pGEM5zf/SC-TcAB. pUC19/Jc-TcAB and pGEM5zf/SC-TcAB were digested with SacI and HindIII. The resulting DNA fragments, Jc-TcAB and SC-TcAB, were ligated with HindIII-digested pUC19 (Invitrogen) by means of three-piece ligation, resulting in the production of pUC19/Jc-SC-TcAB, pUC19/TcAB/Jc-SC-TcAB, and pUC19/TcAB/SC-TcAB, which were transformed into A. thaliana genomes, PCR was performed as previously described23 with the gene-specific primer sets (Table 1). 

Table 1. Primers used for PCR reactions.

| Gene            | Name of primers | DNA sequence                     |
|-----------------|-----------------|----------------------------------|
| J chain         | JCF15           | 5′-CCTTTCTAGAGTGAACAGCAAGATA-3′  |
| J chain         | JCR-Xho         | 5′-CTAGTCCAGAGTGAACAGCAAGATA-3′  |
| SC              | SCF-Xho         | 5′-CCCTCTAGAGTCTCTTATAGTGCAAAAGGC-3′ |
| SC              | SCR-Xba         | 5′-CTAAATCTAGAGAAGATGAGGCTTACT-3′ |
| IgG H chain     | IgG Heavy 2F    | 5′-CTTAATCTAGAGAAGATGAGGCTTACT-3′ |
| IgG L chain     | IgG Light 2R    | 5′-CTTAATCTAGAGAAGATGAGGCTTACT-3′ |
| ACTIN2          | actin2-F        | 5′-GTTGGTGATGAAGCACAATCCAAG-3′   |
| ACTIN2          | actin2-R        | 5′-CTGGAACAAAGACTCTGCGGATC-3′    |

Materials and Methods

Construction of a secretory hybrid-IgG/IgA expression vector. We previously constructed a hybrid-IgG/IgA expression cassette for the expression of antibody heavy and light chains in plants25. To express J chain and SC proteins, a Jc-SC expression cassette was constructed as follows: mouse J chain cDNA was amplified by PCR using J chain-specific primers named JCF15 and JCR-15 (Table 1). The amplified cDNA was digested with XbaI and subcloned into the XbaI/EcoRV-digested pcDNA3.1 vector (Invitrogen, Carlsbad, CA, USA). The subcloned J chain cDNA was digested with XbaI/Xhol and ligated into the pGEM5zf vector (Invitrogen, Madison, WI, USA) harbouring chlorophyll a/b-binding protein terminator (TcAB)25, resulting in the production of pGEM5zf/Jc-TcAB. To subclone mouse SC cDNA, PCR was performed using pcDNA3.1/mouse SC45 as a template, and SC-specific SCF-Xho and SCR-Xba as primers. Following XbaI/Xhol digestion and generation of blunt ends, SC cDNA was inserted into EcoRV-digested pGEM5zf harboured TcAB (pGEM5zf/TcAB), resulting in the production of pGEM5zf/SC-TcAB. pUC19/Jc-TcAB and pGEM5zf/SC-TcAB were digested with SacI and HindIII. The resulting DNA fragments, Jc-TcAB and SC-TcAB, were ligated with HindIII-digested pUC19 (Invitrogen) by means of three-piece ligation, resulting in the production of pUC19/Jc-SC-TcAB, pUC19/TcAB/Jc-SC-TcAB, and pUC19/TcAB/SC-TcAB, which were transformed into A. thaliana genomes, PCR was performed as previously described23 with the gene-specific primer sets (Table 1).

Transformation of Arabidopsis thaliana. Transformation and selection of A. thaliana was performed as previously described25. As a binary vector, pBCH1/secretory hybrid-IgG/IgA was used. Homozygous transformants were obtained by self-pollination.

DNA analysis of transgenic A. thaliana by PCR. To confirm the S-hylga transgenes in the transgenic A. thaliana genomes, PCR was performed as previously described23 with the gene-specific primer sets (Table 1).

mRNA analysis of transgenic A. thaliana by RT-PCR. Total RNA was purified from the transgenic plant (4-wk old) leaves using an RNeasy Mini Kit (QIAGEN, Hilden, Germany). Purified RNA samples were...
analysed by RT-PCR using an AccessQuick RT-PCR System (Promega) and the same gene-specific primer sets as for DNA analysis.

**Protein extraction and ammonium sulphate precipitation.** Leaves from the transgenic plants (4-wk old) were frozen and crushed in liquid nitrogen. A protein extraction buffer (50 mM acetate buffer [pH 5.0], 500 mM NaCl, 0.5 mM EDTA, protease inhibitor cocktail for plant cell and tissue extracts [Sigma-Aldrich, St. Louis, MO, USA]) was added to each leaf sample. The leaf tissue suspension was centrifuged for 10 min at 15,000 × g at 4 °C. The supernatant was collected and neutralized with 1 M Tris–HCl (pH 8.0). After centrifugation, ammonium sulphate was added to the supernatant to 50% saturation on ice, and proteins were allowed to precipitate overnight at 4 °C. A sample was centrifuged for 10 min at 15,000 × g at 4 °C, and the precipitate was re-dissolved in PBS. Remaining insoluble aggregates were removed by centrifugation for 10 min at 21,500 × g at 4 °C. The supernatant was dialyzed against PBS with Mini Dialysis Kit 8 kDa cut-off (GE Healthcare, Buckinghamshire, UK). The resulting sample was used as the 50%-saturated ammonium sulphate precipitated fraction.

**ELISA.** To determine the hybrid-IgG/IgA in samples, a sandwich ELISA was performed as previously described16. To measure total hybrid-IgG/IgA, goat anti-mouse κ (1 μg/mL; SouthernBiotech, Birmingham, AL, USA) was used as a capture antibody and horseradish peroxidase (HRP)-goat anti-mouse IgA (1:1,000 dilution; SouthernBiotech) as a detection antibody. To block non-specific binding, phosphate-buffered saline (PBS) containing 1% bovine serum albumin (BSA) was used. To measure S-hyIgA, the capture antibody was goat anti-mouse κ, but rabbit anti-mouse plgR (0.5 μg/mL; Sino Biological, Beijing, China) was used as the detection primary antibody. To measure total SC, goat anti-mouse plgR (1 μg/mL; R&D Systems, Minneapolis, MN, USA) was used as a capture antibody and the detection primary antibody was rabbit anti-mouse plgR. As standards, TEPC 15 mouse IgA myeloma proteins were purchased from Sigma-Aldrich, and recombinant soluble mouse plgR (extracellular domains) from R&D Systems. To detect plant-specific sugar chains on hybrid-IgG/IgA, the same capture antibody was used for total hybrid-IgG/IgA and rabbit anti-α1,3-fucose (0.5 μg/mL; Agrisera, Vännäs, Sweden) or rabbit anti-β1,2 xylose (1 μg/mL; Agrisera) were used as detection primary antibodies. To eliminate the effects of non-specific adsorption of plant proteins to ELISA wells, signals from wells with BSA-blocking alone were subtracted from those with anti-κ coating. The binding activity of hybrid-IgG/IgA to immobilized Stx1B was analysed by means of ELISA16. As detection primary antibodies, HRP-goat anti-mouse IgA (1:1,000 dilution), goat anti-mouse κ (1 μg/mL) or rabbit anti-mouse plgR (0.5 μg/mL) were used. For all indirect ELISA, HRP-goat anti-rabbit IgG (1:1,000 dilution; SouthernBiotech) or HRP-donkey anti-goat IgG (0.5 μg/mL; Santa Cruz Biotechnology, Santa Cruz, CA, USA) were used as the secondary antibodies.

**SDS-PAGE and immunoblotting.** The hybrid-IgG/IgA protein in the 50%-saturated ammonium sulphate precipitation sample was analysed by SDS-PAGE and immunoblotting as described previously16. Proteins were separated under reducing (12% Mini-PROTEAN TXG Gel, Bio-Rad) or non-reducing (4–20% Mini-PROTEAN TXG Gel, Bio-Rad) conditions, and blotted on PVDF membranes (Bio-Rad). The hybrid-IgG/IgA was detected with HRP-goat anti-mouse IgA (1:1,000 dilution; SouthernBiotech), goat anti-mouse κ (0.2 μg/mL; SouthernBiotech) plus HRP-donkey anti-goat IgG (0.2 μg/mL; Santa Cruz), rabbit anti-mouse IgA (1:1,000; Abcam, Cambridge, UK) and rabbit anti-mouse IgA (1 μg/mL; Jackson Immunoresearch, West Grove, PA, USA) or rabbit anti-mouse plgR (0.2 μg/mL; Sino Biological) plus HRP-anti-rabbit IgG (1:3,000 dilution). HiMark® Pre-Stained High Molecular Weight Protein Standards (Invitrogen) and MagicMark® XP Western Protein Standards (Invitrogen) were used as molecular weight standards.

**Immuno-electron microscopy.** The leaf tissue of *Arabidopsis* plants was fixed in 2% (v/v) glutaraldehyde-0.1 M potassium phosphate buffer (pH7.2) and embedded in LR White (London Resin, Berkshire, UK). Ultrathin sections (100 nm) were blocked with 5% BSA-PBS and then labelled with goat anti-mouse plgR (1 μg/mL; R&D Systems) and rabbit anti-mouse IgA (1:1,000; Abcam, Cambridge, UK). Normal goat IgG and normal rabbit IgG were used as isotype controls. The bound primary antibodies were detected with 12 nm colloidal gold-donkey anti-goat IgG (1:100 dilution; Jackson Immunoresearch, West Grove, PA, USA) and 18 nm colloidal gold-donkey anti-rabbit IgG (1:100 dilution; Jackson Immunoresearch). After electron staining, the sections were examined under a JEM-1011 transmission electron microscope (JEOL, Tokyo, Japan) at 80 kV.

**Cell viability assay.** To measure the cytotoxicity of Stx1 and neutralization by plantibodies, WST-8 cell viability assay was performed as previously described17. To compare the Stx1-sensitivity of Caco-2 cells with or without butyrate treatment, Caco-2 cells were seeded at 2 × 10⁴ cells/well in a 96-well cell culture plate (Falcon® 353072) and cultured for 3 days in 10% foetal bovine serum (FBS; Hyclone, South Logan, UT, USA)- Dulbecco’s modified Eagle’s medium (DMEM; Nissui Pharmaceuticals, Tokyo, Japan) supplemented with 1% non-essential amino acids (GIBCO BRL, Palm Alto, CA) at 37 °C under a humidified atmosphere of 5% CO₂/95% air. The cells were treated with 2 mM sodium butyrate in the medium for 4 days to induce G0/G1 expression on the cell membrane. The butyrate untreated cells were continuously cultured in the medium. Stx1 was added to Caco-2 cells at 0, 10, 30, 100 or 300 μg/mL. After 48 h culture, cell viability was assessed by WST-assay cell viability assaying (Cell Counting Kit-8; DOJINDO, Kumamoto, Japan) following the manufacturer’s instructions. Cell viability was calculated as the percentage of that of control level (no toxin exposure).

For toxin neutralization assaying using Caco-2 cells, 100 μg/mL of Stx1 holotoxin was incubated with the hybrid-IgG/IgA, an extract from wild-type plant leaves or mouse IgA myeloma protein TEPC 15 for 1 h at 37 °C. For plant preparation, proteins after precipitation with 50%-saturated ammonium sulphate were used. The mixture was then added to butyrate-treated Caco-2 cells. After 48 h culture, cell viability was assessed as above.
For toxin neutralization assays using Vero cells, cells were seeded at 2 × 10⁴ cells/well in a 96-well cell culture plate and cultured for 16 h in 10% FBS-medium 199 (M199; Gibco BRL). Preparation of the Stx1/plant-body mixture and Stx1 exposure were the same as in the experiments involving Caco-2 cells except that 10 pg/mL of Stx1 was used.

**DNA fragmentation assay.** Caco-2 cells were cultured in a 24-well cell culture plate (Corning, Costar 3526, NY, USA) at 5 × 10⁴ cells/well for 3 days. Sodium butyrate treatment and Stx1 exposure were carried out as described above. After 48 h incubation with Stx1, genomic DNA was extracted, purified and separated by agarose gel electrophoresis. DNA fragmentation was visualized as described previously.²¹

**References**

1. Woolf, J. M. & Kerr, M. A. The function of immunoglobulin A in immunity. J. Pathol. 208, 270–282 (2006).
2. Cortés, B. Multi-Faceted Functions of Secretory IgA at Mucosal Surfaces. Front. Immunol. 4, 185; 10.3389/fimmu.2013.00185 (2013).
3. Lindh, E. Increased resistance of immunoglobulin A dimers to proteolytic degradation after binding of secretory component. J. Immunol. 114, 284–286 (1975).
4. Loret, S. et al. Human plasma-derived polymeric IgA and IgM antibodies associate with secretory component to yield biologically active secretory-like antibodies. J. Biol. Chem. 288, 4085–4094 (2013).
5. Rogier, E. W. et al. Secretory antibodies in breast milk promote long-term intestinal homeostasis by regulating the gut microbiota and host gene expression. Proc. Natl. Acad. Sci. USA 111, 3074–3079 (2014).
6. Virdi, V. et al. Orally fed seeds producing designer IgAs protect weaned piglets against enterotoxigenic Escherichia coli infection. Proc. Natl. Acad. Sci. USA 110, 11809–11814 (2013).
7. Chintalacharuvu, K. R. & Morrison, S. L. Production of secretory immunoglobulin A by a single mammalian cell. Proc. Natl. Acad. Sci. USA 94, 6364–6368 (1997).
8. Johansen, F. E., Norderhaug, I. N., Røe, M., Sandlie, I. & Brandtzaeg, P. Recombinant expression of polymeric IgA: Incorporation of J chain and secretory component of human origin. Eur. J. Immunol. 29, 1701–1708 (1999).
9. Tesh, V. L. & Tesh, V. L. Shiga Toxins as Multi-Functional Proteins: Induction of Host Cellular Stress Responses, Role in Pathogenesis and Therapeutic Applications. Toxins (Basel) 8, 77, 10.3390/toxins8030077 (2016).
10. Chan, Y. S. & Ng, T. B. Shiga toxins: from structure and mechanism to applications.FEBS Lett. 574–584 (2011).
11. Lee, M.-S., Koo, S., Jeong, D. G. & Tesh, V. L. Shiga Toxins as Multi-Functional Proteins: Induction of Host Cellular Stress Responses, Role in Pathogenesis and Therapeutic Applications. Toxins (Basel) 8, 77, 10.3390/toxins8030077 (2016).
12. Jacewicz, M. S. The intergenic region of divergent genes functions as a bidirectional promoter. Plant Biotechnol. J. 2, 229–235 (2004).
13. Juarez, P. et al. Combinatorial Analysis of Secretory Immunoglobulin A (IgA) Expression in Plants. Int. J. Mol. Sci. 14, 6205–6222 (2013).
14. Juarez, P., Virdi, V., Depicker, A. & Orzaez, D. Bionanufacturing of protective antibodies and other therapeutics in edible plant tissues for oral applications. Plant Biotechnol. J. 14, 1791–1799 (2016).
15. Daniell, H., Streafield, S. J. & Wycoff, K. Medical molecular farming: Production of antibodies, biopharmaceuticals and edible vaccines in plants. Trends Plant Sci. 6, 219–226 (2001).
16. Ockenhouse, O. P., Poopa, J. O., Leelavathi, S. & Reddy, S. V. Advances in plant molecular farming. Biotechnol. Adv. 29, 210–222 (2011).
17. Tobisawa, Y. et al. Establishment of Recombiant Hybrid-IgG/IgA Immunoglobulin Specific for Shiga Toxin. Scand. J. Immunol. 74, 574–584 (2011).
18. Lee, M.-S., Koo, S., Jeong, D. G. & Tesh, V. L. Shiga Toxins as Multi-Functional Proteins: Induction of Host Cellular Stress Responses, Role in Pathogenesis and Therapeutic Applications. Toxins (Basel) 8, 77, 10.3390/toxins8030077 (2016).
19. Chan, Y. S. & Ng, T. B. Shiga toxins: from structure and mechanism to applications. Appl. Microbiol. Biotechnol. 100, 1597–1610 (2012).
20. Tesh, V. L. Induction of apoptosis by Shiga toxins. Future Microbiol. 5, 431–453 (2010).
21. Kovalskij, O. et al. The glycosphingolipid globotriaosylceramide in the metastatic transformation of colon cancer. Proc. Natl. Acad. Sci. USA 102, 19087–19092 (2005).
22. LaCasse, E. C. et al. Shiga-like toxin-1 receptor on human breast cancer, lymphoma, and myeloma and absence from CD34(+) hematopoietic stem cells: implications for cx vivo tumor purging and autologous stem cell transplantation. Blood 94, 2901–2910 (1999).
23. Jacewicz, M. S. et al. Maturational regulation of globotriaosylceramide, the Shiga-like toxin 1 receptor, in cultured human gut epithelial cells. J. Clin. Invest. 96, 1328–1335 (1995).
24. Ito, Y., Iguchi, M. & Kurata, N. KNOX homeobox genes are sufficient in maintaining cultured cells in an undifferentiated state in rice. Genesis 30, 231–238 (2001).
25. Takaiwa, F., Kurohane, K., Nakanishi, K., Miya, M. & Imai, Y. Stable expression and characterization of monomeric and dimeric recombinant hybrid-IgG/IgA immunoglobulins specific for Shiga toxin. Biol. Pharm. Bull. 37, 1510–1515 (2014).
26. Ito, Y., Iguchi, M. & Kurata, N. KNOX homeobox genes are sufficient in maintaining cultured cells in an undifferentiated state in rice. Genesis 30, 231–238 (2001).
27. Takaiwa, F., Yang, L., Maruyama, N., Wakasa, Y. & Ozawa, K. Deposition mode of transforming growth factor-β3 expressed in transgenic rice seed. Plant Cell Rep. in press; 10.1007/s00299-016-2047-2 (2016).
28. Schüller, S., Frankel, G. & Phillips, A. D. Interaction of Shiga toxin from Escherichia coli with human intestinal epithelial cell lines and explants: Stx2 induces epithelial damage in organ culture. Cell. Microbiol. 6, 289–301 (2004).
29. Tanikawa, T., Ishikawa, T., Maekawa, K., Kuronome, K. & Imai, Y. Characterization of monoclonal immunoglobulin A and G against Shiga toxin binding subunits produced by intranasal immunization. Scand. J. Immunol. 68, 414–422 (2008).
30. De Muyck, B., Navarre, C. & Boutry, M. Production of antibodies in plants: Status after twenty years. Plant Biotechnol. J. 10, 529–563 (2012).
31. da Cunha, N. B., Vianna, G. R., da Almeida Lima, T. & Rech, E. Molecular farming of human cytokines and blood products from plants: Challenges in biosynthesis and detection of plant-produced recombinant proteins. Biotechnol. J. 9, 39–50 (2014).
32. Virdi, V., Juarez, P., Boudoul, V. & Depicker, A. Recombinant IgA production for mucosal passive immunization, advancing beyond the hurdles. Cell. Mol. Life Sci. 73, 535–545 (2016).
33. Wycoff, K. L. et al. Recombinant anthrax toxin receptor-Fc fusion proteins produced in plants protect rabbits against inhalational anthrax. Antimicrob. Agents Chemother. 55, 132–139 (2011).
36. Niemer, M. et al. The human anti-HIV antibodies 2F5, 2G12, and PG9 differ in their susceptibility to proteolytic degradation: Down-regulation of endogenous serine and cysteine proteinase activities could improve antibody production in plant-based expression platforms. Biotechnol. J. 9, 493–500 (2014).
37. Hehle, V. K. et al. Site-specific proteolytic degradation of IgG monoclonal antibodies expressed in tobacco plants. Plant Biotechnol. J. 13, 235–245 (2015).
38. Saberianfar, R., Joensuu, J. J., Conley, A. J. & Menassa, R. Protein body formation in leaves of Nicotiana benthamiana: a concentration-dependent mechanism influenced by the presence of fusion tags. Plant Biotechnol. J. 13, 927–937 (2015).
39. Nicholson, L. et al. A recombinant multimeric immunoglobulin expressed in rice shows assembly-dependent subcellular localization in endosperm cells. Plant Biotechnol. J. 3, 115–127 (2005).
40. Jiang, L. et al. The protein storage vacuole: a unique compound organelle. J. Cell Biol. 155, 991–1002 (2001).
41. Schüller, S. Shiga toxin interaction with human intestinal epithelium. Toxins (Basel). 3, 626–639 (2011).
42. Hurley, B. P., Thorpe, C. N. & Acheson, D. W. Shiga toxin translocation across intestinal epithelial cells is enhanced by neutrophil transmigration. Infect. Immun. 69, 6148–6155 (2001).
43. Imai, Y. et al. Restricted expression of shiga toxin binding sites on mucosal epithelium of mouse distal colon. Infect. Immun. 71, 985–990 (2003).
44. Kashiwamura, M. et al. Shiga toxin kills epithelial cells isolated from distal but not proximal part of mouse colon. Biol. Pharm. Bull. 32, 1614–1617 (2009).
45. Shoji, K. et al. Recombinant immunoglobulin A specific for influenza A virus hemagglutinin: production, functional analysis, and formation of secretory immunoglobulin A. Viral Immunol. 28, 170–178 (2015).

Acknowledgements
This work was supported by JSPS KAKENHI Grant Numbers JP15H04660 and JP25670063 to YI; JP16K21277 and JP25-10915 to KN. We thank Mr. N.J. Halewood for language editing services.

Author Contributions
K.N. and Y.I. wrote the main manuscript text and K.N. prepared figures. Y.A., S. Matsubara and H. Kawakami performed immuno-electron microscopy. K.N., S. Morikane, S.I. and Y.N. performed plant biotechnology experiments, and K.N., S. Morikane, and K.K. performed immunological experiments. Y.I. and H. Kobayashi designed and supervised the study. All authors reviewed the manuscript.

Additional Information
Competing Interests: The authors declare no competing financial interests.

How to cite this article: Nakanishi, K. et al. Protection of Human Colon Cells from Shiga Toxin by Plant-based Recombinant Secretory IgA. Sci. Rep. 7, 45843; doi: 10.1038/srep45843 (2017).

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/

© The Author(s) 2017