

Moss-based production of asialo-erythropoietin devoid of Lewis A and other plant-typical carbohydrate determinants

Juliana Parsons¹, Friedrich Altmann², Claudia K. Arrenberg¹, Anna Koprivova^{1†}, Anna K. Beike¹, Christian Stemmer^{3‡}, Gilbert Gorr^{3‡}, Ralf Reski^{1,4,5} and Eva L. Decker^{1*}

¹Plant Biotechnology, Faculty of Biology, University of Freiburg, Freiburg, Germany

²Department of Chemistry, University of Natural Resources and Life Sciences, Vienna, Austria

³Greenovation Biotech, Freiburg, Germany

⁴BIOSS Centre for Biological Signalling Studies, Freiburg, Germany

⁵FRIAS Freiburg Institute for Advanced Studies, Freiburg, Germany

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*Correspondence (fax 49 761 2036945; email eva.decker@biologie.uni-freiburg.de)

†Present address: John Innes Centre, Norwich Research Park, Colney, Norwich NR4 7UH, UK.

‡Present address: Emcid Biotech, Hans-Bunte-Str. 19, 79108 Freiburg, Germany.

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Summary

Protein therapeutics represent one of the most increasing areas in the pharmaceutical industry. Plants gain acceptance as attractive alternatives for high-quality and economical protein production. However, as the majority of biopharmaceuticals are glycoproteins, plant-specific N-glycosylation has to be taken into consideration. In *Physcomitrella patens* (moss), glyco-engineering is an applicable tool, and the removal of immunogenic core xylose and fucose residues was realized before. Here, we present the identification of the enzymes that are responsible for terminal glycosylation (α 1,4 fucosylation and β 1,3 galactosylation) on complex-type N-glycans in moss. The terminal trisaccharide consisting of α 1,4 fucose and β 1,3 galactose linked to N-acetylglucosamine forms the so-called Lewis A epitope. This epitope is rare on moss wild-type proteins, but was shown to be enriched on complex-type N-glycans of moss-produced recombinant human erythropoietin, while unknown from the native human protein. Via gene targeting of moss galactosyltransferase and fucosyltransferase genes, we identified the gene responsible for terminal glycosylation and were able to completely abolish the formation of Lewis A residues on the recombinant biopharmaceutical.

Introduction

Recombinant protein therapeutics constitute the largest class of products currently developed by the pharmaceutical industry. As about 70% of the so-called biopharmaceuticals on the market or in clinical studies are glycoproteins (Durocher and Butler, 2009), the development of alternative expression systems with human-like glycosylation patterns is essential. Protein glycosylation in mammalian expression systems is most similar to human glycans. Likewise, plants, as higher eukaryotes, are able to produce complex-type N-glycans (Lerouge *et al.*, 1998; Wilson, 2002). The core structure of the complex N-glycans, a pentasaccharide of three mannoses and two N-acetylglucosamines (Man3GlcNAc2), is conserved between mammals and plants, as well as up to two terminally attached GlcNAc residues (reviewed in Gomord *et al.*, 2010). Differing from the human structure, most plant N-glycans carry a β 1,2 xylose and an α 1,3 fucose linked to the glycan core. These residues raised concerns about plant-produced biopharmaceuticals as they were shown to be highly immunogenic in mammals (Bardor *et al.*, 2003; Bencúrová *et al.*, 2004; Garcia-Casado *et al.*, 1996; Jin *et al.*, 2008; Van Ree *et al.*, 2000). The creation of transgenic plants, in which expression of the respective glycosyltransferase genes was either abolished (Koprivova *et al.*, 2004; Strasser *et al.*, 2004) or reduced (Cox *et al.*, 2006; Sourrouille *et al.*, 2008;

Strasser *et al.*, 2008), increased the acceptance of plant-made biopharmaceuticals, and several products are currently studied in clinical trials (Davies, 2010; Paul and Ma, 2011). Another difference between human and plant complex-type N-glycans lies in their terminal elongation. While on many human N-glycans, β 1,4 galactose (h1,4 Gal) is linked to the outer GlcNAc residues followed by terminal sialic acid, these two sugars are non-existing on plant glycoproteins. In contrast, β 1,3 galactose (Gal) and α 1,4 fucose (Fuc) may be attached to the terminal GlcNAc residues giving rise to the Fuc α 1-4(Gal β 1-3)GlcNAc structure. This trisaccharide, which was originally described for Lewis-positive histo-blood groups in humans (Henry *et al.*, 1995), is known as Lewis A epitope (Le^a). Although Le^a determinants may be regarded as human-type glycosylation, this structure occurs only rarely on glycoproteins of healthy adult humans. Moreover, Le^a and Sialyl-Le^a were shown to be highly increased in the sera from patients with colon cancer (Zhang *et al.*, 1994). Therefore, the repression of Le^a attachment to recombinant plant-made biopharmaceuticals is desirable.

Le^a is widespread on glycoproteins of several plant species and was estimated to represent about 15% of the total N-glycan population in tobacco (Fitchette *et al.*, 1999; Wilson, 2001). In addition, the Le^a epitope was also described on recombinant proteins produced in tobacco (Petruccioli *et al.*, 2006). In contrast, the appearance of Le^a structures on

glycoproteins in the moss *Physcomitrella patens* or the model plant *Arabidopsis thaliana* was shown to be low (Koprivova et al., 2003; Strasser et al., 2007; Viëtor et al., 2003). In *P. patens*, several biopharmaceuticals were successfully expressed (Baur et al., 2005; Büttner-Mainik et al., 2011; Gorr and Jost, 2005; Schuster et al., 2007; Weise et al., 2007); however, in one case, the human erythropoietin (hEPO), a significant proportion of Le^a structures attached to the N-glycan core was observed (Weise et al., 2007).

Erythropoietin is a hormone, predominantly synthesized by the kidneys, which plays an important role in the maturation of red blood cells (erythropoiesis). It is commonly used to treat and prevent anaemia in nephrology and oncology patients (Jelkmann, 2000). Recombinant hEPO (rhEPO), mainly produced in chinese hamster ovary (CHO) cells, has a worldwide market of about 10 billion euros per year, making it the leading biopharmaceutical in the market (Jelkmann, 2007). Human EPO consists of 165 amino acids and has a molecular mass of approximately 34 kDa. Three N-linked (at Asn 24, Asn 38 and Asn 83) and one O-linked (at Ser 126) carbohydrate chains are attached to the amino acid chain, which in total represent 40% of the molecular mass (Takeuchi et al., 1989). Yet, these oligosaccharides are not essential for the activity of EPO *in vitro*, indicating that solely the polypeptide is involved in binding to the EPO receptor (Dordal et al., 1985). On the other hand, desialylated EPO (asialo-EPO) loses its erythropoietic activity *in vivo*, while it maintains, or even increases, its receptor affinity *in vitro* (Tsuda et al., 1990). This indicates a major role of glycosylation in increasing the half-life of EPO in circulation (Takeuchi and Kobata, 1991).

More recently, EPO was shown to be produced in many tissues under hypoxia or metabolic stress, where it inhibits apoptosis and protects these tissues from collateral damage elicited by the injury (reviewed in Brines and Cerami, 2008). In this context, EPO was found to be neuroprotective, representing a potential treatment for ischaemic stroke, diabetes-induced eye damage and peripheral nerve injury (Brines et al., 2008). For these indications, an increase in red blood cells is not desirable (Masuda et al., 1993). However, asialo-EPO can provide tissue protection without the undesired erythropoietic side effect, making asialo-EPO an advantageous pharmaceutical (Carelli et al., 2011; Erbayraktar et al., 2003).

Recombinant hEPO was already successfully produced in moss (Weise et al., 2007), and its activity was confirmed *in vitro*. Considering that plants naturally lack the required enzymes for glycoprotein sialylation, moss can be assumed as a suitable platform for the production of asialo-EPO. However, to reliably provide a safe pharmaceutical, Le^a residues detected on moss-produced rhEPO have to be abolished.

Here, we identified the genes encoding α 1,4 fucosyl- and β 1,3 galactosyltransferases responsible for terminal plant N-glycan elongation in *P. patens*. In this species, gene functions can be eliminated by precise gene targeting via homologous recombination with a unique efficiency (Kamisugi et al., 2006; Khraiweh et al., 2010; Mosquna et al., 2009; Strepp et al., 1998). Using this method to target the corresponding glycosyltransferase genes in transgenic rhEPO-producing moss lines, we were able to completely abolish Le^a formation on recombinant as well as endogenous moss glycoproteins, thus providing the next generation of plant N-glycan humanization. Furthermore, we successfully demonstrated β 1,4 galactosylation of rhEPO after introduction of human β 1,4 galactosyltransferase to the Le^a-free moss lines.

Results

Identification of α 1,4 fucosyltransferase and β 1,3 galactosyltransferase genes from *Physcomitrella patens*

Via BLAST (Basic Local Alignment Search Tool) searches in the Cosmoss database (<http://www.cosmoss.org>) using human and *Arabidopsis* α 1,4 fucosyltransferases (FucT) and β 1,3 galactosyltransferases (GalT), we were able to identify respective homologous sequences from the *P. patens* genome. The full-length cDNA of Pp*fuct* was obtained by 5' and 3' RACE (rapid amplification of cDNA ends) PCR. In *Physcomitrella*, α 1,4 fucosyltransferase is a single-copy gene (Pp1s151_12V6.1), encoding a protein of 437 amino acids. The PpFucT protein showed 50.1% identity to FucT from *A. thaliana* (FUT13, AAK11728) and 19.6% identity to the human FucT (FUT3; NP_001091110), an enzyme with α 1,3 fucosyltransferase and α 1,4 fucosyltransferase activities, which is responsible for the addition of fucose in the last step of Le^a antigen biosynthesis.

Eleven amino acid sequences of putative galactosyltransferases from *P. patens* resulted from the BLASTP search against the gene models derived from the moss genome sequence: Pp1s251_37V6.1, Pp1s237_65V6.1, Pp1s163_84V6.1, Pp1s65_62V2.1, Pp1s34_213V2.1, Pp1s101_246V0.1, Pp1s152_166V0.1, Pp1s313_19V0.1, Pp1s34_146V0.1, Pp1s14_277V0.1, Pp1s263_36V0.1. All 11 amino acid sequences show sequence identities of 19%–31% and sequence similarities of 39%–49% to all six human sequences, within the galactosyltransferase domain (Pfam 01762). Subsequently, a protein family search in the Pfam 24.0 database (<http://pfam.sanger.ac.uk/>) was performed to identify further galactosyltransferase genes (Pfam 01762) in *P. patens*. Two additional putative galactosyltransferases were found: Pp1s121_83V0.1 and Pp1s111_119V2.1. Three of the GalT homologues, Pp1s251_37V6.1, Pp1s237_65-V6.1 and Pp1s163_84V6.1, were predicted to contain a signal peptide for the secretory pathway and a transmembrane domain, as well as a galactoside-binding lectin domain (Pfam 0037). These features were considered as potential prerequisites for the physiologic function of the GalT on glycoproteins in the Golgi apparatus. A phylogenetic analysis (Figure 1), performed with the putative *Physcomitrella* GalT sequences as well as six lectin-binding domain-containing *Arabidopsis* GalT and six human GalT homologues, shows distinct clades for the putative galactosyltransferases from the moss. One PpGalT sequence (Pp1s251_37V6.1) forms a distinct clade with two sequences from *A. thaliana* (At1g26810 and At3g06440). As Pp1s251_37V6.1 clusters together with GalT1 from *A. thaliana* (Strasser et al., 2007), it was named GalT1 for *Physcomitrella* (PpGalT1). Two additional PpGalT sequences (Pp1s237_65V6.1 and Pp1s163_84V6.1) cluster with four AtGalT sequences (At1g74800, At4g21060, At1g27120 and At5g62620) and were named PpGalT2 and PpGalT3, respectively.

Generation of *fuct* and *galt* single- and double-knockout plants

To reveal their specific function on Le^a formation in moss glycoproteins, we created plasmid constructs for the targeted knockout (KO) of the single *fuct* gene as well as for the three *galt* homologues 1, 2 and 3 (Figure S1a). These constructs were transferred into *P. patens* protoplasts to generate single deletions for *fuct*, *galt1* and *galt3* via homologous recombination, respectively. Owing to the high similarity of GalT2 and GalT3

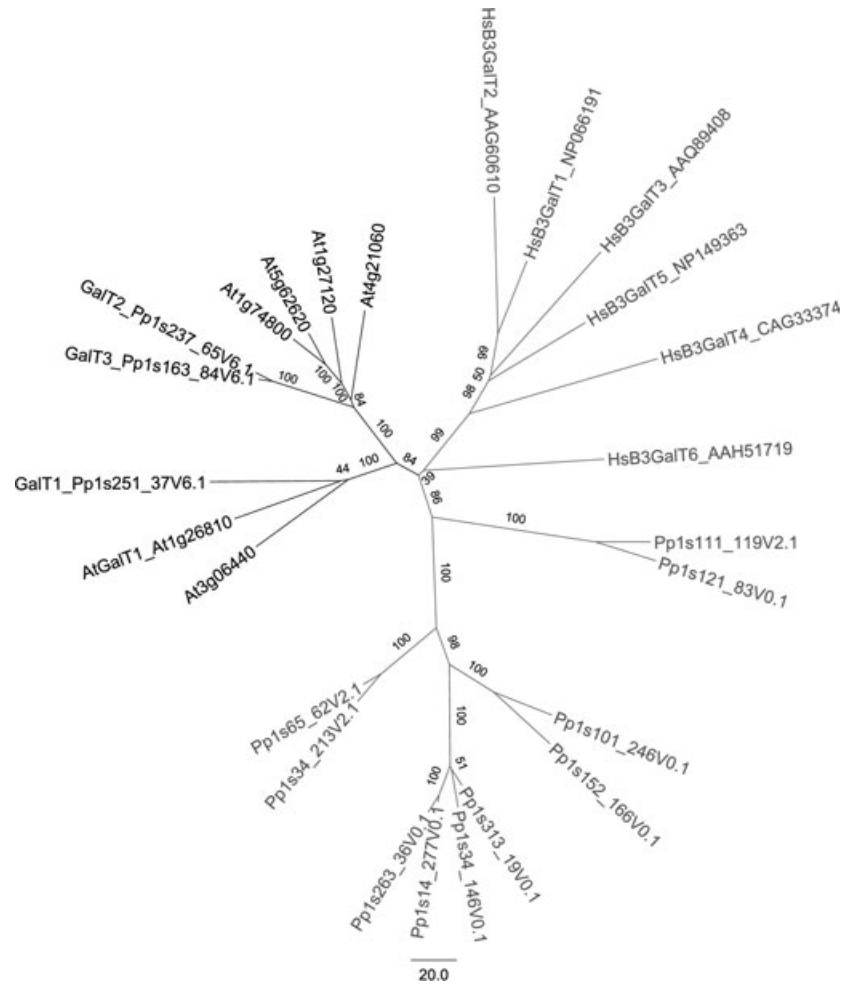


Figure 1 Phylogeny of putative galactosyltransferases from *Physcomitrella patens*. The phylogenetic tree was constructed from 13 *P. patens* amino acid sequences sharing the galactosyltransferase domain (Pfam 01762), six *Arabidopsis thaliana* sequences (At1g26810, At3g06440, At1g74800, At5g62620, At1g27120 and At4g21060) and six *Homo sapiens* sequences (HsB3GalT1, HsB3GalT2, HsB3GalT3, HsB3GalT4, HsB3GalT5 and HsB3GalT6) with the neighbour-joining algorithm implemented in Quicktree (Howe *et al.*, 2002). The tree was visualised with FigTree V 1.1.2. The clades of GalT1_Pp1s251_37V6.1 and its homologues in *A. thaliana*, and GalT2_Pp1s237_65V6.1, GalT3_Pp1s163_84V6.1 and their homologues in *A. thaliana* are shown in black. Bootstrap values were determined from 1000 replicates and are shown at the respective branches in percentage. Bar = 20.0% sequence divergence.

proteins, which share 71.5% identity, additionally $\Delta gal2/\Delta gal3$ double KO lines were created. For this purpose, a $\Delta gal3$ moss line was transformed with the *galt2* KO construct. Because of the observation that the moss-produced rhEPO displays a substantial amount of Le^a, the rhEPO-producing line 174.16 (Weise *et al.*, 2007) was used as parent plant for the *galt* KO lines. Plants that survived two cycles of antibiotic selections were screened for genomic integration of the KO construct in the correct genomic locus. Plants with positive PCR for both 5' and 3' integration were further analysed by RT-PCR to prove the disappearance of the respective transcript (Figure S1b–e). By these means, seven lines were confirmed as targeted KO lines for the *fuct* gene as well as nine $\Delta gal1$ lines. In addition, one $\Delta gal3$ line and two $\Delta gal2/\Delta gal3$ double-KO lines were confirmed. One or two lines, respectively, of each genetic modification were chosen for further analysis and stored in the International Moss Stock Center (<http://www.moss-stock-center.org>; see Table S4).

Terminal N-glycosylation is affected in *fuct* and *galt* KO lines

To explore whether the disruption of the *galt* and *fuct* genes indeed results in changes of the glycosylation pattern, N-glycans of the cellular protein fraction from *Physcomitrella* wild type (WT), the rhEPO-synthesizing parent plant 174.16 as well as the different glycosyltransferase KO lines were analysed by MALDI-TOF mass spectrometry.

The major peak in the glycan spectrum of WT moss represented the complex-type N-glycan structure GnGnXF with terminally attached GlcNAc residues (Gn) and core xylose (X) and fucose (F), respectively (Figure 2a top). In addition, Le^a structures (FA) on one or both of the antennae were detected [Gn(FA)XF_{iso} and (FA)(FA)XF, respectively]. The $\Delta fuct$ plant displays structures lacking the terminal fucose residues; however, β 1,3-linked galactose residues (A) are still present (lower Figure 2a, GnAXF_{iso} and AAXF). The rhEPO-expressing parent plant (174.16), in which core xylose and fucose residues had been abolished by targeted KO (Weise *et al.*, 2007), presents the GnGn structure and additional Le^a-containing glycans, Gn(FA)_{iso} or (FA)(FA) (upper Figure 2b). The $\Delta gal2/\Delta gal3$ double-KO plants did not show any alteration in the terminal antennae glycosylation in comparison with the parent plant (middle Figure 2b). The same was true for the $\Delta gal3$ single KO (data not shown), suggesting that these GalT homologues are not involved in the synthesis of Le^a epitopes in the gametophytic tissue. In contrast, in the $\Delta gal1$ plants, the major peak that appears in the spectrum corresponds to GnGn glycans, and no peak was observed around the Le^a-indicative masses of 1648 or 1956 Da, proving that $\Delta gal1$ is completely devoid of terminal β 1,3 galactosyl residues. Moreover, any terminal fucose is also missing. This finding supports the observation that the α 1,4 fucosyltransferase transfers a fucose residue to the preformed Gal β 1,3GlcNAc substrate (Léonard *et al.*,

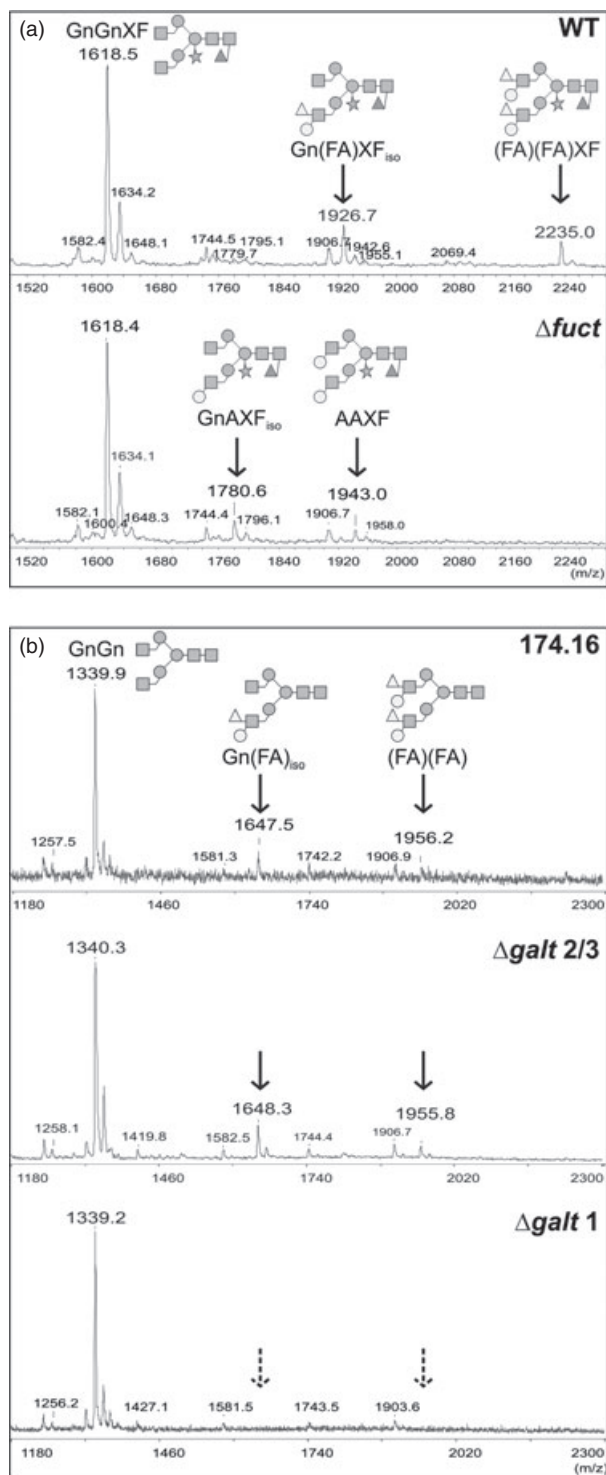


Figure 2 MALDI-TOF mass spectra of the endogenous N-linked glycans from moss. (a) Comparison of the MALDI-TOF mass spectra for the wild-type moss and the $\Delta fuct$ mutant. (b) Comparison of the MALDI-TOF mass spectra for the recombinant human erythropoietin-producing plant 174.16, the double-knockout $\Delta gal2/\Delta gal3$ No. 23 and the $\Delta gal1$ plant No. 58. For the indicated masses, please refer to Table 1. ○: galactose, ☆: xylose, ■: N-acetylglucosamine, ○: mannose, △/△: fucose.

2002; Wilson, 2001), that is, the FucT acts as the last enzyme in the biosynthesis of plant glycoproteins, and preceding GalT

activity is essential to provide the appropriate substrate for the FucT.

Moss-derived Lewis A-free recombinant hEPO

Via glycan analysis of the different moss strains, we proved the disappearance of Le^a residues on total intracellular N-glycans of moss plants lacking GalT1. Most important for biopharmaceutical production, however, is to confirm that the modified plants also produce Le^a-free rhEPO. Therefore, extracellular proteins from the parent plant 174.16, $\Delta gal2/\Delta gal3$ and $\Delta gal1$ lines were precipitated from the culture supernatant, separated via SDS-PAGE and analysed by Western blotting.

Using anti-EPO antibodies, rhEPO could be detected in the medium of all analysed plants (Figure 3a). Different from CHO-derived rhEPO, which was chosen as a standard and migrated with an apparent molecular weight (MW) of about 34 kDa, moss-produced rhEPO migrated with a molecular weight (MW) of approximately 25 kDa. This size difference is because of different glycosylation patterns. CHO-produced rhEPO displays mainly tetra-antennary sialylated N-glycans, while moss-produced rhEPO displays exclusively di-antennary N-glycans as was also observed on human serum EPO (Skibeli *et al.*, 2001). The estimated different amounts of mainly mannose, galactose and sialic acid residues result in a calculated size difference of more than 7 kDa.

Using the anti-Le^a antibody JIM 84 (Horsley *et al.*, 1993), we detected terminal Le^a structures, accumulating at the size of rhEPO, on total secreted proteins of the parent plant (174.16) as well as the $\Delta gal2/\Delta gal3$ line. In contrast, we observed the complete absence of any Le^a signal in the $\Delta gal1$ plants (Figure 3b).

To investigate the structure of the glycans attached to our protein of interest, the moss-produced rhEPO was further analysed via mass spectrometry. For this purpose, total soluble proteins were precipitated from the culture supernatant of the parent plant and $\Delta gal1$ lines and separated by SDS-PAGE. Subsequently, the main EPO-containing band was cut out from the Coomassie-stained gel (Figure S2) and subjected to mass spectrometry for an analysis of the EPO glycopeptides.

While the overall proportion of Le^a residues on moss glycoproteins is very small (Koprivova *et al.*, 2003), it was obviously enriched on the moss-produced rhEPO. Around three-quarters of the rhEPO molecules displayed complex glycans containing Le^a residues (Figure 4). With 3 N-glycosylation sites bound by complex-type bi-antennary glycans, there are several possible combinations where at least one Le^a motif is present. The possible glycosylation pattern for the two tryptic glycopeptides is listed in Table 2. By knocking out the *gal1* gene, the Le^a motif disappeared completely from the rhEPO glycosylation pattern, while the core glycan GnGn remains intact (Figure 4). As already observed for intracellular total moss glycoproteins, the main terminal sugar on moss N-glycans is GlcNAc, while terminal mannoses (M) are far less prominent (Figure 2 and Table 2). This was even more striking in the specific case of secreted rhEPO glycosylation which exhibits a tiny residual GnM peak in addition to the prominent GnGn structure. Therefore, we achieved an outstandingly homogenous product quality, having our protein of interest with almost exclusively the desired glycan pattern.

Phenotype and growth rate of the mutants were comparable to the parent lines, indicating that the overall metabolism is not affected by the mutations (data not shown). The rhEPO produced in different moss lines was measured by means of

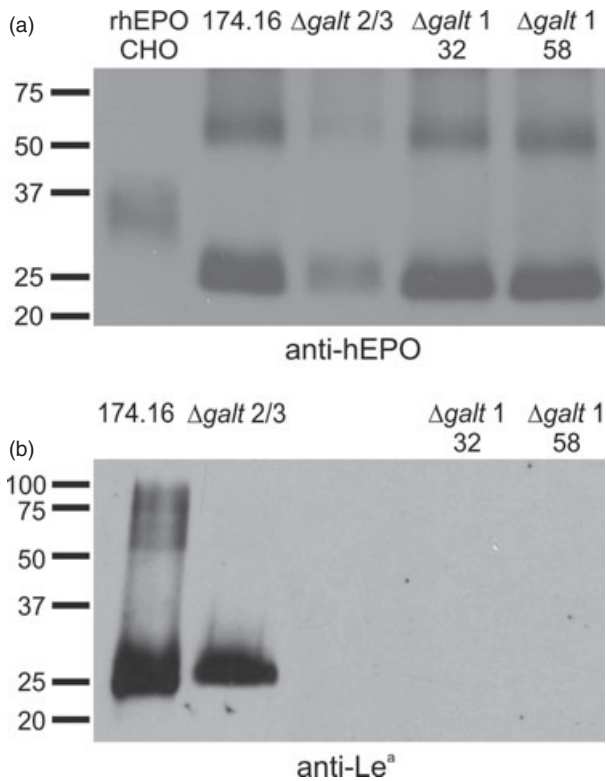


Figure 3 Western blot analysis proved disappearance of Le^a structures from moss-derived recombinant human erythropoietin (rhEPO). Total soluble proteins (TSP) were precipitated from culture supernatant of plants 174.16, Δ *galt2/Δgalt3* No. 23, Δ *galt1* No. 32 and 58. (a) Twenty-five nanograms of extracellular TSP and 5 ng of chinese hamster ovary-produced rhEPO standard were loaded, and detection was performed with anti-hEPO antibodies. (b) Thirty micrograms of TSP were loaded, and detection was performed with the anti-Lewis A (Le^a) antibody JIM 84.

ELISA. The rhEPO production levels observed in the Δ *galt1* plants were comparable to that of the parent plant 174.16 (Table S3), indicating that the genetic modifications did not affect the productivity of the rhEPO or stability of the product, respectively.

Although *in vitro* activity of moss-derived rhEPO has already been proven (Weise *et al.*, 2007), the addition of terminal human-type β 1,4 galactose to the rhEPO molecule might be useful for tissue-protective EPO activity. Therefore, we created rhEPO-producing Δ *galt1* lines that additionally expressed the human β 1,4 galactosyltransferase gene (*h1,4galt*). Some of the resulting plants that showed *h1,4galt* transcription (Figure S3) were further analysed for their ability to produce β 1,4 galactosylated rhEPO. Total soluble proteins from the culture supernatants of *h1,4galt* insertion lines (*h1,4galt* i) were separated via SDS-PAGE and analysed by lectin affinity blotting with *Datura stramonium* agglutinin (DSA) which specifically detects Gal β 1,4GlcNAc residues (Figure 5a). As a control, an anti-EPO Western blot was performed (Figure 5b). While no signal was observed for Δ *galt1* plants, we clearly detected terminal β 1,4 galactose structures accumulating at the size of rhEPO in two *h1,4galt* i lines, proving the activity of the introduced human β 1,4GalT. Comparison with the EPO bands detected by anti-EPO Western blotting suggests that a substantial proportion of the moss-derived rhEPO contains human-type β 1,4 galactose residues.

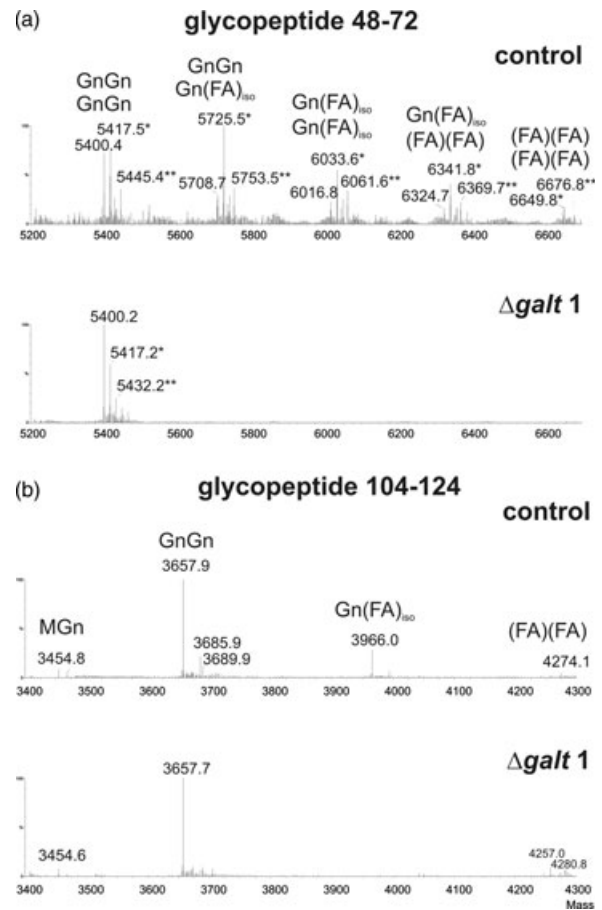


Figure 4 Glycopeptides of moss-produced recombinant human erythropoietin (rhEPO) from plants Δ *galt1* No. 32 and control plant 174.16. Tryptic digestion of rhEPO generated two glycopeptides. The peptide harbouring two glycosylation sites (EAENITTCGAHEHCLSNENITVPDTK) is shown in panel a. The tryptic peptide containing one glycan (GQALLVNSSQPWEPLQLHVDK) is depicted in panel b. Glycopeptides containing Le^a structures are totally absent in the Δ *galt1* line. For the indicated masses, please refer to Table 2. Salt adducts are marked with asterisks.

Discussion

N-glycosylation is one of the most important post-translational modifications of proteins, which may have impact on functionality, stability or compatibility of a recombinant pharmaceutical. It starts in the endoplasmic reticulum with the transfer of the oligosaccharide precursor Glc₃Man₉GlcNAc₂ from a dolichol carrier to specific Asn-X-Ser (or Thr) tripeptides on the polypeptide chain. Subsequently, processing takes place in the Golgi apparatus, where complex-type N-glycans are finally synthesized. Plant complex-type N-glycans are characterized by the attachment of plant-specific β 1,2 xylose and α 1,3 fucose to the glycan core. Besides these modifications, plant complex-type N-glycans may be further processed by addition of terminal fucose and galactose residues yielding Fuc α 1-4(Gal β 1-3)GlcNAc sequences (Fitchette-Lainé *et al.*, 1997; Melo *et al.*, 1997). These residues are synthesized by β 1,3 galactosyltransferase (GalT) and α 1,4 fucosyltransferase (FucT) and named Lewis A antigens (Le^a). Le^a-containing N-glycans are present in a wide spectrum of plants including mosses, ferns, and seed plants (Fitchette *et al.*, 1999;

Table 1 Structures and corresponding calculated masses of the major N-glycans, for the wild-type moss (WT), single-KO $\Delta 1,4fuct$, double-KO $\Delta 1,3fuct/\Delta xy1t$ (glycan background from the plant 174.16) and $\Delta \alpha 1,3fuct/\Delta xy1t/\Delta galt$ triple-KO plants

WT		$\Delta \alpha 1,4fuct$		$\Delta \alpha 1,3fuct + \Delta xy1t$		$\Delta \alpha 1,3fuct + \Delta xy1t + \Delta galt$	
Mass	Structure	Mass	Structure	Mass	Structure	Mass	Structure
1582.4	M7	1582.4	M7	1258.2	M5	1258.2	M5
1618.3	GnGnXF	1618.3	GnGnXF	1340.2	GnGn	1340.2	GnGn
1744.5	M8	1744.5	M8	1420.3	M6	1420.3	M6
1906.7	M9	1906.7	M9	1582.4	M7	1582.4	M7
1926.8	Gn(FA)XF _{iso}	1780.6	Gn(A)XF _{iso}	1648.5	Gn(FA) _{iso}	1486.5	Gn(F) _{iso} *
2235.0	(FA)(FA)XF	1942.8	(A)(A)XF	1744.5	M8	1744.5	M8
				1906.7	M9	1906.7	M9
				1955.7	(FA)(FA)	1631.7	(F)(F)*

*Masses and structures for an hypothetical case, where only terminal galactose disappears but terminal fucose remains in the $\Delta galt$ plants, in addition to the absence of xylose and $\alpha 1,3$ fucose.

KO, knockout.

Table 2 Average masses of ions that could be assigned to tryptically digested glycopeptides of recombinant human erythropoietin

Glycopeptide 48–72				Glycopeptide 104–124	
EAENITGCAEHCSLNENITVPDTK (glycosylation sites A and B)				GQALLVNSQPWEPLQLHVDK (glycosylation site C)	
A	B	M+H	M+NH ₄ ⁺	C	M+H
GnGn	GnM	5197.12	5214	GnM	3454.64
GnGn	GnGn	5400.20	5417	GnGn	3657.72
GnGn	(FA)Gn	5708.31	5725	AGn	3819.77
(FA)Gn	(FA)Gn	6016.42	6033	(FA)Gn	3965.83
(FA)Gn	(FA)(FA)	6324.53	6341	AA	3981.82
(FA)(FA)	(FA)(FA)	6632.65	6649	(FA)(FA)	4273.94

One of the three N-glycosylation sites lays on peptide 104–124, while the other two lay on the same peptide (48–72), giving raise to different combinations of glycopeptides.

Glycosylation sites underlined.

Wilson *et al.*, 2001). They were estimated to represent about 15% of the total N-glycan population in tobacco (Fitchette *et al.*, 1999). As demonstrated by immunolabelling experiments, the Le^a epitope is associated with extracellular and plasma membrane-bound glycoproteins (Fitchette *et al.*, 1999), those compartments in which the majority of recombinant pharmaceutical glycoproteins are located. As Le^a structures are rare on human glycoproteins and increased in connection with certain types of cancer (Zhang *et al.*, 1994), they should be avoided on recombinant therapeutics.

All of the biopharmaceuticals that have already been successfully expressed in the moss *P. patens*, human vascular endothelial growth factor, complement regulatory factor H, IgG immunoglobulins and hEPO (Baur *et al.*, 2005; Büttner-Mainik *et al.*, 2011; Gorr and Jost, 2005; Schuster *et al.*, 2007; Weise *et al.*, 2007) are extracellular glycoproteins. One of them, hEPO, showed a high proportion of N-glycans bearing the Le^a epitope (Weise *et al.*, 2007).

The features that make moss an attractive system for biopharmaceutical production comprise its growth as tissue cultures in strictly contained and scalable photobioreactors, the

possibility of product secretion to the culture supernatant and high flexibility for medium pH and temperature (Decker and Reski, 2007, 2012). Its unique amenability for targeted gene replacements by homologous recombination has been employed in adapting the moss glycosylation pattern to the requirements for biopharmaceutical production by knocking out the core $\beta 1,2$ xylosyltransferase and $\alpha 1,3$ fucosyltransferase genes (Koprivova *et al.*, 2004).

Here, we aimed to identify the enzymes responsible for the biosynthesis of terminal Le^a structures on *P. patens* glycoproteins and to eliminate their function in rhEPO-producing moss lines in order to provide a Le^a-free asialo-EPO as a new interesting pharmaceutical target protein.

While $\alpha 1,4FucT$ is encoded by a single-copy gene in *P. patens*, 13 putative $\beta 1,3GalT$ homologues were identified from the moss genome sequence. On the basis of domain features and sequence similarity to *A. thaliana* GalTs, we took three of the *P. patens* *galt* genes under closer consideration. By generation of single $\Delta fuct$, $\Delta galt1$ and $\Delta galt3$ as well as $\Delta galt2/\Delta galt3$ double-KO lines, we intended to identify those enzymes relevant for Le^a formation in *Physcomitrella*.

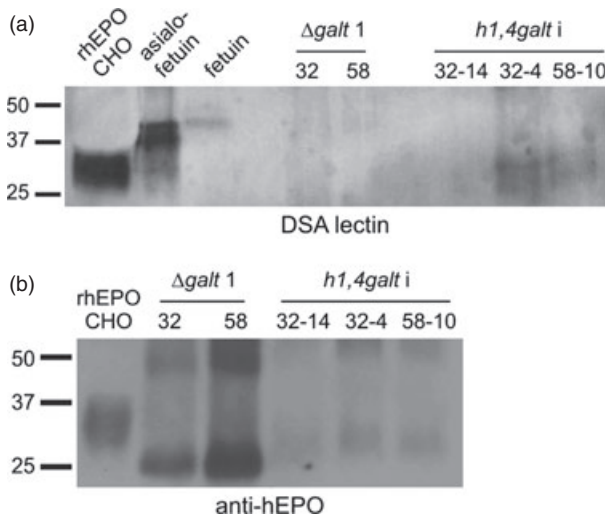


Figure 5 Lectin affinity blot analysis demonstrated specific terminal β 1,4 galactosylation of recombinant human erythropoietin (rhEPO) from *h1,4galt* insertion plants. (a) TSP were precipitated from culture supernatant of plants $\Delta galt1$ No. 32 and 58 and insertion lines *h1,4galt i* 32-14, 32-4 and 58-10. As positive controls, 200 ng of chinese hamster ovary (CHO)-produced rhEPO, 500 ng of β 1,4 galactosylated asialo-fetuin and 200 ng of β 1,4 galactosylated fetuin were loaded, and detection was performed with lectin *Datura stramonium* agglutinin. (b) As a control, CHO-produced rhEPO and culture supernatant of plants $\Delta galt1$ No. 32 and 58 and insertion lines *h1,4galt i* 32-14, 32-4 and 58-10 were analysed by anti-EPO Western blotting.

MALDI-TOF mass spectra analysis confirmed changes in the N-glycosylation pattern in the mutants generated in this work. In comparison to WT with Le^a epitopes on the N-glycans Gn(FA)XF_{iso} and (FA)(FA)XF (Figure 2a), $\Delta fuct$ plants exhibited the corresponding structures with the terminal galactose residues only, GnAXF_{iso} and AAXF, respectively. These structures have been only rarely observed on plant glycoproteins (Wilson, 2001). Steinkellner *et al.* (Strasser *et al.*, 2007) suggested two possible explanations: either the FucT enzyme modifies quickly and quantitatively the glycans with terminal galactose, or the Gal β 1-3GlcNAc oligosaccharide is rapidly degraded by the action of β -galactosidases. In the $\Delta fuct$ plants analysed here, we could clearly show an increase in terminal galactose compared to the WT (Figure 2a). This indicates that FucT modifies the Gal β 1-3GlcNAc substrate very efficiently. However, as the GnAXF_{iso}/AAXF proportion in $\Delta fuct$ is not identical to the Gn(FA)XF_{iso}/(FA)(FA)XF fractions of WT, we cannot exclude additional cleavage of terminal Gal residues.

While the disruption of *galt2* and *galt3* was obviously without any influence on terminal galactosylation, MALDI-TOF mass spectra from $\Delta galt1$ plant material revealed the absence of the complete Le^a epitope, confirming that the FucT enzyme needs the presence of the galactose to transfer a fucose to the GlcNAc residue.

In addition, immunoblotting performed using an antibody binding specifically to Le^a, JIM 84, showed the absence of this antigen from total secreted proteins of the $\Delta galt1$ plants. In contrast, Le^a epitopes were considerably high in the rhEPO-producing parent plant and the $\Delta galt2/\Delta galt3$ double-KO plants at the mass corresponding to moss-produced rhEPO.

A closer look at the glycopeptides of secreted rhEPO from the $\Delta galt1$ plants confirmed the lack of any Le^a epitopes. Moreover,

the high homogeneity of the product was striking, with almost exclusively one type of glycosylation, the desired GnGn structure. Here, a pharmaceutical product is presented with an excellent degree of uniformity.

In vitro activity of moss-derived rhEPO has already been proven (Weise *et al.*, 2007). However, moss-produced rhEPO lacked the terminal β 1,4 galactose present on asialo-EPO which was tested for neuro-protection (Erbayraktar *et al.*, 2003). Therefore, we engineered the moss genome for human β 1,4 galactosyltransferase activity (Figure 5) in a proof-of-concept approach and demonstrated substantial β 1,4 galactosylation of the secreted rhEPO in the resulting insertion lines. However, the degree of human-type galactosylation might be further increased using a hybrid instead of the native human β 1,4GalT (Strasser *et al.*, 2009).

The glycan engineering performed in the production lines presented in this work did not affect either biomass accumulation or rhEPO productivity; therefore, we suggest the moss bioreactor as the system of choice for the production of asialo-EPO as a new exciting biopharmaceutical target.

Experimental procedures

Identification of α 1,4 fucosyltransferase and β 1,3 galactosyltransferase in *Physcomitrella patens*

For the identification of α 1,4 fucosyltransferase homologues in *P. patens*, the amino acid sequence of the human galactoside 3(4)-L-fucosyltransferase (FUT3, NP_001091110.1) and the *A. thaliana* homologue At1g71990.1 were used to perform a BLASTP search in the *P. patens* resource (<http://www.cosmoss.org>). The sequence of the complete cDNA of the Pp*fuct* was obtained via RACE (GeneRacer™; Invitrogen, Karlsruhe, Germany) according to the manufacturer's protocol.

For the identification of Pp β 1,3 galactosyltransferase homologues, amino acid sequences of six human β 1,3 galactosyltransferases HsB3GalT1 (NP066191), HsB3GalT2 (AAG60610), HsB3GalT3 (AAQ89408), HsB3GalT4 (CAG33374), HsB3GalT5 (NP149363) and HsB3GalT6 (AAH51719) were used to perform a BLASTP search in <http://www.cosmoss.org>. Galactosyltransferase amino acid sequences from *A. thaliana* (At1g26810, At3g06440, At1g74800, At5g62620, At1g27120 and At4g21060) were taken from The Arabidopsis Information Resource database (TAIR, <http://www.arabidopsis.org>).

Plant material and transformation procedure

Physcomitrella patens (Hedw.) B.S.G. was cultivated as described previously (Frank *et al.*, 2005). The rhEPO-producing moss line 174.16 (Weise *et al.*, 2007) was used as starting material for the *galt* KOs. In this moss line, the α 1,3 fucosyltransferase and the β 1,2 xylosyltransferase genes are disrupted (Koprivova *et al.*, 2004). Protoplast isolation and polyethylene glycol-mediated transfection were performed as described previously (Frank *et al.*, 2005). For mutant selection, Knop media were supplemented with 50 μ g/mL G418, 50 μ g/mL Zeocin™ (Invitrogen) or 100 μ g/mL sulphadiazine (Sigma). The selection pressure started 14 days after transformation for G418 resistance and 9 days after transformation in the case of Zeocin™ and sulphadiazine selection, respectively.

For rhEPO production, *P. patens* was cultivated in 500 mL flasks with 200 mL basal medium prepared as follows: a solution containing 250 mg/L KCl, 250 mg/L MgSO₄·7H₂O, 1 g/L

Ca(NO₃)₂·4H₂O and 1 g/L 2-(N-morpholino)-ethanesulphonic acid (MES) was adjusted to pH 5.9 with 1 M KOH and autoclaved. Subsequently the following solutions were added per litre of this solution: 10 mL 25 g/L KH₂PO₄ (autoclaved separately), 1 mL 50 mM Fe-EDTA (sterile filtrated) and 1 mL microelement solution (614 mg/L H₃BO₃, 25 mg/L CoCl₂·6H₂O, 25 mg/L CuSO₄·5H₂O, 830 mg/L KI, 390 mg/L MnCl₂·4H₂O, 250 mg/L Na₂MoO₄·2H₂O, 25 mg/L NiCl₂·6H₂O, 10 mg/L Na₂SeO₃·5H₂O, 42 mg/L Zn acetate·2H₂O, pH 7.0) (sterile filtrated). The cultures were subcultivated weekly as described earlier (Reski and Abel, 1985) for at least 3 weeks. Afterwards, the cultures were kept 16 days without subculturing before the supernatant was harvested.

Creation of α 1,4 fucosyltransferase and β 1,3 galactosyltransferase KO constructs and human β 1,4 galactosyltransferase knockin constructs

To generate the *galt1* and *galt2* KO constructs, *P. patens* genomic DNA fragments corresponding to the respective genes were amplified using the primers listed in Table S1 and cloned into pCR[®]4-TOPO[®] (Invitrogen). The *galt1* genomic fragment was first linearized using *NheI* and *NdeI*, thus deleting a 1481-bp fragment, and recircularized by ligating a double-stranded oligonucleotide containing restriction sites for *Bam*HI and *Hind*III. These sites were used for the insertion of the Zeocin[™] selection cassette, obtained from the plasmid pUC-zeo. For creation of pUC-zeo, the *ble* gene was amplified from the vector pMS171 (Schroda *et al.*, 2002) with the primers listed in Table S1 and cloned into the vector pRT101-neo (Girke *et al.*, 1998) using *Xho*I and *Xba*I restriction sites to replace the *nptII* gene, resulting in pRT101-zeo. Subsequently, the Zeocin[™] selection cassette, including the 35S promoter and the 35S terminator, was removed from pRT101-zeo and ligated into the pUC vector (Vieira and Messing, 1982) via *Pst*I digestion, giving rise to the pUC-zeo vector. For the *galt2* KO construct, a 1093-bp fragment was deleted from the genomic sequence cloned into the pCR[®]4-TOPO[®] using *Hind*III and *Eco*RV sites and replaced by the Zeocin[™] selection cassette obtained from the pUC-Zeo plasmid using the flanking restriction sites *Hind*III and *Ecl*136II. In both KO constructs, the homologous regions flanking the selection marker were around 700 bp. The *galt3* KO construct was generated by PCR performed with genomic DNA of *P. patens*. Two fragments were amplified, representing the 5' part and the 3' part of the KO construct, respectively. Both amplified constructs were digested with *Hind*III followed by ligation using T4 DNA ligase. The resulting product comprised a deletion of 270 bp with regard to the genomic *galt3* sequence. Moreover, a stop codon had been introduced within the early 5' part of the corresponding cDNA. To generate the Δ *galt2*/ Δ *galt3* double KOs, the rhEPO-producing Δ *galt3* mutant was transformed with the *galt2* KO construct.

The KO construct for *fuct* was synthesized by PCR amplification of a 2860-bp genomic fragment with the primers listed in Table S1. Then, the *nptII* cartridge was amplified by PCR introducing the *Bst*Z17I restriction sites and was cloned into the *Bst*Z17I in the *fuct* genomic fragment.

The human β 1,4*galt* knockin construct was created by replacement of the Zeocin[™] selection cassette from the *galt1* KO construct with the human β 1,4*galt* expression cassette from plasmid pCR4-XTko-GTKi (Huether *et al.*, 2005), resulting in the

*galt1*KO/*h1,4galt*KI construct. To generate *h1,4galt* insertion lines, Δ *galt1* lines 32 and 58 were co-transfected with either the *galt1*KO/*h1,4galt*KI construct or pCR4-XTko-GTKi together with pUCsul, which confers sulphadiazine resistance (Markmann-Mulisch *et al.*, 2007).

Screening of transgenic plants

Screening for transformants with the 5' and 3' integration of the KO construct in the correct genomic locus was performed via direct PCR (Schween *et al.*, 2002) with genomic DNA extracted using a protocol modified from Edwards *et al.* (1991). Briefly, approximately 5 mg of plant material was homogenized for 1 min in a bead mill (Qiagen, Hilden, Germany) with 200 μ L of extraction buffer (200 mM Tris-HCl, pH 7.5; 250 mM NaCl; 25 mM EDTA; and 0.5% SDS), and the sample was then shortly vortexed and centrifuged at 20 000 *g* for 1 min. Subsequently, 150 μ L of the supernatant was transferred to a fresh tube, mixed with 150 μ L cold (–20 °C) isopropanol and incubated for 2 min. After centrifugation at 20 000 *g* for 5 min at 4 °C, the pellets were washed with 200 μ L cold 70% ethanol, dried at room temperature and resuspended in 30 μ L TE buffer. From these extracts, 2 μ L was used as template for PCR. Plants that in this genomic DNA screening showed the expected PCR products were considered as putative KOs and selected for further analysis.

The absence of the glycosyltransferase transcripts in the KO lines and expression of *h1,4galt* in the *h1,4galt* i lines was analysed via RT-PCR. For this purpose, RNA was extracted from approximately 100 mg fresh weight plant material with TRIzol[®] reagent (Invitrogen) following the manufacturer's recommendations. Synthesis of cDNA was performed with M-MuLV reverse transcriptase (Fermentas, St Leon-Rot, Germany) using 2 μ g RNA previously treated with DNase I (Fermentas), and cDNA quality was checked by PCR with primers C45f and c45r which amplify the constitutively expressed gene of the ribosomal protein L21. The specific transcripts were amplified with the primers listed in Table S2.

Protein and N-glycan analysis

Total soluble proteins were recovered from 160 mL of 7- to 16-day-old culture supernatant by precipitation with 10% (w/v) trichloroacetic acid (TCA; Sigma-Aldrich, Deisenhofen, Germany) as described earlier (Büttner-Mainik *et al.*, 2011). The pellet was resuspended in sample loading buffer (Laemmli, 1970) without bromophenol blue, and total proteins were measured by means of the bicinchoninic acid assay (BCA protein assay kit; Pierce, Rockford, IL), following the manufacturer's recommendations. Bromophenol blue was added later to the samples, and electrophoretic separation of proteins was carried out in 12% SDS-polyacrylamide gels (Ready Gel Tris-HCl; BioRad, Munich, Germany) at 150 V for 1 h under non-reducing conditions.

For glycopeptide analysis, the proteins in the gels were stained with PageBlue[®] Protein Staining Solution (Fermentas), and the bands corresponding to 25 kDa were cut out, S-alkylated and digested with trypsin. Analysis by reversed-phase liquid chromatography coupled to electrospray ionization mass spectrometry (LC-ESI-MS) was performed as described (Grass *et al.*, 2011).

For hEPO Western blot analyses, 100 ng of proteins was loaded for SDS-PAGE as described above, and for Lewis A Western blots, 30 μ g of proteins was used. Gels were then blotted

to polyvinylidene fluoride (PVDF) membranes (Immobilon-P; Millipore, Bedford, MA) in a Trans-Blot SD Semi-Dry Electrophoretic Cell (Bio-Rad) for 1 h with 1 mA/cm² membrane. For the hEPO Western blot, a mouse anti-hEPO monoclonal antibody (MAB2871; R&D Systems, Minneapolis, MN) and an anti-mouse antibody coupled with peroxidase (NA 931V; GE Healthcare, München, Germany) in 1 : 4000 and 1 : 150 000 dilutions, respectively, were used. As standard, 20 ng rhEPO (cat. no 287-TC; R&D Systems) was loaded. For the anti-Le^a immunoblot, a 1 : 10 dilution from the JIM 84 antibody (kindly provided by Prof. Hawes, Oxford Brookes University) and a 1 : 50 000 dilution from a rabbit anti-rat antibody (ab 6250; Abcam, Cambridge, MA) were used. Detection was performed using the ECL Advance Western Blotting Detection kit (GE Healthcare) following the manufacturer's instructions.

For lectin affinity blots, SDS-PAGE and blotting to PVDF membranes were performed as described above. Terminal Galβ1,4GlcNAc residues on blotted glycoproteins were detected with digoxigenin-labelled DSA (DIG Glycan Differentiation kit; Roche, Mannheim, Germany) according to the manufacturer's instructions.

Quantification of the moss-produced rhEPO was performed using a hEPO Quantikine IVD ELISA kit (cat. no DEP00; R&D Systems) according to the manufacturer's protocol.

Structural analysis of N-glycans was performed by MALDI-TOF MS of enzymatically released N-glycans that were isolated as described in detail elsewhere (Altmann *et al.*, 2001).

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Supporting information

Additional Supporting information may be found in the online version of this article:

Figure S1 Schematic representation of the *fuct* and *galt* corresponding knockout constructs and molecular analysis of the knockout mutants.

Figure S2 Extracellular proteins from rhEPO producing moss.

Figure S3 Expression analysis of human $\beta 1,4galt$ in the putative *h1,4galt* insertion lines.

Table S1 Oligonucleotides used for DNA amplification for KO constructs cloning.

Table S2 Oligonucleotides used for glycosyltransferases transcript screening.

Table S3 Quantification of moss produced rhEPO.

Table S4 International Moss Stock Center numbers of plants used in this work.

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