Introduction

Glycosylation is considered the most abundant post-translation al modification and occurs when oligosaccharides (glycans) are covalently attached to other biomolecules such as proteins or lipids [1]. Glycosylation can have a prominent role in several fundamental biological processes such as embryogenesis, development, growth, differentiation and also in diseases such as cancer [2,3,4]. Glycosylation is catalyzed by several glycosyltransferases that act in a step-wise manner. Functional glycomics, in particular, is currently allowing the understanding of the role played by glycosyltransferases and glycans in cell biology and function of organisms [5].

N-acetylglucosaminyltransferase III (GnT-III) is a glycosyltransferase encoded by the Mgat3 gene that catalyzes the transfer of N-acetylglucosamine (GlcNAc) in a 1,4 linkage to mannose on N-glycans forming a bisecting GlcNAc structure [6]. GnT-III is considered a key glycosyltransferase in N-glycan biosynthetic pathway since the introduction of the bisecting GlcNAc residue precludes further processing and elongation of N-glycans catalyzed by GnT-V, suppressing the formation of β1,6 GlcNAc branching structures [7]. In a tumour context, GnT-III and GnT-V have generally a dual role where GnT-III and its bisecting GlcNAc structures act as metastases suppressors whereas GnT-V and its β1,6 GlcNAc branching structures are associated with increased malignancy and metastasis [8].
The detailed molecular mechanism and the most important molecular targets underlying this antagonistic role of Gnt-III and Gnt-V in cancer development and progression have recently been addressed [8,9,10]. Focusing on the role of GnT-III as an important metastases suppressor, Yoshimura et al. have demonstrated that Gnt-I-III gene transfected into B16 mouse melanoma cells with high metastatic capacity led to a suppression in the formation of α1,6 GlcNAc branching structures catalyzed by Gnt-V, together with a significant decrease in lung colonization after mouse intravenous administration of Gnt-I-III transfectants [11]. Further on, the same group showed that in those Gnt-I-III transfectants from mouse melanoma cells, E-cadherin was modified with bisecting GlcNAc structures showing a delayed turnover rate and an increased cell-cell adhesion, which might explain the metastatic impairment induced by Gnt-I-III overexpression [12]. The anti-metastatic role of Gnt-I-III was also described by Isaji et al., in a report showing that modification of Integrin α5β1 with bisecting GlcNAc structures, catalyzed by Gnt-I-III, inhibits cell spreading and migration on fibronectin [13].

According with these results, it was proposed that Gnt-I-III prevents tumour metastasis by at least two mechanisms: enhancement of cell-cell adhesion (through E-cadherin glycosylation) and down-regulation of cell-ECM adhesion (through Integrin glycosylation) [9]. Moreover, it was also demonstrated that the addition of bisecting GlcNAc structures catalyzed by Gnt-I-III to mammary tumour cell glycoprotein receptors inhibits growth factor signalling reducing growth and retarding mammary tumour progression [14]. Recently, our group has proposed a mechanistic model where E-cadherin expression regulates the Mgat3 gene transcription leading to increased expression of Gnt-I-III enzyme that in turn glycosylates E-cadherin by adding bisecting GlcNAc structures [15], which promotes its stability and function at the cell membrane (unpublished results). Interestingly, when this functional feed-back loop between E-cadherin and Gnt-I-III is disturbed, by performing Gnt-I-III knockdown, we observed a dramatic alteration in the cell morphology with formation of filopodia and lamellipodia extensions, together with a delocalization of E-cadherin from the cell membrane into the cytoplasm [15].

E-cadherin is a well-accepted marker of phenotypic plasticity, and is the central target and the most common endpoint of Epithelial to Mesenchymal Transition (EMT) signalling pathways [17]. This is valid for EMT, but also to the apparent reverse process known as Mesenchymal to Epithelial Transition (MET), that occurs during embryonic development, tissue regeneration, wound-healing and thought to occur in cancer initiation/progression [18,19,20]. The most commonly used epithelial molecular markers include E-cadherin, occludin, cytokeratins, whereas N-cadherin and vimentin are classically considered mesenchymal markers [21]. EMT is also characterized by increased production of transcription factors such as Snail, Slug, Twist, ZEB1, ZEB2, and/or E47, some of which also known as EMT inducers. During EMT/MET, cells alternate between pure epithelial and mesenchymal phenotypes characterized by modifications in cell adhesion, polarity, migration and cell shape [16,17]. This EMT/MET associated cell plasticity is mirrored by many features induced by overexpression of Gnt-I-III (associated with epithelial phenotype) and Gnt-V (associated with mesenchymal features) [15,22].

Altogether, some reports as well as our previous results show an effect of Gnt-I-III mediated glycosylation on cellular phenotype and E-cadherin cellular expression [10,15,23]. We therefore hypothesize that during EMT/MET, E-cadherin could be posttranslationally regulated by Gnt-I-III-mediated glycosylation.

In the present study, we analyzed the expression profile and the genetic mechanism controlling Mgat3 gene expression as well as the Gnt-I-III-induced glycosylation levels, in general and specifically on E-cadherin, during EMT/MET. We found that during EMT, Mgat3 gene suffered a dramatic decrease in expression that was significantly recovered when cells re-acquired an epithelial-like phenotype (MET). E-cadherin was specifically targeted and regulated by Gnt-I-III mediated glycosylation during EMT/MET. We propose here, for the first time, Mgat3/Gnt-I-III mediated glycosylation as a novel and major mechanism of E-cadherin regulation during EMT/MET.

**Results**

TGF-β1 induces EMT in EpH-4 non-tumourigenic mammary epithelial cells and TGF-β1 removal initiates the MET program

The EpH-4 cell line derives from a normal mammary gland of a mid-pregnant BALB/c mouse that underwent spontaneously immortalization [24]. To induce EMT, we have selected a previously described approach, using a pleiotropic cytokine, in particular the transforming growth factor-β1 (TGF-β1) [25]. TGF-β1 supply during 7 days efficiently induced EMT in original epithelial EpH-4 cells (E), and thus mesenchymal EpH-4 cells (M) were generated (Figure 1A). Removal of TGF-β1 from the culture medium led to phenotypic reversion from mesenchyme back to epithelia (RE), after four days. The MET process was terminated at day four after which bright field microscopy revealed a widespread recovery of the epithelial-like phenotype (Figure 1A).

Three distinct time-points were created for biological material collection (DNA, RNA, protein and fixed cell monolayers) corresponding to EpH-4 cells with distinct phenotypic features (E, M and RE). To understand whether the observed phenotypic alterations were due to EMT and MET induction, we next analysed the mRNA expression by qRT-PCR of several classical epithelial (CDH1 and Ocln) and mesenchymal (Vim) markers, as well as a well-known EMT inducer (Zeb2). We have observed a slight decrease in CDH1 mRNA expression upon EMT, which was recovered with MET (Figure 1B). Ocln, encoding occludin, was shown to decrease with EMT (E vs. M p = 1.34E-05), being recovered in MET (M vs. RE p = 4.07E-02). The opposite variation was observed for Vim encoding vimentin (E vs. M p = 1.60E-02 and M vs. RE p = 5.20E-02) (Figure 1B). Zeb2 mRNA showed a significant increase in EMT (E vs. M p = 1.91E-02) and no significant variation in MET (Figure 1B).

Given that CDH1 mRNA expression was only slightly decreased when comparing EpH-4 E and M cells, we next analyzed the expression of the encoded protein, E-cadherin, by western blot. We observed that E-cadherin expression decreased with EMT and was partially recovered with MET (Figure 2A and 2B), however these variations were not statistically significant. By immunofluorescence, we observed an E-cadherin de-localization from the cell membrane (Figure 2A and 2C). mRNA expression was only slightly decreased when comparing EpH-4 E and M cells, we next analyzed the expression of the encoded protein, E-cadherin, by western blot. We observed that E-cadherin expression decreased with EMT and was partially recovered with MET (Figure 2A and 2B), however these variations were not statistically significant. By immunofluorescence, we observed an E-cadherin de-localization from the cell membrane (Figure 2A and 2C).
Figure 1. EMT/MET model validation. Panel A shows bright field microscopy still images of the EpH-4 cell line (400×) during EMT and MET induction. E stands for epithelial cells, M stands for mesenchymal cells, and RE stands for reverted epithelial cells. E and RE cells display a cuboid shape associated with the epithelial phenotype while M cells exhibit a fibroblastic morphology associated with the mesenchymal phenotype. Panel B illustrates the quantification of CDH1, Ocln, Vim and Zeb2 relative mRNA expression (n = 3 biological replicas). Data was normalized for results obtained for E cells. Single asterisk corresponds to p≤0.05 and double asterisks stands for p≤0.001. CDH1 expression, an epithelial marker, was not significantly altered during EMT/MET induction; Ocln expression, an epithelial marker, was significantly decreased in M cells (in comparison to E cells) and recovered in RE cells (in comparison to M cells); Vim expression, a mesenchymal marker, was significantly increased in M cells (in comparison to E cells) and decreased in RE cells (in comparison to M cells) and; Zeb2 expression, a classical EMT inducer, was significantly increased in M cells in comparison with E cells, supporting EMT occurrence.

doi:10.1371/journal.pone.0033191.g001

Figure 2. EMT/MET model validation. Panel A shows the Western blot for E-cadherin. Panel B illustrates the quantification of E-cadherin across the EMT/MET induction (n = 3 biological replicas). Data was normalized for E cells. Results are described as mean±standard error mean of 3 biological replicas. No significant differences were observed concerning E-cadherin expression (ns stands for non-significant, p>0.05). Panels A and B show that E-cadherin expression is decreased in M cells (in comparison to E cells) and partially recovered in RE cells (in comparison to M cells). Panel C represents the immunofluorescence for E-cadherin during EMT/MET induction (200×). NC stands for negative control (no E-cadherin antibody used). Panel C illustrates the variation of E-cadherin localization during the EMT/MET induction: E cells display the classical E-cadherin expression at the cell membrane; M cells show a decreased expression of E-cadherin which is only observed in some points of intercellular contacts and in the cytoplasm; RE cells display E-cadherin expression in the cell membrane.

doi:10.1371/journal.pone.0033191.g002
panied by an epithelial transcriptional program together with a recovery of E-cadherin at its classical membranous localization.

*Mgat3* gene expression is significantly down regulated during EMT being significantly recovered in MET and is associated with *Mgat3* promoter methylation/demethylation.

*Mgat3* mRNA expression was evaluated by qRT-PCR during EMT/MET. We observed a significant decrease in *Mgat3* mRNA expression upon EMT ($p = 8.10^{-04}$), followed by a full recovery upon MET ($p = 1.25E^{-02}$) (Figure 3A). To understand whether this down-regulation could be related with a classical inactivating mechanism, we have analysed the methylation status of *Mgat3* promoter region.

Due to the lack of a described mouse *Mgat3* promoter in the literature, we pursued the classical approach of analysing the bioinformatically predicted CpG islands within the gene locus, given the known association between CpG islands and promoter regions [26] (Figure 3B). We observed that the methylation pattern of *Mgat3* CpG island 1 was significantly altered with EMT (Figure 3C); 1) three CpG sites became completely methylated (CpG sites 4, 7 and 8); 2) two CpG sites became completely demethylated (CpG sites 3 and 11). We also observed that the CpG island 1 methylation pattern observed for EpH-4 cells E and RE reproduced the *Mgat3* mRNA expression levels (Figure 3A and C). Concerning CpG island 2, no significant alterations were detected during the EMT/MET. Overall this CpG island was permanently methylated throughout the experiment (Figure 3D). To confirm that methylation of *Mgat3* promoter was a specific feature of this gene, potentially leading to its expression downregulation, we also analysed the methylation status of the promoter of *Mgat5*, encoding the GnT-V glycosyltransferase, for which RNA expression did not vary during the EMT/MET experiment (Figure 3E and F). The methylation status of *Mgat5* did not change along EMT/MET (Figure 3G).

Figure 3. *Mgat3* and *Mgat5* RNA expression and methylation status of their predicted promoter-associated CpG islands. Panel A illustrates the quantification of *Mgat3* relative mRNA expression ($n=3$ biological replicas). Data was normalized for E cells for each biological replica. Single asterisk corresponds to $p \leq 0.05$ and double asterisks stands for $p \leq 0.001$. Results are described as mean ± standard error mean of three biological replicas. Panel A shows that *Mgat3* expression was significantly decreased in M cells (in comparison to E cells) and recovered in RE cells (in comparison to M cells). Schematic representation of *Mgat3* genomic locus is represented in panel B. White squares correspond to exonic untranslated regions and black squares to exonic translated regions. Black line stands for intronic regions. Grey squares represent the position of the bioinformatically predicted CpG islands (classified as 1 and 2). Panel C and D show the schematic representation of the methylation status of several CpG dinucleotides evaluated within CpG islands 1 (C) and 2 (D) of *Mgat3* across the EMT/MET experiment (E, M and RE). White circles correspond to unmethylated CpGs, grey circles correspond to partially methylated CpGs, black circles correspond to methylated CpGs, white circles with a question mark correspond to unknown methylation status. Panel C shows methylation pattern alterations across several CpG sites within *Mgat3*'s CpG island 1 in E, M and RE cells. Panel E illustrates the quantification of *Mgat5* relative mRNA expression ($n=3$ biological replicas). Same legend as in A applies. No significant variation of *Mgat5* RNA expression was observed during EMT/MET. Schematic representation of part of the *Mgat5* genomic locus is represented in panel F. Same legend as in panel B applies. Schematic representation of the methylation status of several CpG dinucleotides evaluated within the annotated *Mgat5* CpG island is represented in panel G. Same legend as in panels C and D applies. The results showed no variation of the methylation status of *Mgat5* promoter during EMT/MET.

doi:10.1371/journal.pone.0033191.g003
These results highlighted a novel regulatory mechanism controlling \textit{Mgat3} expression involving \textit{G6} island promoter methylation associated with EMT and MET.

**Evaluation of the levels of expression and cellular localization of bisecting GlcNAc structures during EMT/MET**

Taking into consideration the results at the transcriptional level of \textit{Mgat3} gene during EMT/MET, we analyzed the product of the activity of the \textit{GnT-III} enzyme, the bisecting GlcNAc structures (Figure 4). We performed E-PHA lectin blot analysis, which specifically recognizes the product of \textit{GnT-III}. The results showed that the total levels of expression of bisecting GlcNAc structures significantly decrease during EMT ($p = 3.3E-03$), with E cells displaying a significant increased expression of bisecting GlcNAc structures than M cells (Figure 4A, 4B). The total levels of expression of bisecting GlcNAc structures was dramatically recovered in RE cells when compared with M cells ($p = 8.0E-04$).

The evaluation of the cellular localization of the product of \textit{GnT-III} (bisecting GlcNAc structures), by performing E-PHA lectin IF, clearly showed that during EMT there was a remarkable decrease in the levels of expression of these structures, as observed in M cells, that were recovered in RE cells. In terms of cellular localization, the expression of the \textit{GnT-III} product in E cells was mainly in the cell membrane with rare cytoplasmic staining; in M cells very few E-PHA lectin staining was observed restricted at the cytoplasm in a perinuclear location, and in RE cells E-PHA lectin staining was observed both at the cell membrane and cytoplasm (Figure 4C).

These results showed that high levels of expression of bisecting GlcNAc structures are associated with epithelial phenotypes (E, RE) and that these structures localize at the cell membrane only in these phenotypes.

**E-cadherin is a specific target of regulation by GnT-III mediated glycosylation during EMT/MET**

As the expression of bisecting GlcNAc structures and E-cadherin was observed at the cell membrane in epithelial phenotypes (E and RE), we next evaluated the co-localization of both molecules by performing double-labelled immunofluorescence (Figure 5A). The results showed that in both E and RE cells there was a co-localization of bisecting GlcNAc structures (green colour) and E-cadherin (red colour) at the cell membrane. During EMT, when cells acquire a mesenchymal phenotype, the co-localization disappears and cells show few bisecting GlcNAc structures in the cytoplasm in a perinuclear position (that may correspond to the Golgi apparatus), whereas E-cadherin expression is limited only to the focal points of intercellular contact (Figure 5A).

![Figure 4](https://plosone.org/images/article/figure/1.png)

**Figure 4. Expression levels and cellular localization of the product of GnT-III enzyme during EMT/MET induction.** Panel A shows the lectin blot analysis using E-PHA lectin, showing the total expression levels of bisecting GlcNAc structures during EMT/MET. Panel B illustrates the quantification of E-PHA lectin normalized to actin ($n = 2$ biological replicas). Single asterisk corresponds to $p = 0.05$ and double asterisks stands for $p = 0.001$. Bisecting GlcNAc structures significantly decrease when comparing E and M cells and their expression is significantly recovered in RE cells. Panel C represents the immunofluorescence for E-PHA lectin during EMT/MET induction (400×). Bisecting GlcNAc structures are preferentially localized in the cell membrane of E cells. M cells exhibit a clear decrease in expression of the bisecting GlcNAc structures that was only observed in focal areas in the perinuclear region. In RE cells, there was a significant increase in the E-PHA staining showing an increase in the expression levels of bisecting GlcNAc structures that are localized in the cell membrane and in the cytoplasm.

doi:10.1371/journal.pone.0033191.g004
We next examined whether the EMT/MET marker, E-cadherin, would be a target of regulation by the GnT-III glycosyltransferase (Figure 5B, 5C). For doing so, we performed E-cadherin IP followed by E-PHA lectin blot analysis to evaluate the specific modification of E-cadherin with bisecting GlcNAc structures, catalyzed by GnT-III during EMT/MET. We performed a normalization of the densities between E-PHA lectin band and the correspondent E-cadherin band in order to allow the comparison of the levels of E-cadherin glycosylation by GnT-III along EMT/MET. The results clearly demonstrated that during EMT E-cadherin suffered a significant decreased modification with bisecting GlcNAc structures of about 50% \((p = 5.1E-02)\) (Figure 5C). On the contrary, during the reverted process (MET) E-cadherin glycosylation with bisecting GlcNAc structures was recovered for levels on average higher than those of the starting point (parental epithelial cells).

These results demonstrate that E-cadherin is being specifically glycosylated by GnT-III and that this post-translational modification varies significantly during EMT/MET.

**Discussion**

The transition between epithelial and mesenchymal states requires alterations in cell morphology, cellular architecture, adhesion, and migration capacity [16,17]. Current interest in these processes (EMT/MET) stem from their developmental importance and their involvement in several adult pathologies including cancer [21], therefore it is crucial to unravel the mechanisms involved in these transitions.

In the present manuscript, we describe for the first time that transcription of *Mgat3* gene, that encodes the GnT-III glycosyltransferase, significantly varies along EMT/MET and that its expression is controlled by promoter methylation. Further, we also showed that the product of GnT-III, the bisecting GlcNAc structures, specifically glycosylates and regulates E-cadherin functions.

GnT-III, has been considered a key glycosyltransferase during the N-glycan biosynthetic pathway [6], and has been also pointed as a metastases suppressor gene [11]. This role in the suppression of tumour invasion and metastases appears to be through glycosylation and regulation of a critical tumour suppressor gene, E-cadherin [12,15,27], as well as through regulation of Integrin-mediated cell-ECM adhesion [13].

Previously, we have described the existence of a functional feed-back loop between E-cadherin and GnT-III, where we found that E-cadherin expression regulates the transcription of *Mgat3* gene, which in turns leads to glycosylation of E-cadherin with bisecting GlcNAc structures [15], promoting its cell surface stability and function (unpublished results). The perturbation of this stable loop either by GnT-III knockdown or by increased competition with GnT-V enzymatic activity was shown to lead to alterations in cell morphology and E-cadherin subcellular localization [15]. These results pointing to a regulation of E-cadherin function through glycosylation [10], led us to postulate that *Mgat3* expression and GnT-III enzymatic product could interfere with E-cadherin during EMT/MET, a process in which changes in E-cadherin function are described to be pivotal [16,17,28].
Using a TGF-β1-induced EMT/MET cell line model, we describe for the first time that Mgat3 gene was significantly downregulated in mesenchymal (M) cells and was significantly recovered in cells that re-acquired an epithelial-like phenotype after removing TGF-β1 (RE cells). This suggests that Mgat3 gene expression and the GnT-III enzymatic product may be critical to maintain the cellular epithelial and differentiated phenotype, while the sharp drop at the mesenchymal state may pinpoint the need for Mgat3 down regulation to the acquisition of this phenotype. Moreover, we have also provided compelling evidence of increased CpG island methylation at the Mgat3 promoter, underlying the down regulation of Mgat3 expression in M cells (Figure 6).

Changes in glycogenes expression leads to alterations in the glycosyltransferases expressions levels that have major impact in the remodelling of cell surface glycosylation of glycoproteins, with consequent impact on several cellular mechanisms such as cell adhesion [5,29]. The observed similar expression variation of the Mgat3 gene and the product of GnT-III, the bisecting GlcNAc structures, along EMT/MET further support that, on one hand the expression of bisecting GlcNAc structures is crucial for the preservation of epithelial cell phenotype, as previously described by Iijima et al [27], and the absence or low levels of bisecting GlcNAc structures are needed for cells to acquire mesenchymal features.

Moreover, our current results are consistent with our previous report that MCF-7/AZ cells with Mgat3 knockdown, displayed a remarkable modification of the cellular phenotype with disruption of the cell-cell contacts, increased lamellipodia and filopodia formation and E-cadherin internalization to the cytoplasm (mesenchymal features) [15].

Given the previously described bi-directional crosstalk between E-cadherin and GnT-III glycosylation [15], in the present study we examined the specific modification of E-cadherin with bisecting GlcNAc structures during EMT/MET. We found that, although the variation of the CDH1/E-cadherin during EMT/MET was not significant, E-cadherin was glycosylated with bisecting GlcNAc glycoforms and they co-localized at the membrane preferentially in E and RE cells.

Altogether, the results herein presented, lead us to propose that loss and recovery of Mgat3 expression and GnT-III mediated glycosylation is a pivotal mechanism during EMT/MET with major consequences on E-cadherin subcellular localization. We showed a novel perspective over the regulation of E-cadherin function along EMT/MET by demonstrating that E-cadherin glycosylation with bisecting GlcNAc structures, catalysed by GnT-III enzyme, has major consequences in the regulation of its function in the EMT/MET biological processes.

These cause-consequence events, involving changes in Mgat3 expression, GnT-III enzymatic product, E-cadherin protein glycosylation and subcellular localization are likely to uncover a mechanism controlling the transition between epithelial and mesenchymal states, therefore revealing a novel regulatory path during EMT/MET (Figure 6).

![Figure 6. Proposed model.](image-url)
Materials and Methods

TGF-β-induced EMT/MET in a non-tumourigenic mammary epithelial cell line

EpH-4 cell line [24] (classified as E for epithelial) was cultured in D-MEM/F12 GlutamaxTM (Invitrogen, Oregon, USA) supplemented with fetal bovine serum (Lonza, Switzerland), penicillin-streptomycin (Invitrogen, Oregon, USA) and recombinant human insulin (Invitrogen, Oregon, USA). Mesenchymal cell cultures (classified as M) were obtained by supplementing the normal culture medium with transforming growth factor-β1 (TGF-β1, Sigma-Aldrich-Aldrich, Missouri, USA) during 7 days [25]. Reverted-epithelial cell cultures (classified as RE) were obtained by replacing the TGF-β1 enriched medium in mesenchymal cell cultures by normal culture medium for another 4 days.

RNA expression quantification

RNA was extracted from 3 biological replicates of EpH-4 cell line stages (E, M and RE) using the mirVana miRNA Isolation Kit (Invitrogen, Oregon, USA), according to the kit’s instruction manual. Using TaqMan Gene Expression Assays (Applied Biosystems, California, USA), we have quantified the mRNA expression levels of Mgat3, CDH1, Ocn, Zeb2, and Vim. Approximately 1000 ng of total RNA were reversed transcribed to single stranded cDNA using Superscript II Reverse Transcriptase and random hexamer primers (Invitrogen, Oregon, USA). Quantitative Real-Time-PCR (qRT-PCR) was carried out in triplicates using source RNA from 3 biological replicates of the EMT/MET experiment, for the target genes Mgat3, CDH1, Ocn, Zeb2, Vim and for the endogenous control GAPDH using as probe sets Mm00447798_s1, Mm_00496909_g1, Mm.PT.47.16166845, Mm.PT.47.13169136, Mm.01333430.m1 and 4352932E (Applied Biosystems, California, USA and Integrated DNA Technologies, Iowa, USA) respectively and an ABI Prism 7000 Sequence Detection System. Data was analysed by the comparative 2-DDACT method [30]. For all data comparisons, the Student’s T-Test was used (two tailed, unequal variance).

Mgat3 and Mgat5 methylation status analysis

DNA from 3 independent biological replicates of EpH-4 cells (E, M and RE) was extracted using the kit Invisorb Spin Tissue Mini Kit following the manufacturer’s instructions (STRATEC Molecular, Berlin, Germany). Approximately 300 ng of DNA from each and every sample were then subjected to complete bisulfite conversion and subsequent cleanup using the Epitect Bisulfite Kit following manufacturer’s instructions (Qiagen, Hilden, Germany). Bisulfite treated DNA was amplified using primers flanking the Mgat3 promoter CpG island (Sigma-Aldrich-Aldrich, Missouri, USA), specifically designed for bisulfite treated DNA sequences without CpG sites. The bisulfite PCR products were sequenced for methylation status determination. Mgat3 promoter methylation analysis was carried out within the two CpG islands bioinformatically predicted to exist within Mgat3 genomic locus [26,31]. Mgat5 promoter methylation analysis was carried out within the single CpG island bioinformatically predicted to exist within Mgat5 genomic locus [26,31]. The criteria used for CpG island prediction was as follows: 1) genomic area with ≥500 bp; 2) a percentage of GC ≥55 and; 3) the observed/expected CpG dinucleotides ≥0.65. Using the Ensembl database [31] and the web tool “CpG Island Searcher” [26] and the described criteria, two distinct CpG islands were predicted within/in the vicinity of Mgat3 genomic locus: CpG island 1, located at chr15: 80003669-80004712; CpG island 2, located at chr15: 80041789-80042386. In particular, the methylation status was analysed at 13 CpG sites for CpG island 1 (chr15: 80004555–80004661) and at 15 CpG sites for CpG island 2 (chr15: 80042610–80042920). Concerning Mgat5, the predicted CpG island was located at chr1:129.101.924–129.102.322. In particular, the methylation status was analysed at 17 CpG sites (chr1:129.101.638–129.101.814). Only results defining the findings obtained for all biological and technical replicates are herein presented.

Western-blot, Immunoprecipitation and Lectin Blot Analysis

EpH-4 cells (E, M and RE) were washed with PBS and then lysed in cold PBS containing 1% TritonX-100, 1% NP40, protease inhibitor cocktail (Roche 1tablet/50 ml buffer) and phosphatase inhibitor cocktail (Sigma-Aldrich, 1:100 dilution). Total protein was quantified using a BCA protein assay kit (Pierce).

Equal amounts of total cell protein lysates (25 µg) from each cell were subjected to 7.5% SDS-PAGE electrophoresis and then transferred to nitrocellulose membranes. After blocking with 5% nonfat milk (for western-blot) or with 5% BSA in PBS (for lectin blot), the membranes were incubated with primary antibodies against human E-cadherin (clone 36, BD Transduction Laboratories and Cell Signaling). For the lectin blot analysis, membranes were incubated with E-PHA lectin (Vector Laboratories). Next, membranes were washed four times with PBST solution, followed by incubation with horseradish peroxidase-linked secondary antibody for the E-cadherin blot analysis. For the lectin blots, the immunoreactive bands were visualized using the Vectorstain ABC kit (Vector Laboratories). Blots were then probed with anti-actin antibody (Santa Cruz Biotechnology) for loading control analysis. Results were obtained from two independent experiments using two different biological replicates.

For immunoprecipitation analysis, equal amounts of total protein from each cell lysate were precleared with 25 ml of protein G-Sepharose beads (Sigma-Aldrich) for 1–2 h. After centrifugation, the supernatant was incubated overnight with 5 µg of monoclonal antibody against E-cadherin (BD Biosciences and Cell Signalling). After that, incubation with protein G-Sepharose for 2 h was performed. Next, the beads were washed three times with immunoprecipitation buffer. The immune complexes were released by boiling for 5 min at 95°C in Laemmli sampling buffer and the immunoprecipitates were subjected to 7.5% SDS-PAGE; the separated proteins were transferred to a nitrocellulose membrane. The blots were then probed with the primary antibody against E-cadherin. For the biotinylated GlnAc analysis on E-cadherin IP, membranes were probed with E-PHA lectin. The results were visualized with ECL detection reagent (Amersham Biosciences). The experiment was reproduced three times using protein extract from two different biological replicates. For all data comparisons, the Student’s T-Test was used (two tailed, unequal variance).

Immunofluorescence

EpH-4 cells at the three different stages (E, M, RE) were plated on six-well plates with coverslips on the bottom of each well. Cells from each stage (E; M; RE) were fixed with Methanol and blocked with BSA 10% in PBS.

For E-cadherin staining, cells were incubated with anti-E-cadherin monoclonal antibody (BD Biosciences and Cell Signalling) for 1 h at room temperature. After three washes with phosphate-buffered saline (PBS), cells were incubated with Alexa Fluor 488 anti-mouse secondary antibody (Invitrogen) for 1 h at room temperature.

For E-PHA staining and in the same way, EpH-4 E, M and RE cells were incubated for 1 h at room temperature with biotinylated
Phasolus vulgaris erythroagglutinin (E-PHA, which binds to biecting GlcNAc structures, product of GnT-III) lectin (Vector Laboratories). Following three washes with PBS, the cells were incubated with streptavidin-FITC for 1 h at room temperature.

For E-cadherin and E-PHA double-staining a double-labeled immunofluorescence, E/PHA E, M and RE cells were incubated with anti-E-cadherin monoclonal antibody (BD Biosciences and Cell Signalling followed by Alexa Fluor 594 anti-mouse secondary antibody. Then cells were incubated with E-PHA biotinylated lectin followed by streptavidin-FITC incubation.

The nuclear staining for all the immunofluorescence experiments was performed and images were then visualized in a fluorescence microscope (Zeiss).

Each experiment was reproduced two times using protein extract from two different biological replicas.

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